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(71) Applicant: **PRESIDENT AND FELLOWS OF HARVARD COLLEGE** [US/US]; 17 Quincy Street, Cambridge, MA 02138 (US).

(72) Inventors: **LIU, David, R.**; 3 Whitman Circle, Lexington, MA 02420 (US). **KOMOR, Alexis Christine**; 1000 Cordova Street, #109, Pasadena, CA 91106 (US). **REES, Holly A.**; 35 Oxford Street, Cambridge, MA 02138 (US). **KIM, Yongjoo**; 5 Cowperthwaite Street, #323, Cambridge, MA 02138 (US).

(74) Agent: **BAKER, C. Hunter**; Wolf, Greenfield & Sacks, P.C., 600 Atlantic Avenue, Boston, MA 02210-2206 (US).

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(57) Abstract: Some aspects of this disclosure provide strategies, systems, reagents, methods, and kits that are useful for the targeted editing of nucleic acids, including editing a single site within the genome of a cell or subject, e.g., within the human genome. In some embodiments, fusion proteins of Cas9 and nucleic acid editing proteins or protein domains, e.g., deaminase domains, are provided. In some embodiments, methods for targeted nucleic acid editing are provided. In some embodiments, reagents and kits for the generation of targeted nucleic acid editing proteins, e.g., fusion proteins of Cas9 and nucleic acid editing proteins or domains, are provided.



NUCLEOBASE EDITORS AND USES THEREOF

GOVERNMENT SUPPORT

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RELATED APPLICATIONS

[0002] This application claims priority under 35 U.S.C. § 119(e) to U.S. provisional patent applications, U.S.S.N. 62/245,828 filed October 23, 2015, U.S.S.N. 62/279,346 filed January 15, 2016, U.S.S.N. 62/311,763 filed March 22, 2016, U.S.S.N. 62/322,178 filed April 13, 2016, U.S.S.N. 62/357,352 filed June 30, 2016, U.S.S.N. 62/370,700 filed August 3, 2016, U.S.S.N. 62/398,490 filed September 22, 2016, U.S.S.N. 62/408,686 filed October 14, 2016, and U.S.S.N. 62/357,332 filed June 30, 2016; each of which is incorporated herein by reference.

BACKGROUND OF THE INVENTION

[0003] Targeted editing of nucleic acid sequences, for example, the targeted cleavage or the targeted introduction of a specific modification into genomic DNA, is a highly promising approach for the study of gene function and also has the potential to provide new therapies for human genetic diseases.¹ An ideal nucleic acid editing technology possesses three characteristics: (1) high efficiency of installing the desired modification; (2) minimal off-target activity; and (3) the ability to be programmed to edit precisely any site in a given nucleic acid, *e.g.*, any site within the human genome.² Current genome engineering tools, including engineered zinc finger nucleases (ZFNs),³ transcription activator like effector nucleases (TALENs),⁴ and most recently, the RNA-guided DNA endonuclease Cas9,⁵ effect sequence-specific DNA cleavage in a genome. This programmable cleavage can result in mutation of the DNA at the cleavage site via non-homologous end joining (NHEJ) or replacement of the DNA surrounding the cleavage site via homology-directed repair (HDR).^{6,7}

[0004] One drawback to the current technologies is that both NHEJ and HDR are stochastic processes that typically result in modest gene editing efficiencies as well as unwanted gene alterations that can compete with the desired alteration.⁸ Since many genetic diseases in principle can be treated by effecting a specific nucleotide change at a specific location in the genome (for example, a C to T change in a specific codon of a gene associated with a disease),⁹ the development of a programmable way to achieve such precision gene editing would represent both a powerful new research tool, as well as a potential new approach to gene editing-based human therapeutics.

SUMMARY OF THE INVENTION

[0005] The clustered regularly interspaced short palindromic repeat (CRISPR) system is a recently discovered prokaryotic adaptive immune system¹⁰ that has been modified to enable robust and general genome engineering in a variety of organisms and cell lines.¹¹ CRISPR-Cas (CRISPR associated) systems are protein-RNA complexes that use an RNA molecule (sgRNA) as a guide to localize the complex to a target DNA sequence *via* base-pairing.¹² In the natural systems, a Cas protein then acts as an endonuclease to cleave the targeted DNA sequence.¹³ The target DNA sequence must be both complementary to the sgRNA, and also contain a “protospacer-adjacent motif” (PAM) at the 3'-end of the complementary region in order for the system to function.¹⁴

[0006] Among the known Cas proteins, *S. pyogenes* Cas9 has been mostly widely used as a tool for genome engineering.¹⁵ This Cas9 protein is a large, multi-domain protein containing two distinct nuclease domains. Point mutations can be introduced into Cas9 to abolish nuclease activity, resulting in a dead Cas9 (dCas9) that still retains its ability to bind DNA in a sgRNA-programmed manner.¹⁶ In principle, when fused to another protein or domain, dCas9 can target that protein to virtually any DNA sequence simply by co-expression with an appropriate sgRNA.

[0007] The potential of the dCas9 complex for genome engineering purposes is immense. Its unique ability to bring proteins to specific sites in a genome programmed by the sgRNA in theory can be developed into a variety of site-specific genome engineering tools beyond nucleases, including transcriptional activators, transcriptional repressors, histone-modifying proteins, integrases, and recombinases.¹¹ Some of these potential applications have recently been implemented through dCas9 fusions with transcriptional activators to afford RNA-guided transcriptional activators,^{17,18} transcriptional repressors,^{16,19,20} and chromatin

modification enzymes.²¹ Simple co-expression of these fusions with a variety of sgRNAs results in specific expression of the target genes. These seminal studies have paved the way for the design and construction of readily programmable sequence-specific effectors for the precise manipulation of genomes.

[0008] Significantly, 80-90% of protein mutations responsible for human disease arise from the substitution, deletion, or insertion of only a single nucleotide.⁶ Most current strategies for single-base gene correction include engineered nucleases (which rely on the creation of double-strand breaks, DSBs, followed by stochastic, inefficient homology-directed repair, HDR), and DNA-RNA chimeric oligonucleotides.²² The latter strategy involves the design of a RNA/DNA sequence to base pair with a specific sequence in genomic DNA except at the nucleotide to be edited. The resulting mismatch is recognized by the cell's endogenous repair system and fixed, leading to a change in the sequence of either the chimera or the genome. Both of these strategies suffer from low gene editing efficiencies and unwanted gene alterations, as they are subject to both the stochasticity of HDR and the competition between HDR and non-homologous end-joining, NHEJ.²³⁻²⁵ HDR efficiencies vary according to the location of the target gene within the genome,²⁶ the state of the cell cycle,²⁷ and the type of cell/tissue.²⁸ The development of a direct, programmable way to install a specific type of base modification at a precise location in genomic DNA with enzyme-like efficiency and no stochasticity therefore represents a powerful new approach to gene editing-based research tools and human therapeutics.

[0009] Some aspects of the disclosure are based on the recognition that certain configurations of a dCas9 domain, and a cytidine deaminase domain fused by a linker are useful for efficiently deaminating target cytidine residues. Other aspects of this disclosure relate to the recognition that a nucleobase editing fusion protein with a cytidine deaminase domain fused to the N-terminus of a nuclease inactive Cas9 (dCas9) via a linker was capable of efficiently deaminating target nucleic acids in a double stranded DNA target molecule. See for example, Examples 3 and 4 below, which demonstrate that the fusion proteins, which are also referred to herein as base editors, generate less indels and more efficiently deaminate target nucleic acids than other base editors, such as base editors without a UGI domain. In some embodiments, the fusion protein comprises a nuclease-inactive Cas9 (dCas9) domain and an apolipoprotein B mRNA-editing complex 1 (APOBEC1) deaminase domain, where the deaminase domain is fused to the N-terminus of the dCas9 domain via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7). In some embodiments, the nuclease-inactive Cas9 (dCas9) domain of comprises the amino acid

sequence set forth in SEQ ID NO: 263. In some embodiments, the deaminase is rat APOBEC1 (SEQ ID NO: 284). In some embodiments, the deaminase is human APOBEC1 (SEQ ID NO: 282). In some embodiments, the deaminase is pmCDA1 (SEQ ID NO: 5738). In some embodiments, the deaminase is human APOBEC3G (SEQ ID NO: 275). In some embodiments, the deaminase is a human APOBEC3G variant of any one of (SEQ ID NOs: 5739-5741).

[0010] Some aspects of the disclosure are based on the recognition that certain configurations of a dCas9 domain, and a cytidine deaminase domain fused by a linker are useful for efficiently deaminating target cytidine residues. Other aspects of this disclosure relate to the recognition that a nucleobase editing fusion protein with an apolipoprotein B mRNA-editing complex 1 (APOBEC1) deaminase domain fused to the N-terminus of a nuclease inactive Cas9 (dCas9) via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7) was capable of efficiently deaminating target nucleic acids in a double stranded DNA target molecule. In some embodiments, the fusion protein comprises a nuclease-inactive Cas9 (dCas9) domain and an apolipoprotein B mRNA-editing complex 1 (APOBEC1) deaminase domain, where the deaminase domain is fused to the N-terminus of the dCas9 domain via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).

[0011] In some embodiments, the fusion protein comprises the amino acid residues 11-1629 of the amino acid sequence set forth in SEQ ID NO: 591. In some embodiments, the fusion protein comprises the amino acid sequence set forth in SEQ ID NO: 591. In some embodiments, the fusion protein comprises the amino acid sequence of any one of SEQ ID NOs: 5737, 5743, 5745, and 5746.

[0012] Some aspects of this disclosure provide strategies, systems, reagents, methods, and kits that are useful for the targeted editing of nucleic acids, including editing a single site within a subject's genome, *e.g.*, a human's genome. In some embodiments, fusion proteins of Cas9 (*e.g.*, dCas9, nuclease active Cas9, or Cas9 nickase) and deaminases or deaminase domains, are provided. In some embodiments, methods for targeted nucleic acid editing are provided. In some embodiments, reagents and kits for the generation of targeted nucleic acid editing proteins, *e.g.*, fusion proteins of Cas9 and deaminases or deaminase domains, are provided.

[0013] Some aspects of this disclosure provide fusion proteins comprising a Cas9 protein as provided herein that is fused to a second protein (*e.g.*, an enzymatic domain such as a cytidine deaminase domain), thus forming a fusion protein. In some embodiments, the

second protein comprises an enzymatic domain, or a binding domain. In some embodiments, the enzymatic domain is a nuclease, a nickase, a recombinase, a deaminase, a methyltransferase, a methylase, an acetylase, an acetyltransferase, a transcriptional activator, or a transcriptional repressor domain. In some embodiments, the enzymatic domain is a nucleic acid editing domain. In some embodiments, the nucleic acid editing domain is a deaminase domain. In some embodiments, the deaminase is a cytosine deaminase or a cytidine deaminase. In some embodiments, the deaminase is an apolipoprotein B mRNA-editing complex (APOBEC) family deaminase. In some embodiments, the deaminase is an APOBEC1 deaminase. In some embodiments, the deaminase is an APOBEC2 deaminase. In some embodiments, the deaminase is an APOBEC3 deaminase. In some embodiments, the deaminase is an APOBEC3A deaminase. In some embodiments, the deaminase is an APOBEC3B deaminase. In some embodiments, the deaminase is an APOBEC3C deaminase. In some embodiments, the deaminase is an APOBEC3D deaminase. In some embodiments, the deaminase is an APOBEC3E deaminase. In some embodiments, the deaminase is an APOBEC3F deaminase. In some embodiments, the deaminase is an APOBEC3G deaminase. In some embodiments, the deaminase is an APOBEC3H deaminase. In some embodiments, the deaminase is an APOBEC4 deaminase. In some embodiments, the deaminase is an activation-induced deaminase (AID). It should be appreciated that the deaminase may be from any suitable organism (*e.g.*, a human or a rat). In some embodiments, the deaminase is from a human, chimpanzee, gorilla, monkey, cow, dog, rat, or mouse. In some embodiments, the deaminase is rat APOBEC1 (SEQ ID NO: 284). In some embodiments, the deaminase is human APOBEC1 (SEQ ID NO: 282). In some embodiments, the deaminase is pmCDA1.

[0014] Some aspects of this disclosure provide fusion proteins comprising: (i) a nuclease-inactive Cas9 (dCas9) domain comprising the amino acid sequence of SEQ ID NO: 263; and (ii) an apolipoprotein B mRNA-editing complex 1 (APOBEC1) deaminase domain, wherein the deaminase domain is fused to the N-terminus of the dCas9 domain via a linker comprising the amino acid sequence of SGSETPGTSESATPES (SEQ ID NO: 7). In some embodiments, the deaminase is rat APOBEC1 (SEQ ID NO: 284). In some embodiments, the deaminase is human APOBEC1 (SEQ ID NO: 282). In some embodiments, the fusion protein comprises the amino acid sequence of SEQ ID NO: 591. In some embodiments, the fusion protein comprises the amino acid sequence of SEQ ID NO: 5737. In some embodiments, the deaminase is pmCDA1 (SEQ ID NO: 5738). In some embodiments, the deaminase is human APOBEC3G (SEQ ID NO: 275). In some embodiments, the deaminase is a human APOBEC3G variant of any one of SEQ ID NOs: 5739-5741.

[0015] Some aspects of this disclosure provide fusion proteins comprising: (i) a Cas9 nickase domain and (ii) an apolipoprotein B mRNA-editing complex 1 (APOBEC1) deaminase domain, wherein the deaminase domain is fused to the N-terminus of the Cas9 nickase domain. In some embodiments, the Cas9 nickase domain comprises a D10X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for D. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises a D10A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises a histidine at amino acid position 840 of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding amino acid position in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises the amino acid sequence as set forth in SEQ ID NO: 267. In some embodiments, the deaminase is rat APOBEC1 (SEQ ID NO: 284). In some embodiments, the deaminase is human APOBEC1 (SEQ ID NO: 282). In some embodiments, the deaminase is pmCDA1.

[0016] Some aspects of this disclosure provide fusion proteins comprising: (i) a Cas9 nickase domain and (ii) an apolipoprotein B mRNA-editing complex 1 (APOBEC1) deaminase domain, wherein the deaminase domain is fused to the N-terminus of the Cas9 nickase domain. In some embodiments, the Cas9 nickase domain comprises a D10X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for D. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises a D10A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises a histidine at amino acid position 840 of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding amino acid position in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises the amino acid sequence as set forth in SEQ ID NO: 267. In some embodiments, the deaminase is rat APOBEC1 (SEQ ID NO: 284). In some embodiments, the deaminase is human APOBEC1 (SEQ ID NO: 282). In some embodiments, the deaminase is pmCDA1.

[0017] Other aspects of this disclosure relate to the recognition that fusion proteins comprising a deaminase domain, a dCas9 domain and a uracil glycosylase inhibitor (UGI) domain demonstrate improved efficiency for deaminating target nucleotides in a nucleic acid molecule. Without wishing to be bound by any particular theory, cellular DNA-repair response to the presence of U:G heteroduplex DNA may be responsible for a decrease in nucleobase editing efficiency in cells. Uracil DNA glycosylase (UDG) catalyzes removal of U from DNA in cells, which may initiate base excision repair, with reversion of the U:G pair to a C:G pair as the most common outcome. As demonstrated herein, Uracil DNA Glycosylase Inhibitor (UGI) may inhibit human UDG activity. Without wishing to be bound by any particular theory, base excision repair may be inhibited by molecules that bind the single strand, block the edited base, inhibit UGI, inhibit base excision repair, protect the edited base, and/or promote “fixing” of the non-edited strand, *etc.* Thus, this disclosure contemplates fusion proteins comprising a dCas9-cytidine deaminase domain that is fused to a UGI domain.

[0018] In some embodiments, the fusion protein comprises a nuclease-inactive Cas9 (dCas9) domain; a nucleic acid editing domain; and a uracil glycosylase inhibitor (UGI) domain. In some embodiments, the dCas9 domain comprises a D10X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for D. In some embodiments, the amino acid sequence of the dCas9 domain comprises a D10A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the amino acid sequence of the dCas9 domain comprises an H840X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for H. In some embodiments, the amino acid sequence of the dCas9 domain comprises an H840A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the dCas9 domain comprises the amino acid sequence as set forth in SEQ ID NO: 263.

[0019] Further aspects of this disclosure relate to the recognition that fusion proteins using a Cas9 nickase as the Cas9 domain demonstrate improved efficiency for editing nucleic acids. For example, aspects of this disclosure relate to the recognition that fusion proteins comprising a Cas9 nickase, a deaminase domain and a UGI domain demonstrate improved

efficiency for editing nucleic acids. For example, the improved efficiency for editing nucleotides is described below in the Examples section.

[0020] Some aspects of the disclosure are based on the recognition that any of the base editors provided herein are capable of modifying a specific nucleotide base without generating a significant proportion of indels. An “indel”, as used herein, refers to the insertion or deletion of a nucleotide base within a nucleic acid. Such insertions or deletions can lead to frame shift mutations within a coding region of a gene. In some embodiments, it is desirable to generate base editors that efficiently modify (*e.g.* mutate or deaminate) a specific nucleotide within a nucleic acid, without generating a large number of insertions or deletions (*i.e.*, indels) in the nucleic acid. In certain embodiments, any of the base editors provided herein are capable of generating a greater proportion of intended modifications (*e.g.*, point mutations or deaminations) versus indels.

[0021] Some aspects of the disclosure are based on the recognition that any of the base editors provided herein are capable of efficiently generating an intended mutation, such as a point mutation, in a nucleic acid (*e.g.* a nucleic acid within a genome of a subject) without generating a significant number of unintended mutations, such as unintended point mutations.

[0022] In some embodiments, a fusion protein comprises a Cas9 nickase domain, a nucleic acid editing domain; and a uracil glycosylase inhibitor (UGI) domain. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises a D10X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOS: 11-260, wherein X is any amino acid except for D. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises a D10A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOS: 11-260. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises a histidine at amino acid position 840 of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding amino acid position in any of the amino acid sequences provided in SEQ ID NOS: 11-260. In some embodiments, the amino acid sequence of the Cas9 nickase domain comprises the amino acid sequence as set forth in SEQ ID NO: 267.

[0023] In some embodiments, the deaminase domain of the fusion protein is fused to the N-terminus of the dCas9 domain or the Cas9 nickase. In some embodiments, the UGI domain is fused to the C-terminus of the dCas9 domain or the Cas9 nickase. In some embodiments, the dCas9 domain or the Cas9 nickase and the nucleic acid editing domain are fused via a linker.

In some embodiments, the dCas9 domain or the Cas9 nickase and the UGI domain are fused via a linker.

[0024] In certain embodiments, linkers may be used to link any of the peptides or peptide domains of the invention. The linker may be as simple as a covalent bond, or it may be a polymeric linker many atoms in length. In certain embodiments, the linker is a polypeptide or based on amino acids. In other embodiments, the linker is not peptide-like. In certain embodiments, the linker is a covalent bond (*e.g.*, a carbon-carbon bond, disulfide bond, carbon-heteroatom bond, *etc.*). In certain embodiments, the linker is a carbon-nitrogen bond of an amide linkage. In certain embodiments, the linker is a cyclic or acyclic, substituted or unsubstituted, branched or unbranched aliphatic or heteroaliphatic linker. In certain embodiments, the linker is polymeric (*e.g.*, polyethylene, polyethylene glycol, polyamide, polyester, *etc.*). In certain embodiments, the linker comprises a monomer, dimer, or polymer of aminoalkanoic acid. In certain embodiments, the linker comprises an aminoalkanoic acid (*e.g.*, glycine, ethanoic acid, alanine, beta-alanine, 3-aminopropanoic acid, 4-aminobutanoic acid, 5-pentanoic acid, *etc.*). In certain embodiments, the linker comprises a monomer, dimer, or polymer of aminohexanoic acid (Ahx). In certain embodiments, the linker is based on a carbocyclic moiety (*e.g.*, cyclopentane, cyclohexane). In other embodiments, the linker comprises a polyethylene glycol moiety (PEG). In other embodiments, the linker comprises amino acids. In certain embodiments, the linker comprises a peptide. In certain embodiments, the linker comprises an aryl or heteroaryl moiety. In certain embodiments, the linker is based on a phenyl ring. The linker may include functionalized moieties to facilitate attachment of a nucleophile (*e.g.*, thiol, amino) from the peptide to the linker. Any electrophile may be used as part of the linker. Exemplary electrophiles include, but are not limited to, activated esters, activated amides, Michael acceptors, alkyl halides, aryl halides, acyl halides, and isothiocyanates.

[0025] In some embodiments, the linker comprises the amino acid sequence (GGGGS)_n (SEQ ID NO: 5), (G)_n, (EAAAK)_n (SEQ ID NO: 6), (GGS)_n, (SGGS)_n (SEQ ID NO: 4288), SGSETPGTSESATPES (SEQ ID NO: 7), (XP)_n, or any combination thereof, wherein n is independently an integer between 1 and 30, and wherein X is any amino acid. In some embodiments, the linker comprises the amino acid sequence (GGS)_n, wherein n is 1, 3, or 7. In some embodiments, the linker comprises the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).

[0026] In some embodiments, the fusion protein comprises the structure [nucleic acid editing domain]-[optional linker sequence]-[dCas9 or Cas9 nickase]-[optional linker

sequence]-[UGI]. In some embodiments, the fusion protein comprises the structure [nucleic acid editing domain]-[optional linker sequence]-[UGI]-[optional linker sequence]-[dCas9 or Cas9 nickase]; [UGI]-[optional linker sequence]-[nucleic acid editing domain]-[optional linker sequence]-[dCas9 or Cas9 nickase]; [UGI]-[optional linker sequence]-[dCas9 or Cas9 nickase]-[optional linker sequence]-[nucleic acid editing domain]; [dCas9 or Cas9 nickase]-[optional linker sequence]-[UGI]-[optional linker sequence]-[nucleic acid editing domain]; or [dCas9 or Cas9 nickase]-[optional linker sequence]-[nucleic acid editing domain]-[optional linker sequence]-[UGI].

[0027] In some embodiments, the nucleic acid editing domain comprises a deaminase. In some embodiments, the nucleic acid editing domain comprises a deaminase. In some embodiments, the deaminase is a cytidine deaminase. In some embodiments, the deaminase is an apolipoprotein B mRNA-editing complex (APOBEC) family deaminase. In some embodiments, the deaminase is an APOBEC1 deaminase, an APOBEC2 deaminase, an APOBEC3A deaminase, an APOBEC3B deaminase, an APOBEC3C deaminase, an APOBEC3D deaminase, an APOBEC3F deaminase, an APOBEC3G deaminase, an APOBEC3H deaminase, or an APOBEC4 deaminase. In some embodiments, the deaminase is an activation-induced deaminase (AID). In some embodiments, the deaminase is a Lamprey CDA1 (pmCDA1) deaminase.

[0028] In some embodiments, the deaminase is from a human, chimpanzee, gorilla, monkey, cow, dog, rat, or mouse. In some embodiments, the deaminase is from a human. In some embodiments the deaminase is from a rat. In some embodiments, the deaminase is a rat APOBEC1 deaminase comprising the amino acid sequence set forth in (SEQ ID NO: 284). In some embodiments, the deaminase is a human APOBEC1 deaminase comprising the amino acid sequence set forth in (SEQ ID NO: 282). In some embodiments, the deaminase is pmCDA1 (SEQ ID NO: 5738). In some embodiments, the deaminase is human APOBEC3G (SEQ ID NO: 275). In some embodiments, the deaminase is a human APOBEC3G variant of any one of (SEQ ID NOs: 5739-5741). In some embodiments, the deaminase is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of the amino acid sequences set forth in SEQ ID NOs: 266-284 or 5725-5741.

[0029] In some embodiments, the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to SEQ ID NO: 600. In some

embodiments, the UGI domain comprises the amino acid sequence as set forth in SEQ ID NO: 600.

[0030] Some aspects of this disclosure provide complexes comprising a Cas9 protein or a Cas9 fusion protein as provided herein, and a guide RNA bound to the Cas9 protein or the Cas9 fusion protein.

[0031] Some aspects of this disclosure provide methods of using the Cas9 proteins, fusion proteins, or complexes provided herein. For example, some aspects of this disclosure provide methods comprising contacting a DNA molecule (a) with a Cas9 protein or a fusion protein as provided herein and with a guide RNA, wherein the guide RNA is about 15-100 nucleotides long and comprises a sequence of at least 10 contiguous nucleotides that is complementary to a target sequence; or (b) with a Cas9 protein, a Cas9 fusion protein, or a Cas9 protein or fusion protein complex with a gRNA as provided herein.

[0032] Some aspects of this disclosure provide kits comprising a nucleic acid construct, comprising (a) a nucleotide sequence encoding a Cas9 protein or a Cas9 fusion protein as provided herein; and (b) a heterologous promoter that drives expression of the sequence of (a). In some embodiments, the kit further comprises an expression construct encoding a guide RNA backbone, wherein the construct comprises a cloning site positioned to allow the cloning of a nucleic acid sequence identical or complementary to a target sequence into the guide RNA backbone.

[0033] Some aspects of this disclosure provide polynucleotides encoding a Cas9 protein or a fusion protein as provided herein. Some aspects of this disclosure provide vectors comprising such polynucleotides. In some embodiments, the vector comprises a heterologous promoter driving expression of polynucleotide.

[0034] Some aspects of this disclosure provide cells comprising a Cas9 protein, a fusion protein, a nucleic acid molecule, and/or a vector as provided herein.

[0035] The description of exemplary embodiments of the reporter systems above is provided for illustration purposes only and not meant to be limiting. Additional reporter systems, *e.g.*, variations of the exemplary systems described in detail above, are also embraced by this disclosure.

[0036] The summary above is meant to illustrate, in a non-limiting manner, some of the embodiments, advantages, features, and uses of the technology disclosed herein. Other embodiments, advantages, features, and uses of the technology disclosed herein will be apparent from the Detailed Description, the Drawings, the Examples, and the Claims.

BRIEF DESCRIPTION OF THE DRAWINGS

[0037] **Figure 1** shows the deaminase activity of deaminases on single stranded DNA substrates. Single stranded DNA substrates using randomized PAM sequences (NNN PAM) were used as negative controls. Canonical PAM sequences used (NGG PAM)

[0038] **Figure 2** shows activity of Cas9:deaminase fusion proteins on single stranded DNA substrates.

[0039] **Figure 3** illustrates double stranded DNA substrate binding by Cas9:deaminase:sgRNA complexes.

[0040] **Figure 4** illustrates a double stranded DNA deamination assay.

[0041] **Figure 5** demonstrates that Cas9 fusions can target positions 3-11 of double-stranded DNA target sequences (numbered according to the schematic in **Figure 5**). Upper Gel: 1 μ M rAPOBEC1-GGS-dCas9, 125 nM dsDNA, 1 equivalent sgRNA. Mid Gel: 1 μ M rAPOBEC1-(GGS)₃(SEQ ID NO: 596)-dCas9, 125 nM dsDNA, 1 equivalent sgRNA. Lower Gel: 1.85 μ M rAPOBEC1-XTEN-dCas9, 125 nM dsDNA, 1 equivalent sgRNA.

[0042] **Figure 6** demonstrates that the correct guide RNA, *e.g.*, the correct sgRNA, is required for deaminase activity.

[0043] **Figure 7** illustrates the mechanism of target DNA binding of *in vivo* target sequences by deaminase-dCas9:sgRNA complexes.

[0044] **Figure 8** shows successful deamination of exemplary disease-associated target sequences.

[0045] **Figure 9** shows *in vitro* C→T editing efficiencies using His6-rAPOBEC1-XTEN-dCas9.

[0046] **Figure 10** shows C→T editing efficiencies in HEK293T cells is greatly enhanced by fusion with UGI.

[0047] **Figures 11A to 11C** show NBE1 mediates specific, guide RNA-programmed C to U conversion *in vitro*. **Figure 11A**: Nucleobase editing strategy. DNA with a target C (red) at a locus specified by a guide RNA (green) is bound by dCas9 (blue), which mediates the local denaturation of the DNA substrate. Cytidine deamination by a tethered APOBEC1 enzyme (orange) converts the target C to U. The resulting G:U heteroduplex can be permanently converted to an A:T base pair following DNA replication or repair. If the U is in the template DNA strand, it will also result in an RNA transcript containing a G to A mutation following transcription. **Figure 11B**: Deamination assay showing an activity window of approximately five nucleotides. Following incubation of NBE1-sgRNA complexes with dsDNA substrates

at 37 °C for 2 h, the 5' fluorophore-labeled DNA was isolated and incubated with USER enzyme (uracil DNA glycosylase and endonuclease VIII) at 37 °C for 1 h to induce DNA cleavage at the site of any uracils. The resulting DNA was resolved on a denaturing polyacrylamide gel, and any fluorophore-linked strands were visualized. Each lane is labeled according to the position of the target C within the protospacer, or with “–” if no target C is present, counting the base distal from the PAM as position 1. Figure 11C: Deaminase assay showing the sequence specificity and sgRNA-dependence of NBE1. The DNA substrate with a target C at position 7 was incubated with NBE1 as in Figure 11B with either the correct sgRNA, a mismatched sgRNA, or no sgRNA. No C to U editing is observed with the mismatched sgRNA or with no sgRNA. The positive control sample contains a DNA sequence with a U synthetically incorporated at position 7.

[0048] Figures 12A to 12B show effects of sequence context and target C position on nucleobase editing efficiency *in vitro*. Figure 12A: Effect of changing the sequence surrounding the target C on editing efficiency *in vitro*. The deamination yield of 80% of targeted strands (40% of total sequencing reads from both strands) for C₇ in the protospacer sequence 5'-TTATTTTCGTGGATTATTTA-3'(SEQ ID NO: 264) was defined as 1.0, and the relative deamination efficiencies of substrates containing all possible single-base mutations at positions 1-6 and 8-13 are shown. Values and error bars reflect the mean and standard deviation of two or more independent biological replicates performed on different days. Figure 12B: Positional effect of each NC motif on editing efficiency *in vitro*. Each NC target motif was varied from positions 1 to 8 within the protospacer as indicated in the sequences shown on the right (the PAM shown in red, the protospacer plus one base 5' to the protospacer are also shown). The percentage of total sequence reads containing T at each of the numbered target C positions following incubation with NBE1 is shown in the graph. Note that the maximum possible deamination yield *in vitro* is 50% of total sequencing reads (100% of targeted strands). Values and error bars reflect the mean and standard deviation of two or three independent biological replicates performed on different days. Figure 12B depicts SEQ ID NOs: 285 through 292 from top to bottom, respectively.

[0049] Figures 13A to 13C show nucleobase editing in human cells. Figure 13A: Protospacer (black) and PAM (red) sequences of the six mammalian cell genomic loci targeted by nucleobase editors. Target Cs are indicated with subscripted numbers corresponding to their positions within the protospacer. Figure 13A depicts SEQ ID NOs: 293 through 298 from top to bottom, respectively. Figure 13B: HEK293T cells were transfected with plasmids expressing NBE1, NBE2, or NBE3 and an appropriate sgRNA. Three days

after transfection, genomic DNA was extracted and analyzed by high-throughput DNA sequencing at the six loci. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with Ts at the target positions indicated, are shown for NBE1, NBE2, and NBE3 at all six genomic loci, and for wt Cas9 with a donor HDR template at three of the six sites (EMX1, HEK293 site 3, and HEK293 site 4). Values and error bars reflect the mean and standard deviation of three independent biological replicates performed on different days. Figure 13C: Frequency of indel formation, calculated as described in the Methods, is shown following treatment of HEK293T cells with NBE2 and NBE3 for all six genomic loci, or with wt Cas9 and a single-stranded DNA template for HDR at three of the six sites (EMX1, HEK293 site 3, and HEK293 site 4). Values reflect the mean of at least three independent biological replicates performed on different days.

[0050] Figures 14A to 14C show NBE2- and NBE3-mediated correction of three disease-relevant mutations in mammalian cells. For each site, the sequence of the protospacer is indicated to the right of the name of the mutation, with the PAM highlighted in green and the base responsible for the mutation indicated in bold with a subscripted number corresponding to its position within the protospacer. The amino acid sequence above each disease-associated allele is shown, together with the corrected amino acid sequence following nucleobase editing in red. Underneath each sequence are the percentages of total sequencing reads with the corresponding base. Cells were nucleofected with plasmids encoding NBE2 or NBE3 and an appropriate sgRNA. Two days after nucleofection, genomic DNA was extracted and analyzed by HTS to assess pathogenic mutation correction. Figure 14A: The Alzheimer's disease-associated *APOE4* allele is converted to *APOE3* in mouse astrocytes by NBE3 in 11% of total reads (44% of nucleofected astrocytes). Two nearby Cs are also converted to Ts, but with no change to the predicted sequence of the resulting protein (SEQ ID NO: 299). Figure 14B The cancer-associated p53 N239D mutation is corrected by NBE2 in 11% of treated human lymphoma cells (12% of nucleofected cells) that are heterozygous for the mutation (SEQ ID NO: 300). Figure 14C The p53 Y163C mutation is corrected by NBE3 in 7.6% of nucleofected human breast cancer cells (SEQ ID NO: 301).

[0051] Figures 15A to 15D show effects of deaminase-dCas9 linker length and composition on nucleobase editing. Gel-based deaminase assay showing the deamination window of nucleobase editors with deaminase-Cas9 linkers of GGS (Figure 15A), (GGS)₃ (SEQ ID NO: 596) (Figure 15B), XTEN (Figure 15C), or (GGS)₇ (SEQ ID NO: 597) (Figure 15D). Following incubation of 1.85 μM editor-sgRNA complexes with 125 nM dsDNA substrates at 37 °C for 2 h, the dye-conjugated DNA was isolated and incubated with USER

enzyme (uracil DNA glycosylase and endonuclease VIII) at 37 °C for an additional hour to cleave the DNA backbone at the site of any uracils. The resulting DNA was resolved on a denaturing polyacrylamide gel, and the dye-conjugated strand was imaged. Each lane is numbered according to the position of the target C within the protospacer, or with – if no target C is present. 8U is a positive control sequence with a U synthetically incorporated at position 8.

[0052] **Figures 16A to 16B** show NBE1 is capable of correcting disease-relevant mutations *in vitro*. Figure 16A: Protospacer and PAM sequences (red) of seven disease-relevant mutations. The disease-associated target C in each case is indicated with a subscripted number reflecting its position within the protospacer. For all mutations except both *APOE4* SNPs, the target C resides in the template (non-coding) strand. Figure 16A depicts SEQ ID NOs: 302 through 308 from top to bottom, respectively. Figure 16B: Deaminase assay showing each dsDNA oligonucleotide before (–) and after (+) incubation with NBE1, DNA isolation, and incubation with USER enzymes to cleave DNA at positions containing U. Positive control lanes from incubation of synthetic oligonucleotides containing U at various positions within the protospacer with USER enzymes are shown with the corresponding number indicating the position of the U.

[0053] **Figure 17** shows processivity of NBE1. The protospacer and PAM (red) of a 60-mer DNA oligonucleotide containing eight consecutive Cs is shown at the top. The oligonucleotide (125 nM) was incubated with NBE1 (2 μM) for 2 h at 37 °C. The DNA was isolated and analyzed by high-throughput sequencing. Shown are the percent of total reads for the most frequent nine sequences observed. The vast majority of edited strands (>93%) have more than one C converted to T. This figure depicts SEQ ID NO: 309.

[0054] **Figures 18A to 18H** show the effect of fusing UGI to NBE1 to generate NBE2. Figure 18A: Protospacer and PAM (red) sequences of the six mammalian cell genomic loci targeted with nucleobase editors. Editable Cs are indicated with labels corresponding to their positions within the protospacer. Figure 18A depicts SEQ ID NOs: 293 through 298 from top to bottom, respectively. Figures 18B to 18G: HEK293T cells were transfected with plasmids expressing NBE1, NBE2, or NBE1 and UGI, and an appropriate sgRNA. Three days after transfection, genomic DNA was extracted and analyzed by high-throughput DNA sequencing at the six loci. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with Ts at the target positions indicated, are shown for NBE1, NBE1 and UGI, and NBE2 at all six genomic loci. Figure 18H: C to T mutation rates at 510 Cs surrounding the protospacers of interest for NBE1, NBE1 plus UGI on a separate plasmid,

NBE2, and untreated cells are shown. The data show the results of 3,000,000 DNA sequencing reads from 1.5×10^6 cells. Values reflect the mean of at least two biological experiments conducted on different days.

[0055] **Figure 19** shows nucleobase editing efficiencies of NBE2 in U2OS and HEK293T cells. Cellular C to T conversion percentages by NBE2 are shown for each of the six targeted genomic loci in HEK293T cells and U2OS cells. HEK293T cells were transfected using lipofectamine 2000, and U2OS cells were nucleofected. U2OS nucleofection efficiency was 74%. Three days after plasmid delivery, genomic DNA was extracted and analyzed for nucleobase editing at the six genomic loci by HTS. Values and error bars reflect the mean and standard deviation of at least two biological experiments done on different days.

[0056] **Figure 20** shows nucleobase editing persists over multiple cell divisions. Cellular C to T conversion percentages by NBE2 are displayed at two genomic loci in HEK293T cells before and after passaging the cells. HEK293T cells were transfected using Lipofectamine 2000. Three days post transfection, the cells were harvested and split in half. One half was subjected to HTS analysis, and the other half was allowed to propagate for approximately five cell divisions, then harvested and subjected to HTS analysis.

[0057] **Figure 21** shows genetic variants from ClinVar that can be corrected in principle by nucleobase editing. The NCBI ClinVar database of human genetic variations and their corresponding phenotypes⁶⁸ was searched for genetic diseases that can be corrected by current nucleobase editing technologies. The results were filtered by imposing the successive restrictions listed on the left. The x-axis shows the number of occurrences satisfying that restriction and all above restrictions on a logarithmic scale.

[0058] **Figure 22** shows *in vitro* identification of editable Cs in six genomic loci. Synthetic 80-mers with sequences matching six different genomic sites were incubated with NBE1 then analyzed for nucleobase editing *via* HTS. For each site, the sequence of the protospacer is indicated to the right of the name of the site, with the PAM highlighted in red. Underneath each sequence are the percentages of total DNA sequencing reads with the corresponding base. A target C was considered as “editable” if the *in vitro* conversion efficiency is >10%. Note that maximum yields are 50% of total DNA sequencing reads since the non-targeted strand is not a substrate for nucleobase editing. This figure depicts SEQ ID NOs: 293 through 298 from top to bottom, respectively.

[0059] **Figure 23** shows activities of NBE1, NBE2, and NBE3 at EMX1 off-targets. HEK293T cells were transfected with plasmids expressing NBE1, NBE2, or NBE3 and a sgRNA matching the EMX1 sequence using Lipofectamine 2000. Three days after

transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci, plus the top ten known Cas9 off-target loci for the EMX1 sgRNA, as previously determined using the GUIDE-seq method⁵⁵. EMX1 off-target 5 locus did not amplify and is not shown. Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for NBE1, NBE2, and NBE3. On the far right are displayed the total number of sequencing reads reported for each sequence. This figure depicts SEQ ID NOs: 293, and 310 through 318 from top to bottom, respectively.

[0060] **Figure 24** shows activities of NBE1, NBE2, and NBE3 at FANCF off-targets. HEK293T cells were transfected with plasmids expressing NBE1, NBE2, or NBE3 and a sgRNA matching the FANCF sequence using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci, plus all of the known Cas9 off-target loci for the FANCF sgRNA, as previously determined using the GUIDE-seq method⁵⁵. Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for NBE1, NBE2, and NBE3. On the far right are displayed the total number of sequencing reads reported for each sequence. This figure depicts SEQ ID NOs: 294 and 319 through 326 from top to bottom, respectively.

[0061] **Figure 25** shows activities of NBE1, NBE2, and NBE3 at HEK293 site 2 off-targets. HEK293T cells were transfected with plasmids expressing NBE1, NBE2, or NBE3 and a sgRNA matching the HEK293 site 2 sequence using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci, plus all of the known Cas9 off-target loci for the HEK293 site 2 sgRNA, as previously determined using the GUIDE-seq method⁵⁵. Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for NBE1, NBE2, and NBE3. On the far right are displayed the total number of sequencing reads reported for each sequence. This figure depicts SEQ ID NOs: 295, 327, and 328 from top to bottom, respectively.

[0062] **Figure 26** shows activities of NBE1, NBE2, and NBE3 at HEK293 site 3 off-targets. HEK293T cells were transfected with plasmids expressing NBE1, NBE2, or NBE3 and a sgRNA matching the HEK293 site 3 sequence using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci, plus all of the known Cas9 off-target loci for the HEK293 site 3 sgRNA, as previously determined using the GUIDE-seq method⁵⁵. Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for NBE1, NBE2, and NBE3. On the far right are displayed the total number of sequencing reads reported for each sequence. This figure depicts SEQ ID NOs: 296 and 659 through 663 from top to bottom, respectively.

[0063] **Figure 27** shows activities of NBE1, NBE2, and NBE3 at HEK293 site 4 off-targets. HEK293T cells were transfected with plasmids expressing NBE1, NBE2, or NBE3 and a sgRNA matching the HEK293 site 4 sequence using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci, plus the top ten known Cas9 off-target loci for the HEK293 site 4 sgRNA, as previously determined using the GUIDE-seq method⁵⁵. Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for NBE1, NBE2, and NBE3. On the far right are displayed the total number of sequencing reads reported for each sequence. This figure depicts SEQ ID NOs: 297 and 664 through 673 from top to bottom, respectively.

[0064] **Figure 28** shows non-target C mutation rates. Shown here are the C to T mutation rates at 2,500 distinct cytosines surrounding the six on-target and 34 off-target loci tested, representing a total of 14,700,000 sequence reads derived from approximately 1.8×10^6 cells.

[0065] **Figures 29A to 29C** show base editing in human cells. Figure 29A shows possible base editing outcomes in mammalian cells. Initial editing resulted in a U:G mismatch. Recognition and excision of the U by uracil DNA glycosylase (UDG) initiated base excision repair (BER), which lead to reversion to the C:G starting state. BER was impeded by BE2 and BE3, which inhibited UDG. The U:G mismatch was also processed by mismatch repair (MMR), which preferentially repaired the nicked strand of a mismatch. BE3 nicked the non-edited strand containing the G, favoring resolution of the U:G mismatch to the desired U:A or

T:A outcome. Figure 29B shows HEK293T cells treated as described in the Materials and Methods in the Examples below. The percentage of total DNA sequencing read with Ts at the target positions indicated show treatment with BE1, BE2, or BE3, or for treatment with wt Cas9 with a donor HDR template. Figure 29C shows frequency of indel formation following the treatment in Figure 29B. Values are listed in Figure 34. For Figures 29B and 29C, values and error bars reflect the mean and s.d. of three independent biological replicates performed on different days.

[0066] Figures 30A to 30B show BE3-mediated correction of two disease-relevant mutations in mammalian cells. The sequence of the protospacer is shown to the right of the mutation, with the PAM in blue and the target base in red with a subscripted number indicating its position within the protospacer. Underneath each sequence are the percentages of total sequencing reads with the corresponding base. Cells were treated as described in the Materials and Methods. Figure 30A shows the Alzheimer's disease-associated APOE4 allele converted to APOE3r in mouse astrocytes by BE3 in 74.9% of total reads. Two nearby Cs were also converted to Ts, but with no change to the predicted sequence of the resulting protein. Identical treatment of these cells with wt Cas9 and donor ssDNA results in only 0.3% correction, with 26.1% indel formation. Figure 30B shows the cancer associated p53 Y163C mutation corrected by BE3 in 7.6% of nucleofected human breast cancer cells with 0.7% indel formation. Identical treatment of these cells with wt Cas9 and donor ssDNA results in no mutation correction with 6.1% indel formation. This figure depicts SEQ ID NOs: 675 to 680 from top to bottom, respectively.

[0067] Figure 31 shows activities of BE1, BE2, and BE3 at HEK293 site 2 off-targets. HEK293T cells were transfected with plasmids expressing BE1, BE2, or BE3 and a sgRNA matching the HEK293 site 2 sequence using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci, plus all of the known Cas9 and dCas9 off-target loci for the HEK293 site 2 sgRNA, as previously determined by Joung and coworkers using the GUIDE-seq method (63), and Adli and coworkers using chromatin immunoprecipitation high-throughput sequencing (ChIP-seq) experiments (18). Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for BE1, BE2, and BE3. On the far right are displayed the total number of sequencing reads

reported, and the ChIP-seq signal intensity reported for each sequence. This figure depicts SEQ ID NOs: 681 to 688 from top to bottom, respectively.

[0068] **Figure 32** shows activities of BE1, BE2, and BE3 at HEK293 site 3 off-targets. HEK293T cells were transfected with plasmids expressing BE1, BE2, or BE3 and a sgRNA matching the HEK293 site 3 sequence using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci, plus all of the known Cas9 off-target loci and the top five known dCas9 off-target loci for the HEK293 site 3 sgRNA, as previously determined by Joung and coworkers using the GUIDE-seq method⁵⁴, and using chromatin immunoprecipitation high-throughput sequencing (ChIP-seq) experiments⁶¹. Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for BE1, BE2, and BE3. On the far right are displayed the total number of sequencing reads reported, and the ChIP-seq signal intensity reported for each sequence. This figure depicts SEQ ID NOs: 689 to 699 from top to bottom, respectively.

[0069] **Figure 33** shows activities of BE1, BE2, and BE3 at HEK293 site 4 off-targets. HEK293T cells were transfected with plasmids expressing BE1, BE2, or BE3 and a sgRNA matching the HEK293 site 4 sequence using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci, plus the top ten known Cas9 off-target loci and the top five known dCas9 off-target loci for the HEK293 site 4 sgRNA, as previously determined using the GUIDE-seq method⁵⁴, and using chromatin immunoprecipitation high-throughput sequencing (ChIP-seq) experiments⁶¹. Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for BE1, BE2, and BE3. On the far right are displayed the total number of sequencing reads reported, and the ChIP-seq signal intensity reported for each sequence. This figure depicts SEQ ID NOs: 700 to 712 from top to bottom, respectively.

[0070] **Figure 34** shows mutation rates of non-protospacer bases following BE3-mediated correction of the Alzheimer's disease-associated *APOE4* allele to *APOE3r* in mouse astrocytes. The DNA sequence of the 50 bases on either side of the protospacer from Figure 30A and Figure 34B is shown with each base's position relative to the protospacer. The side

of the protospacer distal to the PAM is designated with positive numbers, while the side that includes the PAM is designated with negative numbers, with the PAM shown in blue. Underneath each sequence are the percentages of total DNA sequencing reads with the corresponding base for untreated cells, for cells treated with BE3 and an sgRNA targeting the *APOE4* C158R mutation, or for cells treated with BE3 and an sgRNA targeting the *VEGFA* locus. Neither BE3-treated sample resulted in mutation rates above those of untreated controls. This figure depicts SEQ ID NOs: 713 to 716 from top to bottom, respectively.

[0071] Figure 35 shows mutation rates of non-protospacer bases following BE3-mediated correction of the cancer-associated p53 Y163C mutation in HCC1954 human cells. The DNA sequence of the 50 bases on either side of the protospacer from Figure 30B and Figure 39B is shown with each base's position relative to the protospacer. The side of the protospacer distal to the PAM is designated with positive numbers, while the side that includes the PAM is designated with negative numbers, with the PAM shown in blue. Underneath each sequence are the percentages of total sequencing reads with the corresponding base for untreated cells, for cells treated with BE3 and an sgRNA targeting the *TP53* Y163C mutation, or for cells treated with BE3 and an sgRNA targeting the *VEGFA* locus. Neither BE3-treated sample resulted in mutational rates above those of untreated controls. This figure depicts SEQ ID NOs: 717 to 720 from top to bottom, respectively.

[0072] Figures 36A to 36F show the effects of deaminase, linker length, and linker composition on base editing. Figure 36A shows a gel-based deaminase assay showing activity of rAPOBEC1, pmCDA1, hAID, hAPOBEC3G, rAPOBEC1-GGS-dCas9, rAPOBEC1-(GGS)₃(SEQ ID NO: 596)-dCas9, and dCas9-(GGS)₃(SEQ ID NO: 596)-rAPOBEC1 on ssDNA. Enzymes were expressed in a mammalian cell lysate-derived *in vitro* transcription-translation system and incubated with 1.8 μM dye-conjugated ssDNA and USER enzyme (uracil DNA glycosylase and endonuclease VIII) at 37 °C for 2 hours. The resulting DNA was resolved on a denaturing polyacrylamide gel and imaged. The positive control is a sequence with a U synthetically incorporated at the same position as the target C. Figure 36B shows coomassie-stained denaturing PAGE gel of the expressed and purified proteins used in Figures 36C to 36F. Figures 36C to 36F show gel-based deaminase assay showing the deamination window of base editors with deaminase-Cas9 linkers of GGS (Figure 36C), (GGS)₃ (SEQ ID NO: 596) (Figure 36D), XTEN (Figure 36E), or (GGS)₇ (SEQ ID NO: 597) (Figure 36F). Following incubation of 1.85 μM deaminase-dCas9 fusions complexed with sgRNA with 125 nM dsDNA substrates at 37 °C for 2 hours, the dye-conjugated DNA was isolated and incubated with USER enzyme at 37 °C for 1 hour to cleave

the DNA backbone at the site of any uracils. The resulting DNA was resolved on a denaturing polyacrylamide gel, and the dye-conjugated strand was imaged. Each lane is numbered according to the position of the target C within the protospacer, or with – if no target C is present. 8U is a positive control sequence with a U synthetically incorporated at position 8. .

[0073] Figures 37A to 37C show BE1 base editing efficiencies are dramatically decreased in mammalian cells. Figure 37A Protospacer (black and red) and PAM (blue) sequences of the six mammalian cell genomic loci targeted by base editors. Target Cs are indicated in red with subscripted numbers corresponding to their positions within the protospacer. Figure 37B shows synthetic 80-mers with sequences matching six different genomic sites were incubated with BE1 then analyzed for base editing by HTS. For each site, the sequence of the protospacer is indicated to the right of the name of the site, with the PAM highlighted in blue. Underneath each sequence are the percentages of total DNA sequencing reads with the corresponding base. We considered a target C as “editable” if the *in vitro* conversion efficiency is >10%. Note that maximum yields are 50% of total DNA sequencing reads since the non-targeted strand is unaffected by BE1. Values are shown from a single experiment. Figure 37C shows HEK293T cells were transfected with plasmids expressing BE1 and an appropriate sgRNA. Three days after transfection, genomic DNA was extracted and analyzed by high-throughput DNA sequencing at the six loci. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with Ts at the target positions indicated, are shown for BE1 at all six genomic loci. Values and error bars of all data from HEK293T cells reflect the mean and standard deviation of three independent biological replicates performed on different days. Figure 37A depicts SEQ ID NOS: 721 to 726 from top to bottom, respectively. Figure 37B depicts SEQ ID NOS: 727 to 732 from top to bottom, respectively.

[0074] Figure 38 shows base editing persists over multiple cell divisions. Cellular C to T conversion percentages by BE2 and BE3 are shown for HEK293 sites 3 and 4 in HEK293T cells before and after passaging the cells. HEK293T cells were nucleofected with plasmids expressing BE2 or BE3 and an sgRNA targeting HEK293 site 3 or 4. Three days after nucleofection, the cells were harvested and split in half. One half was subjected to HTS analysis, and the other half was allowed to propagate for approximately five cell divisions, then harvested and subjected to HTS analysis. Values and error bars reflect the mean and standard deviation of at least two biological experiments.

[0075] **Figures 39A to 39C** show non-target C/G mutation rates. Shown here are the C to T and G to A mutation rates at 2,500 distinct cytosines and guanines surrounding the six on-target and 34 off-target loci tested, representing a total of 14,700,000 sequence reads derived from approximately 1.8×10^6 cells. Figures 39A and 39B show cellular non-target C to T and G to A conversion percentages by BE1, BE2, and BE3 are plotted individually against their positions relative to a protospacer for all 2,500 cytosines/guanines. The side of the protospacer distal to the PAM is designated with positive numbers, while the side that includes the PAM is designated with negative numbers. Figure 39C shows average non-target cellular C to T and G to A conversion percentages by BE1, BE2, and BE3 are shown, as well as the highest and lowest individual conversion percentages.

[0076] **Figures 40A to 40B** show additional data sets of BE3-mediated correction of two disease-relevant mutations in mammalian cells. For each site, the sequence of the protospacer is indicated to the right of the name of the mutation, with the PAM highlighted in blue and the base responsible for the mutation indicated in red bold with a subscripted number corresponding to its position within the protospacer. The amino acid sequence above each disease-associated allele is shown, together with the corrected amino acid sequence following base editing in green. Underneath each sequence are the percentages of total sequencing reads with the corresponding base. Cells were nucleofected with plasmids encoding BE3 and an appropriate sgRNA. Two days after nucleofection, genomic DNA was extracted from the nucleofected cells and analyzed by HTS to assess pathogenic mutation correction. Figure 40A shows the Alzheimer's disease-associated *APOE4* allele is converted to *APOE3r* in mouse astrocytes by BE3 in 58.3% of total reads only when treated with the correct sgRNA. Two nearby Cs are also converted to Ts, but with no change to the predicted sequence of the resulting protein. Identical treatment of these cells with wt Cas9 and donor ssDNA results in 0.2% correction, with 26.7% indel formation. Figure 40B shows the cancer-associated p53 Y163C mutation is corrected by BE3 in 3.3% of nucleofected human breast cancer cells only when treated with the correct sgRNA. Identical treatment of these cells with wt Cas9 and donor ssDNA results in no detectable mutation correction with 8.0% indel formation. Figures 40A to 40B depict SEQ ID NOs: 733 to 740 from top to bottom, respectively.

[0077] **Figure 41** shows a schematic representation of an exemplary USER (Uracil-Specific Excision Reagent) Enzyme-based assay, which may be used to test the activity of various deaminases on single-stranded DNA (ssDNA) substrates.

[0078] **Figure 42** is a schematic of the pmCDA-nCas9-UGI-NLS construct and its activity at the HeK-3 site relative to the base editor (rAPOBEC1) and the negative control (untreated).

[0079] **Figure 43** is a schematic of the pmCDA1-XTEN-nCas9-UGI-NLS construct and its activity at the HeK-3 site relative to the base editor (rAPOBEC1) and the negative control (untreated).

[0080] **Figure 44** shows the percent of total sequencing reads with target C converted to T using cytidine deaminases (CDA) or APOBEC.

[0081] **Figure 45** shows the percent of total sequencing reads with target C converted to A using deaminases (CDA) or APOBEC.

[0082] **Figure 46** shows the percent of total sequencing reads with target C converted to G using deaminases (CDA) or APOBEC.

[0083] **Figure 47** is a schematic of the huAPOBEC3G-XTEN-nCas9-UGI-NLS construct and its activity at the HeK-2 site relative to a mutated form (huAPOBEC3G*(D316R_D317R)-XTEN-nCas9-UGI-NLS, the base editor (rAPOBEC1) and the negative control (untreated).

[0084] **Figure 48** shows the schematic of the LacZ construct used in the selection assay of Example 7.

[0085] **Figure 49** shows reversion data from different plasmids and constructs.

[0086] **Figure 50** shows the verification of lacZ reversion and the purification of reverted clones.

[0087] **Figure 51** is a schematic depicting a deamination selection plasmid used in Example 7.

[0088] **Figure 52** shows the results of a chloramphenicol reversion assay (pmCDA1 fusion).

[0089] **Figures 53A to 53B** demonstrated DNA correction induction of two constructs.

[0090] **Figure 54** shows the results of a chloramphenicol reversion assay (huAPOBEC3G fusion).

[0091] **Figure 55** shows the activities of BE3 and HF-BE3 at EMX1 off-targets. The sequences, from top to bottom, correspond to SEQ ID NOs: 286-292, 299-301.

[0092] **Figure 56** shows on-target base editing efficiencies of BE3 and HF-BE3.

[0093] **Figure 57** is a graph demonstrating that mutations affect cytidine deamination with varying degrees. Combinations of mutations that each slightly impairs catalysis allow

selective deamination at one position over others. The FANCF site was GGAATC₆C₇C₈TTC₁₁TGCAGCACCTGG (SEQ ID NO: 303).

[0094] **Figure 58** is a schematic depicting next generation base editors.

[0095] **Figure 59** is a schematic illustrating new base editors made from Cas9 variants.

[0096] **Figure 60** shows the base-edited percentage of different NGA PAM sites.

[0097] **Figure 61** shows the base-edited percentage of cytidines using NGCG PAM EMX (VRER BE3) and the C₁TC₃C₄C₅ATC₈AC₁₀ATCAACCGGT (SEQ ID NO: 304) spacer.

[0098] **Figure 62** shows the based-edited percentages resulting from different NNGRRT PAM sites.

[0099] **Figure 63** shows the based-edited percentages resulting from different NNHRRT PAM sites.

[00100] **Figures 64A to 64C** show the base-edited percentages resulting from different TTTN PAM sites using Cpf1 BE2. The spacers used were:

TTTCCTC₃C₄C₅C₆C₇C₈C₉AC₁₁AGGTAGAACAT (Figure 64A, SEQ ID NO: 305),

TTTCC₁C₂TC₄TGTC₈C₉AC₁₁ACCCTCATCCTG (Figure 64B, SEQ ID NO: 306), and

TTTCC₁C₂C₃AGTC₇C₈TC₁₀C₁₁AC₁₃AC₁₅C₁₆C₁₇TGAAAC (Figure 64C, SEQ ID NO: 307).

[00101] **Figure 65** is a schematic depicting selective deamination as achieved through kinetic modulation of cytidine deaminase point mutagenesis.

[00102] **Figure 66** is a graph showing the effect of various mutations on the deamination window probed in cell culture with multiple cytidines in the spacer. The spacer used was: TGC₃C₄C₅C₆TC₈C₉C₁₀TC₁₂C₁₃C₁₄TGGCCC (SEQ ID NO: 308).

[00103] **Figure 67** is a graph showing the effect of various mutations on the deamination window probed in cell culture with multiple cytidines in the spacer. The spacer used was: AGAGC₅C₆C₇C₈C₉C₁₀C₁₁TC₁₃AAAGAGA (SEQ ID NO: 309).

[00104] **Figure 68** is a graph showing the effect of various mutations on the FANCF site with a limited number of cytidines. The spacer used was: GGAATC₆C₇C₈TTC₁₁TGCAGCACCTGG (SEQ ID NO: 303). Note that the triple mutant (W90Y, R126E, R132E) preferentially edits the cytidine at the sixth position.

[00105] **Figure 69** is a graph showing the effect of various mutations on the HEK3 site with a limited number of cytidines. The spacer used was: GGCC₄C₅AGACTGAGCACGTGATGG (SEQ ID NO: 310). Note that the double and triple mutants preferentially edit the cytidine at the fifth position over the cytidine in the fourth position.

[00106] **Figure 70** is a graph showing the effect of various mutations on the EMX1 site with a limited number of cytidines. The spacer used was:

GAGTC₅C₆GAGCAGAAGAAGAAGGG (SEQ ID NO: 311). Note that the triple mutant only edits the cytidine at the fifth position, not the sixth.

[00107] **Figure 71** is a graph showing the effect of various mutations on the HEK2 site with a limited number of cytidines. The spacer used was:

GAAC₄AC₆AAAGCATAGACTGCGGG (SEQ ID NO: 312).

[00108] **Figure 72** shows on-target base editing efficiencies of BE3 and BE3 comprising mutations W90Y R132E in immortalized astrocytes.

[00109] **Figure 73** depicts a schematic of three Cpf1 fusion constructs.

[00110] **Figures 74** shows a comparison of plasmid delivery of BE3 and HF-BE3 (EMX1, FANCF, and RNF2).

[00111] **Figure 75** shows a comparison of plasmid delivery of BE3 and HF-BE3 (HEK3 and HEK 4).

[00112] **Figure 76** shows off-target editing of EMX-1 at all 10 sites.

[00113] **Figure 77** shows deaminase protein lipofection to HEK cells using a GAGTCCGAGCAGAAGAAGAAG (SEQ ID NO: 313) spacer. The EMX-1 on-target and EMX-1 off target site 2 were examined.

[00114] **Figure 78** shows deaminase protein lipofection to HEK cells using a GGAATCCCTTCTGCAGCACCTGG (SEQ ID NO: 314) spacer. The FANCF on target and FANCF off target site 1 were examined.

[00115] **Figure 79** shows deaminase protein lipofection to HEK cells using a GGCCAGACTGAGCACGTGA (SEQ ID NO: 315) spacer. The HEK-3 on target site was examined.

[00116] **Figure 80** shows deaminase protein lipofection to HEK cells using a GGCAGTGCAGGCTGGAGGTGGGGG (SEQ ID NO: 316) spacer. The HEK-4 on target, off target site 1, site 3, and site 4.

[00117] **Figure 81** shows the results of an *in vitro* assay for sgRNA activity for sgHR_13 (GTCAGGTCGAGGGTTCTGTC (SEQ ID NO: 317) spacer; C8 target: G51 to STOP), sgHR_14 (GGGCCGAGTATCCTCACTC (SEQ ID NO: 318) spacer; C7 target; C7 target: Q68 to STOP), and sgHR_15 (CCGCCAGTCCCAGTACGGGA (SEQ ID NO: 319) spacer; C10 and C11 are targets: W239 or W237 to STOP).

[00118] **Figure 82** shows the results of an *in vitro* assay for sgHR_17 (CAACCACTGCTCAAAGATGC (SEQ ID NO: 320) spacer; C4 and C5 are targets: W410

to STOP), and sgHR_16 (CTTCCAGGATGAGAACACAG (SEQ ID NO: 321) spacer; C4 and C5 are targets: W273 to STOP).

[00119] **Figure 83** shows the direct injection of BE3 protein complexed with sgHR_13 in zebrafish embryos.

[00120] **Figure 84** shows the direct injection of BE3 protein complexed with sgHR_16 in zebrafish embryos.

[00121] **Figure 85** shows the direct injection of BE3 protein complexed with sgHR_17 in zebrafish embryos.

[00122] **Figure 86** shows exemplary nucleic acid changes that may be made using base editors that are capable of making a cytosine to thymine change.

[00123] **Figure 87** shows an illustration of apolipoprotein E (APOE) isoforms, demonstrating how a base editor (*e.g.*, BE3) may be used to edit one APOE isoform (*e.g.*, APOE4) into another APOE isoform (*e.g.*, APOE3r) that is associated with a decreased risk of Alzheimer's disease.

[00124] **Figure 88** shows base editing of APOE4 to APOE3r in mouse astrocytes.

[00125] **Figure 89** shows base editing of PRNP to cause early truncation of the protein at arginine residue 37.

[00126] **Figure 90** shows that knocking out UDG (which UGI inhibits) dramatically improves the cleanliness of efficiency of C to T base editing.

[00127] **Figure 91** shows that use of a base editor with the nickase but without UGI leads to a mixture of outcomes, with very high indel rates.

[00128] **Figures 92A to 92G** show that SaBE3, SaKKH-BE3, VQR-BE3, EQR-BE3, and VRER-BE3 mediate efficient base editing at target sites containing non-NGG PAMs in human cells. Figure 92A shows base editor architectures using *S. pyogenes* and *S. aureus* Cas9. Figure 92B shows recently characterized Cas9 variants with alternate or relaxed PAM requirements. Figures 92C and 92D show HEK293T cells treated with the base editor variants shown as described in Example 12. The percentage of total DNA sequencing reads (with no enrichment for transfected cells) with C converted to T at the target positions indicated are shown. The PAM sequence of each target tested is shown below the X-axis. The charts show the results for SaBE3 and SaKKH-BE3 at genomic loci with NNGRRT PAMs (Figure 92C), SaBE3 and SaKKH-BE3 at genomic loci with NNNRRT PAMs (Figure 92D), VQR-BE3 and EQR-BE3 at genomic loci with NGAG PAMs (Figure 92E), and with NGAH PAMs (Figure 92F), and VRER-BE3 at genomic loci with NGCG PAMs (Figure 92G).

Values and error bars reflect the mean and standard deviation of at least two biological replicates.

[00129] **Figures 93A to 93C** demonstrate that base editors with mutations in the cytidine deaminase domain exhibit narrowed editing windows. Figures 93A to 93C show HEK293T cells transfected with plasmids expressing mutant base editors and an appropriate sgRNA. Three days after transfection, genomic DNA was extracted and analyzed by high-throughput DNA sequencing at the indicated loci. The percentage of total DNA sequencing reads (without enrichment for transfected cells) with C changed to T at the target positions indicated are shown for the EMX1 site, HEK293 site 3, FANCF site, HEK293 site 2, site A, and site B loci. Figure 93A illustrates certain cytidine deaminase mutations which narrow the base editing window. See Figure 98 for the characterization of additional mutations. Figure 93B shows the effect of cytidine deaminase mutations which effect the editing window width on genomic loci. Combining beneficial mutations has an additive effect on narrowing the editing window. Figure 93C shows that YE1-BE3, YE2-BE3, EE-BE3, and YEE-BE3 effect the product distribution of base editing, producing predominantly singly-modified products in contrast with BE3. Values and error bars reflect the mean and standard deviation of at least two biological replicates.

[00130] **Figures 94A and 94B** show genetic variants from ClinVar that in principle can be corrected by the base editors developed in this work. The NCBI ClinVar database of human genetic variations and their corresponding phenotypes was searched for genetic diseases that in theory can be corrected by base editing. Figure 94A demonstrates improvement in base editing targeting scope among all pathogenic T→C mutations in the ClinVar database through the use of base editors with altered PAM specificities. The white fractions denote the proportion of pathogenic T→C mutations accessible on the basis of the PAM requirements of either BE3, or BE3 together with the five modified-PAM base editors developed in this work. Figure 94B shows improvement in base editing targeting scope among all pathogenic T→C mutations in the ClinVar database through the use of base editors with narrowed activity windows. BE3 was assumed to edit Cs in positions 4-8 with comparable efficiency as shown in Figures 93A to 93C. YEE-BE3 was assumed to edit with C5>C6>C7>others preference within its activity window. The white fractions denote the proportion of pathogenic T→C mutations that can be edited BE3 without comparable editing of other Cs (left), or that can be edited BE3 or YEE-BE3 without comparable editing of other Cs (right).

[00131] **Figures 95A to 95C** show the effect of truncated guide RNAs on base editing window width. HEK293T cells were transfected with plasmids expressing BE3 and sgRNAs

of different 5' truncation lengths. The treated cells were analyzed as described in the Examples. **Figure 95A** shows protospacer and PAM sequence (top, SEQ ID NO: 4270) and cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with Ts at the target positions indicated, at a site within the EMX1 genomic locus. At this site, the base editing window was altered through the use of a 17-nt truncated gRNA. **Figure 95B** shows protospacer and PAM sequences (top, SEQ ID NO: 4270) and cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with Ts at the target positions indicated, at sites within the HEK site 3 and site 4 genomic loci. At these sites, no change in the base editing window was observed, but a linear decrease in editing efficiency for all substrate bases as the sgRNA is truncated was noted.

[00132] **Figure 96** shows the effect of APOBEC1-Cas9 linker lengths on base editing window width. HEK293T cells were transfected with plasmids expressing base editors with rAPOBEC1-Cas9 linkers of XTEN, GGS, (GGS)₃ (SEQ ID NO: 596), (GGS)₅ (SEQ ID NO: 4271), or (GGS)₇ (SEQ ID NO: 597) and an sgRNA. The treated cells were analyzed as described in the Examples. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with Ts at the target positions indicated, are shown for the various base editors with different linkers.

[00133] **Figures 97A to 97C** show the effect of rAPOBEC mutations on base editing window width. **Figure 97C** shows HEK293T cells transfected with plasmids expressing an sgRNA targeting either Site A or Site B and the BE3 point mutants indicated. The treated cells were analyzed as described in the Examples. All C's in the protospacer and within three basepairs of the protospacer are displayed and the cellular C to T conversion percentages are shown. The 'editing window widths', defined as the calculated number of nucleotides within which editing efficiency exceeds the half-maximal value, are displayed for all tested mutants.

[00134] **Figure 98** shows the effect of APOBEC1 mutation on product distributions of base editing in mammalian cells. HEK293T cells were transfected with plasmids expressing BE3 or its mutants and an appropriate sgRNAs. The treated cells were analyzed as described in the Examples. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with Ts at the target positions indicated, are shown (left). Percent of total sequencing reads containing the C to T conversion is shown on the right. The BE3 point mutants do not significantly affect base editing efficiencies at HEK site 4, a site with only one target cytidine.

[00135] **Figure 99** shows a comparison of on-target editing plasma delivery in BE3 and HF-BE3.

[00136] **Figure 100** shows a comparison of on-target editing in protein and plasma delivery of BE3.

[00137] **Figure 101** shows a comparison of on-target editing in protein and plasma delivery of HF-BE3.

[00138] **Figure 102** shows that both lipofection and installing HF mutations decrease off-target deamination events. The diamond indicates no off targets were detected and the specificity ratio was set to 100.

[00139] **Figure 103** shows *in vitro* C to T editing on a synthetic substrate with Cs placed at even positions in the protospacer (NNNNTC₂TC₄TC₆TC₈TC₁₀TC₁₂TC₁₄TC₁₆TC₁₈TC₂₀NGG, SEQ ID NO: 4272).

[00140] **Figure 104** shows *in vitro* C to T editing on a synthetic substrate with Cs placed at odd positions in the protospacer (NNNNTC₂TC₄TC₆TC₈TC₁₀TC₁₂TC₁₄TC₁₆TC₁₈TC₂₀NGG, SEQ ID NO: 4272).

[00141] **Figure 105** includes two graphs depicting the specificity ratio of base editing with plasmid vs. protein delivery.

[00142] **Figures 106A to 106B** shows BE3 activity on non-NGG PAM sites. HEK293T cells were transfected with plasmids expressing BE3 and appropriate sgRNA. The treated cells were analyzed as described in the Examples. **Figure 106A** shows BE3 activity on sites can be efficiently targeted by SaBE3 or SaKKH-BE3. BE3 shows low but significant activity on the NAG PAM. **Figure 106B** shows BE3 has significantly reduced editing at sites with NGA or NGCG PAMs, in contrast to VQR-BE3 or VRER-BE3.

[00143] **Figures 107A to 107B** show the effect of APOBEC1 mutations on VQR-BE3 and SaKKH-BE3. HEK293T cells were transfected with plasmids expressing VQR-BE3, SaKKH-BE3 or its mutants and an appropriate sgRNAs. The treated cells were analyzed as described in the Methods. Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with Ts at the target positions indicated, are shown. **Figure 107A** shows that the window-modulating mutations can be applied to VQR-BE3 to enable selective base editing at sites targetable by NGA PAM. **Figure 107B** shows that, when applied to SaKKH-BE3, the mutations cause overall decrease in base editing efficiency without conferring base selectivity within the target window.

[00144] **Figure 108** shows a schematic representation of nucleotide editing. The following abbreviations are used: (MMR) - mismatch repair, (BE3 Nickase) - refers to base editor 3, which comprises a Cas9 nickase domain, (UGI) - uracil glycosylase inhibitor, (UDG) - uracil DNA glycosylase, (APOBEC) – refers to an APOBEC cytidine deaminase.

DEFINITIONS

[00145] As used herein and in the claims, the singular forms “a,” “an,” and “the” include the singular and the plural reference unless the context clearly indicates otherwise. Thus, for example, a reference to “an agent” includes a single agent and a plurality of such agents.

[00146] The term “Cas9” or “Cas9 nuclease” refers to an RNA-guided nuclease comprising a Cas9 protein, or a fragment thereof (*e.g.*, a protein comprising an active, inactive, or partially active DNA cleavage domain of Cas9, and/or the gRNA binding domain of Cas9). A Cas9 nuclease is also referred to sometimes as a casn1 nuclease or a CRISPR (clustered regularly interspaced short palindromic repeat)-associated nuclease. CRISPR is an adaptive immune system that provides protection against mobile genetic elements (viruses, transposable elements and conjugative plasmids). CRISPR clusters contain spacers, sequences complementary to antecedent mobile elements, and target invading nucleic acids. CRISPR clusters are transcribed and processed into CRISPR RNA (crRNA). In type II CRISPR systems correct processing of pre-crRNA requires a trans-encoded small RNA (tracrRNA), endogenous ribonuclease 3 (*rnc*) and a Cas9 protein. The tracrRNA serves as a guide for ribonuclease 3-aided processing of pre-crRNA. Subsequently, Cas9/crRNA/tracrRNA endonucleolytically cleaves linear or circular dsDNA target complementary to the spacer. The target strand not complementary to crRNA is first cut endonucleolytically, then trimmed 3’-5’ exonucleolytically. In nature, DNA-binding and cleavage typically requires protein and both RNAs. However, single guide RNAs (“sgRNA”, or simply “gRNA”) can be engineered so as to incorporate aspects of both the crRNA and tracrRNA into a single RNA species. See, *e.g.*, Jinek M., Chylinski K., Fonfara I., Hauer M., Doudna J.A., Charpentier E. *Science* 337:816-821(2012), the entire contents of which is hereby incorporated by reference. Cas9 recognizes a short motif in the CRISPR repeat sequences (the PAM or protospacer adjacent motif) to help distinguish self versus non-self. Cas9 nuclease sequences and structures are well known to those of skill in the art (see, *e.g.*, “Complete genome sequence of an M1 strain of *Streptococcus pyogenes*.” Ferretti *et al.*, J.J., McShan W.M., Ajdic D.J., Savic D.J., Savic G., Lyon K., Primeaux C., Sezate S., Suvorov A.N., Kenton S., Lai H.S., Lin S.P., Qian Y., Jia H.G., Najjar F.Z., Ren Q., Zhu H., Song L., White J., Yuan X., Clifton S.W., Roe B.A., McLaughlin R.E., *Proc. Natl. Acad. Sci. U.S.A.* 98:4658-4663(2001); “CRISPR RNA maturation by trans-encoded small RNA and host factor RNase III.” Deltcheva E., Chylinski K., Sharma C.M., Gonzales K., Chao Y., Pirzada Z.A., Eckert M.R., Vogel J., Charpentier E., *Nature* 471:602-607(2011); and “A

programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity.” Jinek M., Chylinski K., Fonfara I., Hauer M., Doudna J.A., Charpentier E. *Science* 337:816-821(2012), the entire contents of each of which are incorporated herein by reference). Cas9 orthologs have been described in various species, including, but not limited to, *S. pyogenes* and *S. thermophilus*. Additional suitable Cas9 nucleases and sequences will be apparent to those of skill in the art based on this disclosure, and such Cas9 nucleases and sequences include Cas9 sequences from the organisms and loci disclosed in Chylinski, Rhun, and Charpentier, “The tracrRNA and Cas9 families of type II CRISPR-Cas immunity systems” (2013) *RNA Biology* 10:5, 726-737; the entire contents of which are incorporated herein by reference. In some embodiments, a Cas9 nuclease has an inactive (*e.g.*, an inactivated) DNA cleavage domain, that is, the Cas9 is a nickase.

[00147] A nuclease-inactivated Cas9 protein may interchangeably be referred to as a “dCas9” protein (for nuclease-“dead” Cas9). Methods for generating a Cas9 protein (or a fragment thereof) having an inactive DNA cleavage domain are known (See, *e.g.*, Jinek *et al.*, *Science*. 337:816-821(2012); Qi *et al.*, “Repurposing CRISPR as an RNA-Guided Platform for Sequence-Specific Control of Gene Expression” (2013) *Cell*. 28;152(5):1173-83, the entire contents of each of which are incorporated herein by reference). For example, the DNA cleavage domain of Cas9 is known to include two subdomains, the HNH nuclease subdomain and the RuvC1 subdomain. The HNH subdomain cleaves the strand complementary to the gRNA, whereas the RuvC1 subdomain cleaves the non-complementary strand. Mutations within these subdomains can silence the nuclease activity of Cas9. For example, the mutations D10A and H840A completely inactivate the nuclease activity of *S. pyogenes* Cas9 (Jinek *et al.*, *Science*. 337:816-821(2012); Qi *et al.*, *Cell*. 28;152(5):1173-83 (2013)). In some embodiments, proteins comprising fragments of Cas9 are provided. For example, in some embodiments, a protein comprises one of two Cas9 domains: (1) the gRNA binding domain of Cas9; or (2) the DNA cleavage domain of Cas9. In some embodiments, proteins comprising Cas9 or fragments thereof are referred to as “Cas9 variants.” A Cas9 variant shares homology to Cas9, or a fragment thereof. For example a Cas9 variant is at least about 70% identical, at least about 80% identical, at least about 90% identical, at least about 95% identical, at least about 96% identical, at least about 97% identical, at least about 98% identical, at least about 99% identical, at least about 99.5% identical, or at least about 99.9% identical to wild type Cas9. In some embodiments, the Cas9 variant may have 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 21, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50 or more amino acid

changes compared to wild type Cas9. In some embodiments, the Cas9 variant comprises a fragment of Cas9 (e.g., a gRNA binding domain or a DNA-cleavage domain), such that the fragment is at least about 70% identical, at least about 80% identical, at least about 90% identical, at least about 95% identical, at least about 96% identical, at least about 97% identical, at least about 98% identical, at least about 99% identical, at least about 99.5% identical, or at least about 99.9% identical to the corresponding fragment of wild type Cas9. In some embodiments, the fragment is at least 30%, at least 35%, at least 40%, at least 45%, at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95% identical, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% of the amino acid length of a corresponding wild type Cas9.

[00148] In some embodiments, the fragment is at least 100 amino acids in length. In some embodiments, the fragment is at least 100, 150, 200, 250, 300, 350, 400, 450, 500, 550, 600, 650, 700, 750, 800, 850, 900, 950, 1000, 1050, 1100, 1150, 1200, 1250, or at least 1300 amino acids in length. In some embodiments, wild type Cas9 corresponds to Cas9 from *Streptococcus pyogenes* (NCBI Reference Sequence: NC_017053.1, SEQ ID NO:1 (nucleotide); SEQ ID NO:2 (amino acid)).

ATGGATAAGAAATACTCAATAGGCTTAGATATCGGCACAAATAGCGTCGGATGG
 GCGGTGATCACTGATGATTATAAGGTTCCGTCTAAAAAGTTCAAGGTTCTGGGAA
 ATACAGACCGCCACAGTATCAAAAAAATCTTATAGGGGCTCTTTTATTTGGCAG
 TGGAGAGACAGCGGAAGCGACTCGTCTCAAACGGACAGCTCGTAGAAGGTATAC
 ACGTCGGAAGAATCGTATTTGTTATCTACAGGAGATTTTTTCAAATGAGATGGCG
 AAAGTAGATGATAGTTTCTTTCATCGACTTGAAGAGTCTTTTTTGGTGAAGAAG
 ACAAGAAGCATGAACGTCATCCTATTTTTGGAAATATAGTAGATGAAGTTGCTTA
 TCATGAGAAATATCCAACATCTATCATCTGCGAAAAAATTGGCAGATTCTACT
 GATAAAGCGGATTTGCGCTTAATCTATTTGGCCTTAGCGCATATGATTAAGTTTC
 GTGGTCATTTTTTATTGAGGGAGATTTAAATCCTGATAATAGTGATGTGGACAA
 ACTATTTATCCAGTTGGTACAAATCTACAATCAATTATTTGAAGAAAACCCTATT
 AACGCAAGTAGAGTAGATGCTAAAGCGATTCTTTCTGCACGATTGAGTAAATCA
 AGACGATTAGAAAATCTCATTGCTCAGCTCCCCGGTGAGAAGAGAAATGGCTTG
 TTTGGGAATCTCATTGCTTTGTCATTGGGATTGACCCCTAATTTTAAATCAAATTT
 TGATTTGGCAGAAGATGCTAAATTACAGCTTTCAAAAGATACTTACGATGATGAT
 TTAGATAATTTATTGGCGCAAATTGGAGATCAATATGCTGATTTGTTTTTGGCAG
 CTAAGAATTTATCAGATGCTATTTTACTTTCAGATATCCTAAGAGTAAATAGTGA
 AATAACTAAGGCTCCCCTATCAGCTTCAATGATTAAGCGCTACGATGAACATCAT
 CAAGACTTGACTCTTTTAAAAGCTTTAGTTTCGACAACAACCTCCAGAAAAGTATA
 AAGAAATCTTTTTTGATCAATCAAAAAACGGATATGCAGGTTATATTGATGGGGG
 AGCTAGCCAAGAAGAATTTTATAAATTTATCAAACCAATTTTAGAAAAAATGGAT
 GGTACTGAGGAATTATTGGTGAAACTAAATCGTGAAGATTTGCTGCGCAAGCAA
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(SEQ ID NO:1)

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 DLNPDNSDVKLFIQLVQIYNQLFEENPINASRVDAKAILSARLSKSRLENLIAQLPG
 EKRNLFGNLIASLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLDNLLAQIGDQYAD
 LFLAAKNLSDAILLSDILRVNSEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEK
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 WMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEK VLPKHSLLYEYFTV
 YNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFD
 SVEISGVEDRFNASLGAYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDRGMIEER
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 VEK GKSKKLKSVKELLGITIMERSSEFKNPIDFLEAKGYKEVKKDLIKLPKYSLFELE
 NGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGKSPEDNEQKQLFVEQH
 HYLDEIIEQISEFSKR VILADANLDKVL SAYNKH RDKPIREQAENIIHLFTLTNLGAPA
 AFKYFDTTIDRKRYTSTKEVLDATLIHQ SITGLYETRIDLSQLGGD (SEQ ID NO:2)
 (single underline: HNH domain; double underline: RuvC domain)

[00149] In some embodiments, wild type Cas9 corresponds to, or comprises SEQ ID NO:3
(nucleotide) and/or SEQ ID NO: 4 (amino acid):

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 TGGCGAAACGGCAGAGGGCGACTCGCCTGAAACGAACCGCTCGGAGAAGGTATAC
 ACGTCGCAAGAACCGAATATGTTACTTACAAGAAATTTT TAGCAATGAGATGGCC
 AAAGTTGACGATTCTTTCTTTCACCGTTTGGAAAGAGTCCTTCCTTGTCGAAGAGG
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 CCGTGGGCACTTTCTCATTGAGGGTGATCTAAATCCGGACA ACTCGGATGTCGAC
 AA ACTGTT CATCCAGTTAGTACAAACCTATAATCAGTTGTTTGAAGAGAACCCTA
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 CCGACGGCTAGAAAACCTGATCGCACAAATTACCCGGAGAGAAGAAAAATGGGTT
 GTTCGGTAACCTTATAGCGCTCTCACTAGGCCTGACACCAAATTTTAAGTCGAAC
 TTCGACTTAGCTGAAGATGCCAAATTGCAGCTTAGTAAGGACACGTACGATGAC
 GATCTCGACAATCTACTGGCACAAATTGGAGATCAGTATGCGGACTTATTTTTGG
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GCATGCTATACTTAGAAGGCAGGAGGATTTTTATCCGTTCCCTCAAAGACAATCGT
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HKHYLDEIIEQISEFSKRVLADANLDKVLSAYNKHRDKPIREQAENIIHLFTLTNLGA
PAAFKYFDTTIDRKRYTSTKEVLDTLIHQSI TGLYETRIDLSQLGGD (SEQ ID NO:4)
 (single underline: HNH domain; double underline: RuvC domain)

[00150] In some embodiments, wild type Cas9 corresponds to Cas9 from *Streptococcus pyogenes* (NCBI Reference Sequence: NC_002737.2, SEQ ID NO: 8 (nucleotide); and Uniport Reference Sequence: Q99ZW2, SEQ ID NO: 10 (amino acid)).

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 TGA (SEQ ID NO: 8)

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POVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVV
 AKVEKKGKSKKLKSVKELLGITIMERSSSFENPIDFLEAKGYKEVKKDLIIKLPKYSLFE
 LENGKRMLASAGELQKGNELALPSKYVNFYLYLASHYEKLGKSPEDNEQKQLFVEQ
 HKHYLDEIIEQISEFSKRVLADANLDKVLSAYNKHRDKPIREQAENIIHLFTLTNLGA
 PAAFKYFDTTIDRKRYTSTKEVLDATLIHQSTGLYETRIDLSQLGGD (SEQ ID NO:
 10) (single underline: HNH domain; double underline: RuvC domain)

[00151] In some embodiments, Cas9 refers to Cas9 from: *Corynebacterium ulcerans* (NCBI Refs: NC_015683.1, NC_017317.1); *Corynebacterium diphtheria* (NCBI Refs: NC_016782.1, NC_016786.1); *Spiroplasma syrphidicola* (NCBI Ref: NC_021284.1); *Prevotella intermedia* (NCBI Ref: NC_017861.1); *Spiroplasma taiwanense* (NCBI Ref: NC_021846.1); *Streptococcus iniae* (NCBI Ref: NC_021314.1); *Belliella baltica* (NCBI Ref: NC_018010.1); *Psychroflexus torquis* (NCBI Ref: NC_018721.1); *Streptococcus thermophilus* (NCBI Ref: YP_820832.1), *Listeria innocua* (NCBI Ref: NP_472073.1), *Campylobacter jejuni* (NCBI Ref: YP_002344900.1) or *Neisseria meningitidis* (NCBI Ref: YP_002342100.1) or to a Cas9 from any of the organisms listed in Example 5.

[00152] In some embodiments, dCas9 corresponds to, or comprises in part or in whole, a Cas9 amino acid sequence having one or more mutations that inactivate the Cas9 nuclease activity. For example, in some embodiments, a dCas9 domain comprises D10A and/or H840A mutation.

dCas9 (D10A and H840A):

MDKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLIGALLFDS
GETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFFHRLEESFLVEEDKK
 HERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLI
 EGDLNPDNSDVDFLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQ
 LPGEKKNGLFGNLIASLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLDNLLAQIGDQ
 YADFLAAKNLSDAILSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQL
 PEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRK
 QRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILFRIPYYVGPLARGNSR
 FAWMTRKSEETITPWNFEVVDKGGASAQSFIERMTNFDKNLPNEKVLPHSLLYEYF
 TVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIEC
 FDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEE
 RLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFAN
 RNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGILQTVKVVDEL
VKVMGRHKPENIVIEMARENQTTQKGOKNSRERMKRIEEGIKELGSQILKEHPVE
NTOLONEKLYLYLQNGRDMYVDQELDINRLSDYDVDAIVPOSFLKDDSIDNKV
LTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLS
ELDKAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSKLVSDFR

KDFQFYK VREINNYHHAHDAYLNAVVG TALIKKYPKLESEFVYGDYKVYDVRKMI
AKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDF
ATVRKVL SMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGF DSP
TVAYSVLVVAKVEKGKSKKLLKSVKELLGITIMERSSEFEKNPIDFLEAKGYKEVKKDLI
IKLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGKSPEDN
EQKQLFVEQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKH RDKPIREQAENIIH
LFTLTNLGAPAAF KYFDTTIDRKRYTSTKEVLDATLIHQ SITGLYETRIDL SQLGGD
(SEQ ID NO: 9) (single underline: HNH domain; double underline: RuvC domain).

[00153] In some embodiments, the Cas9 domain comprises a D10A mutation, while the residue at position 840 remains a histidine in the amino acid sequence provided in SEQ ID NO: 10, or at corresponding positions in any of the amino acid sequences provided in SEQ ID NOs: 11-260. Without wishing to be bound by any particular theory, the presence of the catalytic residue H840 restores the activity of the Cas9 to cleave the non-edited (*e.g.*, non-deaminated) strand containing a G opposite the targeted C. Restoration of H840 (*e.g.*, from A840) does not result in the cleavage of the target strand containing the C. Such Cas9 variants are able to generate a single-strand DNA break (nick) at a specific location based on the gRNA-defined target sequence, leading to repair of the non-edited strand, ultimately resulting in a G to A change on the non-edited strand. A schematic representation of this process is shown in Figure 108. Briefly, the C of a C-G basepair can be deaminated to a U by a deaminase, *e.g.*, an APOBEC deaminase. Nicking the non-edited strand, having the G, facilitates removal of the G via mismatch repair mechanisms. UGI inhibits UDG, which prevents removal of the U.

[00154] In other embodiments, dCas9 variants having mutations other than D10A and H840A are provided, which, *e.g.*, result in nuclease inactivated Cas9 (dCas9). Such mutations, by way of example, include other amino acid substitutions at D10 and H820, or other substitutions within the nuclease domains of Cas9 (*e.g.*, substitutions in the HNH nuclease subdomain and/or the RuvC1 subdomain). In some embodiments, variants or homologues of dCas9 (*e.g.*, variants of SEQ ID NO: 10) are provided which are at least about 70% identical, at least about 80% identical, at least about 90% identical, at least about 95% identical, at least about 98% identical, at least about 99% identical, at least about 99.5% identical, or at least about 99.9% identical to SEQ ID NO: 10. In some embodiments, variants of dCas9 (*e.g.*, variants of SEQ ID NO: 10) are provided having amino acid sequences which are shorter, or longer than SEQ ID NO: 10, by about 5 amino acids, by about 10 amino acids, by about 15 amino acids, by about 20 amino acids, by about 25 amino

acids, by about 30 amino acids, by about 40 amino acids, by about 50 amino acids, by about 75 amino acids, by about 100 amino acids or more.

[00155] In some embodiments, Cas9 fusion proteins as provided herein comprise the full-length amino acid sequence of a Cas9 protein, *e.g.*, one of the Cas9 sequences provided herein. In other embodiments, however, fusion proteins as provided herein do not comprise a full-length Cas9 sequence, but only a fragment thereof. For example, in some embodiments, a Cas9 fusion protein provided herein comprises a Cas9 fragment, wherein the fragment binds crRNA and tracrRNA or sgRNA, but does not comprise a functional nuclease domain, *e.g.*, in that it comprises only a truncated version of a nuclease domain or no nuclease domain at all. Exemplary amino acid sequences of suitable Cas9 domains and Cas9 fragments are provided herein, and additional suitable sequences of Cas9 domains and fragments will be apparent to those of skill in the art.

[00156] In some embodiments, Cas9 refers to Cas9 from: *Corynebacterium ulcerans* (NCBI Refs: NC_015683.1, NC_017317.1); *Corynebacterium diphtheria* (NCBI Refs: NC_016782.1, NC_016786.1); *Spiroplasma syrphidicola* (NCBI Ref: NC_021284.1); *Prevotella intermedia* (NCBI Ref: NC_017861.1); *Spiroplasma taiwanense* (NCBI Ref: NC_021846.1); *Streptococcus iniae* (NCBI Ref: NC_021314.1); *Belliella baltica* (NCBI Ref: NC_018010.1); *Psychroflexus torquisI* (NCBI Ref: NC_018721.1); *Streptococcus thermophilus* (NCBI Ref: YP_820832.1); *Listeria innocua* (NCBI Ref: NP_472073.1); *Campylobacter jejuni* (NCBI Ref: YP_002344900.1); or *Neisseria meningitidis* (NCBI Ref: YP_002342100.1).

[00157] The term “deaminase” or “deaminase domain,” as used herein, refers to a protein or enzyme that catalyzes a deamination reaction. In some embodiments, the deaminase or deaminase domain is a cytidine deaminase, catalyzing the hydrolytic deamination of cytidine or deoxycytidine to uridine or deoxyuridine, respectively. In some embodiments, the deaminase or deaminase domain is a cytidine deaminase domain, catalyzing the hydrolytic deamination of cytosine to uracil. In some embodiments, the deaminase or deaminase domain is a naturally-occurring deaminase from an organism, such as a human, chimpanzee, gorilla, monkey, cow, dog, rat, or mouse. In some embodiments, the deaminase or deaminase domain is a variant of a naturally-occurring deaminase from an organism, that does not occur in nature. For example, in some embodiments, the deaminase or deaminase domain is at least 50%, at least 55%, at least 60%, at least 65%, at least 70%, at least 75% at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to a naturally-occurring deaminase from an organism.

[00158] The term “effective amount,” as used herein, refers to an amount of a biologically active agent that is sufficient to elicit a desired biological response. For example, in some embodiments, an effective amount of a nuclease may refer to the amount of the nuclease that is sufficient to induce cleavage of a target site specifically bound and cleaved by the nuclease. In some embodiments, an effective amount of a fusion protein provided herein, *e.g.*, of a fusion protein comprising a nuclease-inactive Cas9 domain and a nucleic acid editing domain (*e.g.*, a deaminase domain) may refer to the amount of the fusion protein that is sufficient to induce editing of a target site specifically bound and edited by the fusion protein. As will be appreciated by the skilled artisan, the effective amount of an agent, *e.g.*, a fusion protein, a nuclease, a deaminase, a recombinase, a hybrid protein, a protein dimer, a complex of a protein (or protein dimer) and a polynucleotide, or a polynucleotide, may vary depending on various factors as, for example, on the desired biological response, *e.g.*, on the specific allele, genome, or target site to be edited, on the cell or tissue being targeted, and on the agent being used.

[00159] The term “linker,” as used herein, refers to a chemical group or a molecule linking two molecules or moieties, *e.g.*, two domains of a fusion protein, such as, for example, a nuclease-inactive Cas9 domain and a nucleic acid editing domain (*e.g.*, a deaminase domain). In some embodiments, a linker joins a gRNA binding domain of an RNA-programmable nuclease, including a Cas9 nuclease domain, and the catalytic domain of a nucleic-acid editing protein. In some embodiments, a linker joins a dCas9 and a nucleic-acid editing protein. Typically, the linker is positioned between, or flanked by, two groups, molecules, or other moieties and connected to each one via a covalent bond, thus connecting the two. In some embodiments, the linker is an amino acid or a plurality of amino acids (*e.g.*, a peptide or protein). In some embodiments, the linker is an organic molecule, group, polymer, or chemical moiety. In some embodiments, the linker is 5-100 amino acids in length, for example, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 30-35, 35-40, 40-45, 45-50, 50-60, 60-70, 70-80, 80-90, 90-100, 100-150, or 150-200 amino acids in length. Longer or shorter linkers are also contemplated.

[00160] The term “mutation,” as used herein, refers to a substitution of a residue within a sequence, *e.g.*, a nucleic acid or amino acid sequence, with another residue, or a deletion or insertion of one or more residues within a sequence. Mutations are typically described herein by identifying the original residue followed by the position of the residue within the sequence and by the identity of the newly substituted residue. Various methods for making the amino acid substitutions (mutations) provided herein are well known in the art, and are provided by,

for example, Green and Sambrook, *Molecular Cloning: A Laboratory Manual* (4th ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (2012)).

[00161] The terms “nucleic acid” and “nucleic acid molecule,” as used herein, refer to a compound comprising a nucleobase and an acidic moiety, *e.g.*, a nucleoside, a nucleotide, or a polymer of nucleotides. Typically, polymeric nucleic acids, *e.g.*, nucleic acid molecules comprising three or more nucleotides are linear molecules, in which adjacent nucleotides are linked to each other via a phosphodiester linkage. In some embodiments, “nucleic acid” refers to individual nucleic acid residues (*e.g.* nucleotides and/or nucleosides). In some embodiments, “nucleic acid” refers to an oligonucleotide chain comprising three or more individual nucleotide residues. As used herein, the terms “oligonucleotide” and “polynucleotide” can be used interchangeably to refer to a polymer of nucleotides (*e.g.*, a string of at least three nucleotides). In some embodiments, “nucleic acid” encompasses RNA as well as single and/or double-stranded DNA. Nucleic acids may be naturally occurring, for example, in the context of a genome, a transcript, an mRNA, tRNA, rRNA, siRNA, snRNA, a plasmid, cosmid, chromosome, chromatid, or other naturally occurring nucleic acid molecule. On the other hand, a nucleic acid molecule may be a non-naturally occurring molecule, *e.g.*, a recombinant DNA or RNA, an artificial chromosome, an engineered genome, or fragment thereof, or a synthetic DNA, RNA, DNA/RNA hybrid, or including non-naturally occurring nucleotides or nucleosides. Furthermore, the terms “nucleic acid,” “DNA,” “RNA,” and/or similar terms include nucleic acid analogs, *e.g.*, analogs having other than a phosphodiester backbone. Nucleic acids can be purified from natural sources, produced using recombinant expression systems and optionally purified, chemically synthesized, *etc.* Where appropriate, *e.g.*, in the case of chemically synthesized molecules, nucleic acids can comprise nucleoside analogs such as analogs having chemically modified bases or sugars, and backbone modifications. A nucleic acid sequence is presented in the 5' to 3' direction unless otherwise indicated. In some embodiments, a nucleic acid is or comprises natural nucleosides (*e.g.* adenosine, thymidine, guanosine, cytidine, uridine, deoxyadenosine, deoxythymidine, deoxyguanosine, and deoxycytidine); nucleoside analogs (*e.g.*, 2-aminoadenosine, 2-thiothymidine, inosine, pyrrolo-pyrimidine, 3-methyl adenosine, 5-methylcytidine, 2-aminoadenosine, C5-bromouridine, C5-fluorouridine, C5-iodouridine, C5-propynyl-uridine, C5-propynyl-cytidine, C5-methylcytidine, 2-aminoadenosine, 7-deazaadenosine, 7-deazaguanosine, 8-oxoadenosine, 8-oxoguanosine, O(6)-methylguanine, and 2-thiocytidine); chemically modified bases; biologically modified bases (*e.g.*, methylated bases); intercalated bases; modified sugars (*e.g.*, 2'-fluororibose, ribose, 2'-deoxyribose,

arabinose, and hexose); and/or modified phosphate groups (*e.g.*, phosphorothioates and 5'-*N*-phosphoramidite linkages).

[00162] The term “nucleic acid editing domain,” as used herein refers to a protein or enzyme capable of making one or more modifications (*e.g.*, deamination of a cytidine residue) to a nucleic acid (*e.g.*, DNA or RNA). Exemplary nucleic acid editing domains include, but are not limited to a deaminase, a nuclease, a nickase, a recombinase, a methyltransferase, a methylase, an acetylase, an acetyltransferase, a transcriptional activator, or a transcriptional repressor domain. In some embodiments the nucleic acid editing domain is a deaminase (*e.g.*, a cytidine deaminase, such as an APOBEC or an AID deaminase).

[00163] The term “proliferative disease,” as used herein, refers to any disease in which cell or tissue homeostasis is disturbed in that a cell or cell population exhibits an abnormally elevated proliferation rate. Proliferative diseases include hyperproliferative diseases, such as pre-neoplastic hyperplastic conditions and neoplastic diseases. Neoplastic diseases are characterized by an abnormal proliferation of cells and include both benign and malignant neoplasias. Malignant neoplasia is also referred to as cancer.

[00164] The terms “protein,” “peptide,” and “polypeptide” are used interchangeably herein, and refer to a polymer of amino acid residues linked together by peptide (amide) bonds. The terms refer to a protein, peptide, or polypeptide of any size, structure, or function. Typically, a protein, peptide, or polypeptide will be at least three amino acids long. A protein, peptide, or polypeptide may refer to an individual protein or a collection of proteins. One or more of the amino acids in a protein, peptide, or polypeptide may be modified, for example, by the addition of a chemical entity such as a carbohydrate group, a hydroxyl group, a phosphate group, a farnesyl group, an isofarnesyl group, a fatty acid group, a linker for conjugation, functionalization, or other modification, *etc.* A protein, peptide, or polypeptide may also be a single molecule or may be a multi-molecular complex. A protein, peptide, or polypeptide may be just a fragment of a naturally occurring protein or peptide. A protein, peptide, or polypeptide may be naturally occurring, recombinant, or synthetic, or any combination thereof. The term “fusion protein” as used herein refers to a hybrid polypeptide which comprises protein domains from at least two different proteins. One protein may be located at the amino-terminal (N-terminal) portion of the fusion protein or at the carboxy-terminal (C-terminal) protein thus forming an “amino-terminal fusion protein” or a “carboxy-terminal fusion protein,” respectively. A protein may comprise different domains, for example, a nucleic acid binding domain (*e.g.*, the gRNA binding domain of Cas9 that directs the binding of the protein to a target site) and a nucleic acid cleavage domain or a catalytic domain of a

nucleic-acid editing protein. In some embodiments, a protein comprises a proteinaceous part, *e.g.*, an amino acid sequence constituting a nucleic acid binding domain, and an organic compound, *e.g.*, a compound that can act as a nucleic acid cleavage agent. In some embodiments, a protein is in a complex with, or is in association with, a nucleic acid, *e.g.*, RNA. Any of the proteins provided herein may be produced by any method known in the art. For example, the proteins provided herein may be produced via recombinant protein expression and purification, which is especially suited for fusion proteins comprising a peptide linker. Methods for recombinant protein expression and purification are well known, and include those described by Green and Sambrook, *Molecular Cloning: A Laboratory Manual* (4th ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. (2012)), the entire contents of which are incorporated herein by reference.

[00165] The term “RNA-programmable nuclease,” and “RNA-guided nuclease” are used interchangeably herein and refer to a nuclease that forms a complex with (*e.g.*, binds or associates with) one or more RNA that is not a target for cleavage. In some embodiments, an RNA-programmable nuclease, when in a complex with an RNA, may be referred to as a nuclease:RNA complex. Typically, the bound RNA(s) is referred to as a guide RNA (gRNA). gRNAs can exist as a complex of two or more RNAs, or as a single RNA molecule. gRNAs that exist as a single RNA molecule may be referred to as single-guide RNAs (sgRNAs), though “gRNA” is used interchangeably to refer to guide RNAs that exist as either single molecules or as a complex of two or more molecules. Typically, gRNAs that exist as single RNA species comprise two domains: (1) a domain that shares homology to a target nucleic acid (*e.g.*, and directs binding of a Cas9 complex to the target); and (2) a domain that binds a Cas9 protein. In some embodiments, domain (2) corresponds to a sequence known as a tracrRNA, and comprises a stem-loop structure. For example, in some embodiments, domain (2) is identical or homologous to a tracrRNA as provided in Jinek *et al.*, *Science* 337:816-821(2012), the entire contents of which is incorporated herein by reference. Other examples of gRNAs (*e.g.*, those including domain 2) can be found in U.S. Provisional Patent Application, U.S.S.N. 61/874,682, filed September 6, 2013, entitled “Switchable Cas9 Nucleases And Uses Thereof,” and U.S. Provisional Patent Application, U.S.S.N. 61/874,746, filed September 6, 2013, entitled “Delivery System For Functional Nucleases,” the entire contents of each are hereby incorporated by reference in their entirety. In some embodiments, a gRNA comprises two or more of domains (1) and (2), and may be referred to as an “extended gRNA.” For example, an extended gRNA will, *e.g.*, bind two or more Cas9 proteins and bind a target nucleic acid at two or more distinct regions, as

described herein. The gRNA comprises a nucleotide sequence that complements a target site, which mediates binding of the nuclease/RNA complex to said target site, providing the sequence specificity of the nuclease:RNA complex. In some embodiments, the RNA-programmable nuclease is the (CRISPR-associated system) Cas9 endonuclease, for example Cas9 (Csn1) from *Streptococcus pyogenes* (see, e.g., “Complete genome sequence of an M1 strain of *Streptococcus pyogenes*.” Ferretti J.J., McShan W.M., Ajdic D.J., Savic D.J., Savic G., Lyon K., Primeaux C., Sezate S., Suvorov A.N., Kenton S., Lai H.S., Lin S.P., Qian Y., Jia H.G., Najar F.Z., Ren Q., Zhu H., Song L., White J., Yuan X., Clifton S.W., Roe B.A., McLaughlin R.E., Proc. Natl. Acad. Sci. U.S.A. 98:4658-4663(2001); “CRISPR RNA maturation by trans-encoded small RNA and host factor RNase III.” Deltcheva E., Chylinski K., Sharma C.M., Gonzales K., Chao Y., Pirzada Z.A., Eckert M.R., Vogel J., Charpentier E., Nature 471:602-607(2011); and “A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity.” Jinek M., Chylinski K., Fonfara I., Hauer M., Doudna J.A., Charpentier E. Science 337:816-821(2012), the entire contents of each of which are incorporated herein by reference.

[00166] Because RNA-programmable nucleases (e.g., Cas9) use RNA:DNA hybridization to target DNA cleavage sites, these proteins are able to be targeted, in principle, to any sequence specified by the guide RNA. Methods of using RNA-programmable nucleases, such as Cas9, for site-specific cleavage (e.g., to modify a genome) are known in the art (see e.g., Cong, L. *et al.* Multiplex genome engineering using CRISPR/Cas systems. *Science* **339**, 819-823 (2013); Mali, P. *et al.* RNA-guided human genome engineering via Cas9. *Science* **339**, 823-826 (2013); Hwang, W.Y. *et al.* Efficient genome editing in zebrafish using a CRISPR-Cas system. *Nature biotechnology* **31**, 227-229 (2013); Jinek, M. *et al.* RNA-programmed genome editing in human cells. *eLife* **2**, e00471 (2013); Dicarlo, J.E. *et al.* Genome engineering in *Saccharomyces cerevisiae* using CRISPR-Cas systems. *Nucleic acids research* (2013); Jiang, W. *et al.* RNA-guided editing of bacterial genomes using CRISPR-Cas systems. *Nature biotechnology* **31**, 233-239 (2013); the entire contents of each of which are incorporated herein by reference).

[00167] The term “subject,” as used herein, refers to an individual organism, for example, an individual mammal. In some embodiments, the subject is a human. In some embodiments, the subject is a non-human mammal. In some embodiments, the subject is a non-human primate. In some embodiments, the subject is a rodent. In some embodiments, the subject is a sheep, a goat, a cattle, a cat, or a dog. In some embodiments, the subject is a vertebrate, an amphibian, a reptile, a fish, an insect, a fly, or a nematode. In some

embodiments, the subject is a research animal. In some embodiments, the subject is genetically engineered, *e.g.*, a genetically engineered non-human subject. The subject may be of either sex and at any stage of development.

[00168] The term “target site” refers to a sequence within a nucleic acid molecule that is deaminated by a deaminase or a fusion protein comprising a deaminase, (*e.g.*, a dCas9-deaminase fusion protein provided herein).

[00169] The terms “treatment,” “treat,” and “treating,” refer to a clinical intervention aimed to reverse, alleviate, delay the onset of, or inhibit the progress of a disease or disorder, or one or more symptoms thereof, as described herein. As used herein, the terms “treatment,” “treat,” and “treating” refer to a clinical intervention aimed to reverse, alleviate, delay the onset of, or inhibit the progress of a disease or disorder, or one or more symptoms thereof, as described herein. In some embodiments, treatment may be administered after one or more symptoms have developed and/or after a disease has been diagnosed. In other embodiments, treatment may be administered in the absence of symptoms, *e.g.*, to prevent or delay onset of a symptom or inhibit onset or progression of a disease. For example, treatment may be administered to a susceptible individual prior to the onset of symptoms (*e.g.*, in light of a history of symptoms and/or in light of genetic or other susceptibility factors). Treatment may also be continued after symptoms have resolved, for example, to prevent or delay their recurrence.

[00170] The term “recombinant” as used herein in the context of proteins or nucleic acids refers to proteins or nucleic acids that do not occur in nature, but are the product of human engineering. For example, in some embodiments, a recombinant protein or nucleic acid molecule comprises an amino acid or nucleotide sequence that comprises at least one, at least two, at least three, at least four, at least five, at least six, or at least seven mutations as compared to any naturally occurring sequence.

[00171] The term “nucleobase editors (NBEs)” or “base editors (BEs),” as used herein, refers to the Cas9 fusion proteins described herein. In some embodiments, the fusion protein comprises a nuclease-inactive Cas9 (dCas9) fused to a deaminase. In some embodiments, the fusion protein comprises a Cas9 nickase fused to a deaminase. In some embodiments, the fusion protein comprises a nuclease-inactive Cas9 fused to a deaminase and further fused to a UGI domain. In some embodiments, the fusion protein comprises a Cas9 nickase fused to a deaminase and further fused to a UGI domain. In some embodiments, the dCas9 of the fusion protein comprises a D10A and a H840A mutation of SEQ ID NO: 10, or a corresponding mutation in any of SEQ ID NOs: 11-260, which inactivates nuclease activity

of the Cas9 protein. In some embodiments, the fusion protein comprises a D10A mutation and comprises a histidine at residue 840 of SEQ ID NO: 10, or a corresponding mutation in any of SEQ ID NOs: 11-260, which renders Cas9 capable of cleaving only one strand of a nucleic acid duplex. An example of a Cas9 nickase is shown below in SEQ ID NO: 674. The terms “nucleobase editors (NBEs)” and “base editors (BEs)” may be used interchangeably.

[00172] The term “uracil glycosylase inhibitor” or “UGI,” as used herein, refers to a protein that is capable of inhibiting a uracil-DNA glycosylase base-excision repair enzyme.

[00173] The term “Cas9 nickase,” as used herein, refers to a Cas9 protein that is capable of cleaving only one strand of a duplexed nucleic acid molecule (*e.g.*, a duplexed DNA molecule). In some embodiments, a Cas9 nickase comprises a D10A mutation and has a histidine at position H840 of SEQ ID NO: 10, or a corresponding mutation in any of SEQ ID NOs: 11-260. For example, a Cas9 nickase may comprise the amino acid sequence as set forth in SEQ ID NO: 674. Such a Cas9 nickase has an active HNH nuclease domain and is able to cleave the non-targeted strand of DNA, *i.e.*, the strand bound by the gRNA. Further, such a Cas9 nickase has an inactive RuvC nuclease domain and is not able to cleave the targeted strand of the DNA, *i.e.*, the strand where base editing is desired.

[00174] Exemplary Cas9 nickase (Cloning vector pPlatTET-gRNA2; Accession No. BAV54124).

MDKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLIGALLFDSGE
TAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRLEESFLVEEDKKHE
RHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEG
DLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLP
GEKKNGLFGNLIASLGLTPNFKSNFDLAEDAQLSKDLYDDDLNLLAQIGDQYA
DLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPE
KYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQR
TFDNGSIPHQIHLGELHAILRRQEDFYPFKDNREKIEKILTFRIPYYVGPLARGNSRFA
WMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPHSLLYFYFTV
YNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFD
SVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERL
KTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRN
FMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIAKKGILQTVKVVDELVK
VMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIEIEGKELGSQILKEHPVENTQL
QNEKLYLYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDK

NRGKSDNVPSEEVVKMKNYWRQLLNAKLITQRKFDNLTKAERGGELSELDKAGFIK
 RQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKV
 REINNYHHAHDAYLNAVVGTAIIKKYPKLESEFVYGDYKVYDVRKMIKSEQEIGK
 ATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLMS
 PQVNIKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVV
 AKVEKGKSKKLKSVKELLGITIMERSSSFENPIDFLEAKGYKEVKKDLIIKLPKYSLFE
 LENGKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQ
 HKHYLDEIIEQISEFSKRVLADANLDKVLSAYNKHRDKPIREQAENIIHLFTLTNLGA
 PAAFKYFDTTIDRKRYTSTKEVLDATLIHQSTGLYETRIDLSQLGGD (SEQ ID NO:
 674)

DETAILED DESCRIPTION OF CERTAIN EMBODIMENTS OF THE INVENTION

[00175] Some aspects of this disclosure provide fusion proteins that comprise a domain capable of binding to a nucleotide sequence (*e.g.*, a Cas9, or a Cpf1 protein) and an enzyme domain, for example, a DNA-editing domain, such as, *e.g.*, a deaminase domain. The deamination of a nucleobase by a deaminase can lead to a point mutation at the respective residue, which is referred to herein as nucleic acid editing. Fusion proteins comprising a Cas9 variant or domain and a DNA editing domain can thus be used for the targeted editing of nucleic acid sequences. Such fusion proteins are useful for targeted editing of DNA *in vitro*, *e.g.*, for the generation of mutant cells or animals; for the introduction of targeted mutations, *e.g.*, for the correction of genetic defects in cells *ex vivo*, *e.g.*, in cells obtained from a subject that are subsequently re-introduced into the same or another subject; and for the introduction of targeted mutations, *e.g.*, the correction of genetic defects or the introduction of deactivating mutations in disease-associated genes in a subject. Typically, the Cas9 domain of the fusion proteins described herein does not have any nuclease activity but instead is a Cas9 fragment or a dCas9 protein or domain. Methods for the use of Cas9 fusion proteins as described herein are also provided.

Cas9 domains of Nucleobase Editors

[00176] Non-limiting, exemplary Cas9 domains are provided herein. The Cas9 domain may be a nuclease active Cas9 domain, a nuclease inactive Cas9 domain, or a Cas9 nickase. In some embodiments, the Cas9 domain is a nuclease active domain. For example, the Cas9 domain may be a Cas9 domain that cuts both strands of a duplexed nucleic acid (*e.g.*, both strands of a duplexed DNA molecule). In some embodiments, the Cas9 domain comprises

any one of the amino acid sequences as set forth in SEQ ID NOs: 10-263. In some embodiments the Cas9 domain comprises an amino acid sequence that is at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of the amino acid sequences set forth in SEQ ID NOs: 10-263. In some embodiments, the Cas9 domain comprises an amino acid sequence that has 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50 or more or more mutations compared to any one of the amino acid sequences set forth in SEQ ID NOs: 10-263. In some embodiments, the Cas9 domain comprises an amino acid sequence that has at least 10, at least 15, at least 20, at least 30, at least 40, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 300, at least 350, at least 400, at least 500, at least 600, at least 700, at least 800, at least 900, at least 1000, at least 1100, or at least 1200 identical contiguous amino acid residues as compared to any one of the amino acid sequences set forth in SEQ ID NOs: 10-263.

[00177] In some embodiments, the Cas9 domain is a nuclease-inactive Cas9 domain (dCas9). For example, the dCas9 domain may bind to a duplexed nucleic acid molecule (*e.g.*, via a gRNA molecule) without cleaving either strand of the duplexed nucleic acid molecule. In some embodiments, the nuclease-inactive dCas9 domain comprises a D10X mutation and a H840X mutation of the amino acid sequence set forth in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid change. In some embodiments, the nuclease-inactive dCas9 domain comprises a D10A mutation and a H840A mutation of the amino acid sequence set forth in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. As one example, a nuclease-inactive Cas9 domain comprises the amino acid sequence set forth in SEQ ID NO: 263 (Cloning vector pPlatTET-gRNA2, Accession No. BAV54124).

MDKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLIGALLFDSGE
TAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFLVEEDKKHE
RHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEG
DLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLP
GEKKNGLFGNLIASLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLNLLAQIGDQYA
DLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPE

KYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQR
TFDNGSIPHQIHLGELHAILRRQEDFYPLKDNREKIEKILTFRIPYYVGPLARGNSRFA
WMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPKHSLLEYEYFTV
YNELTKVKYVTEGMRKPAFLSSEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFD
SVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERL
KTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRN
FMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIAKKGILQTVKVVDELVK
VMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGKELGSQILKEHPVENTQL
QNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDAIVPQSFLKDDSIDNKVLTRSDK
NRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDKAGFIK
RQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKV
REINNYHHAHDAYLNAVVGTAIIKKYPKLESEFVYGDYKVYDVRKMIASEQEIGK
ATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLMS
PQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVV
AKVEKKGKSKKLKSVKELLGITIMERSSEKPNIDFLEAKGYKEVKKDLIIKLPKYSLFE
LENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQ
HKHYLDEIIEQISEFSKRVLADANLDKVLSAYNKHRDKPIREQAENIIHLFTLTNLGA
PAAFKYFDTTIDRKRYTSTKEVLDATLIHQSIITGLYETRIDLSQLGGD (SEQ ID NO:

263; see, *e.g.*, Qi *et al.*, Repurposing CRISPR as an RNA-guided platform for sequence-specific control of gene expression. *Cell*. 2013; 152(5):1173-83, the entire contents of which are incorporated herein by reference).

[00178] Additional suitable nuclease-inactive dCas9 domains will be apparent to those of skill in the art based on this disclosure and knowledge in the field, and are within the scope of this disclosure. Such additional exemplary suitable nuclease-inactive Cas9 domains include, but are not limited to, D10A/H840A, D10A/D839A/H840A, and D10A/D839A/H840A/N863A mutant domains (See, *e.g.*, Prashant *et al.*, CAS9 transcriptional activators for target specificity screening and paired nickases for cooperative genome engineering. *Nature Biotechnology*. 2013; 31(9): 833-838, the entire contents of which are incorporated herein by reference). In some embodiments the dCas9 domain comprises an amino acid sequence that is at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of the dCas9 domains provided herein. In some embodiments, the Cas9 domain comprises an amino acid sequences that has 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 21, 24, 25, 26, 27, 28,

29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50 or more or more mutations compared to any one of the amino acid sequences set forth in SEQ ID NOs: 10-263. In some embodiments, the Cas9 domain comprises an amino acid sequence that has at least 10, at least 15, at least 20, at least 30, at least 40, at least 50, at least 60, at least 70, at least 80, at least 90, at least 100, at least 150, at least 200, at least 250, at least 300, at least 350, at least 400, at least 500, at least 600, at least 700, at least 800, at least 900, at least 1000, at least 1100, or at least 1200 identical contiguous amino acid residues as compared to any one of the amino acid sequences set forth in SEQ ID NOs: 10-263.

[00179] In some embodiments, the Cas9 domain is a Cas9 nickase. The Cas9 nickase may be a Cas9 protein that is capable of cleaving only one strand of a duplexed nucleic acid molecule (*e.g.*, a duplexed DNA molecule). In some embodiments the Cas9 nickase cleaves the target strand of a duplexed nucleic acid molecule, meaning that the Cas9 nickase cleaves the strand that is base paired to (complementary to) a gRNA (*e.g.*, an sgRNA) that is bound to the Cas9. In some embodiments, a Cas9 nickase comprises a D10A mutation and has a histidine at position 840 of SEQ ID NO: 10, or a mutation in any of SEQ ID NOs: 11-260. For example, a Cas9 nickase may comprise the amino acid sequence as set forth in SEQ ID NO: 674. In some embodiments the Cas9 nickase cleaves the non-target, non-base-edited strand of a duplexed nucleic acid molecule, meaning that the Cas9 nickase cleaves the strand that is not base paired to a gRNA (*e.g.*, an sgRNA) that is bound to the Cas9. In some embodiments, a Cas9 nickase comprises an H840A mutation and has an aspartic acid residue at position 10 of SEQ ID NO: 10, or a corresponding mutation in any of SEQ ID NOs: 11-260. In some embodiments the Cas9 nickase comprises an amino acid sequence that is at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of the Cas9 nickases provided herein. Additional suitable Cas9 nickases will be apparent to those of skill in the art based on this disclosure and knowledge in the field, and are within the scope of this disclosure.

Cas9 Domains with Reduced PAM Exclusivity

[00180] Some aspects of the disclosure provide Cas9 domains that have different PAM specificities. Typically, Cas9 proteins, such as Cas9 from *S. pyogenes* (spCas9), require a canonical NGG PAM sequence to bind a particular nucleic acid region. This may limit the ability to edit desired bases within a genome. In some embodiments, the base editing fusion

proteins provided herein may need to be placed at a precise location, for example where a target base is placed within a 4 base region (*e.g.*, a “deamination window”), which is approximately 15 bases upstream of the PAM. See Komor, A.C., *et al.*, “Programmable editing of a target base in genomic DNA without double-stranded DNA cleavage” *Nature* 533, 420-424 (2016), the entire contents of which are hereby incorporated by reference. Accordingly, in some embodiments, any of the fusion proteins provided herein may contain a Cas9 domain that is capable of binding a nucleotide sequence that does not contain a canonical (*e.g.*, NGG) PAM sequence. Cas9 domains that bind to non-canonical PAM sequences have been described in the art and would be apparent to the skilled artisan. For example, Cas9 domains that bind non-canonical PAM sequences have been described in Kleinstiver, B. P., *et al.*, “Engineered CRISPR-Cas9 nucleases with altered PAM specificities” *Nature* 523, 481-485 (2015); and Kleinstiver, B. P., *et al.*, “Broadening the targeting range of *Staphylococcus aureus* CRISPR-Cas9 by modifying PAM recognition” *Nature Biotechnology* 33, 1293-1298 (2015); the entire contents of each are hereby incorporated by reference.

[00181] In some embodiments, the Cas9 domain is a Cas9 domain from *Staphylococcus aureus* (SaCas9). In some embodiments, the SaCas9 domain is a nuclease active SaCas9, a nuclease inactive SaCas9 (SaCas9d), or a SaCas9 nickase (SaCas9n). In some embodiments, the SaCas9 comprises the amino acid sequence SEQ ID NO: 4273. In some embodiments, the SaCas9 comprises a N579X mutation of SEQ ID NO: 4273, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for N. In some embodiments, the SaCas9 comprises a N579A mutation of SEQ ID NO: 4273, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the SaCas9 domain, the SaCas9d domain, or the SaCas9n domain can bind to a nucleic acid sequence having a non-canonical PAM. In some embodiments, the SaCas9 domain, the SaCas9d domain, or the SaCas9n domain can bind to a nucleic acid sequence having a NNGRRT PAM sequence. In some embodiments, the SaCas9 domain comprises one or more of a E781X, a N967X, and a R1014X mutation of SEQ ID NO: 4273, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid. In some embodiments, the SaCas9 domain comprises one or more of a E781K, a N967K, and a R1014H mutation of SEQ ID NO: 4273, or one or more corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the SaCas9 domain comprises a

E781K, a N967K, or a R1014H mutation of SEQ ID NO: 4273, or corresponding mutations in any of the amino acid sequences provided in SEQ ID NOs: 11-260.

[00182] In some embodiments, the Cas9 domain of any of the fusion proteins provided herein comprises an amino acid sequence that is at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of SEQ ID NOs: 4273-4275. In some embodiments, the Cas9 domain of any of the fusion proteins provided herein comprises the amino acid sequence of any one of SEQ ID NOs: 4273-4275. In some embodiments, the Cas9 domain of any of the fusion proteins provided herein consists of the amino acid sequence of any one of SEQ ID NOs: 4273-4275.

Exemplary SaCas9 sequence

KRNYILGLDIGITSVGYGIIDYETRDVIDAGVRLFKEANVENNEGRRSKRGARRLKRR
 RRHRIQRVKKLLFDYNLLTDHSELGINPYEARVKGLSQKLSEEFSAALLHLAKRRG
 VHVNEVEEDTGNELSTKEQISRNSKALEEKYVAELQLERLKKDGEVRGSINRFKTS
 DYVKEAKQLLKVQKAYHQLDQSFIDTYIDLETRRYYEGPGEGSPFGWKDIKEWY
 EMLMGHCTYFPEELRSVKYAYNADLYNALNDLNNLVITRDENEKLEYEKFQIIENV
 FKQKKKPTLKQIAKEILVNEEDIKGYRVTSTGKPEFTNLKVYHDIKDITARKEIENAE
 LLDQIAKILTIYQSSEDIQEELTNLNSELTQEEIEQISNLKGYTGTHNLCLKAINLILDEL
 WHTNDNQIAIFNRLKLVPKKVDLSQQKEIPTTLVDDFILSPVVKRSFIQSIKVINAIKK
 YGLPNDIIIELAREKNSKDAQKMINEMQKRNRQTNERIEEIIIRTTGKENAKYLIEKIKL
 HDMQEGKCLYSLEAIPLEDLLNPNFYEVVDHIIPRSVSFDNSFNKVLVKQEEN**SK**KG
 NRTPFQYLSSSDSKISYETFKKHILNLAKGKGRISKTKKEYLLEERDINRFVQKDFIN
 RNLVDTRYATRGLMNLRSYFRVNNLDVKVKSINGGFTSFLRRKWKFKKERNKGY
 KHHAEDALIANADFIFKEWKKLDKAKKVMENQMFEKQAESMPEIETEQEYKEIFIT
 PHQIKHIKDFKDYKYSHRVDKKNRELINDTLYSTRKDDKGNTLIVNNLNGLYDKDN
 DKLKKLINKSPEKLLMYHHPQTYQKLKLIMEQYGDEKNPLYKYEEETGNLYLTKYS
 KKDNGPVIKKIKYYGNKLNHLAHLDDYDYPNSRNKVVKLSLKPFRFDVYLDNGVYKF
 VTVKNLDVIKKENYYEVNSKCYEEAKKLLKISNQAEFIASFYNNDLIKINGELYRVIG
 VNNDLLNRIEVNMIDITYREYLENMNDKRPPRIIKTIASKTQSIKKYSTDILGNLYEVK
 SKKHPQIIKKG (SEQ ID NO: 4273)

Residue N579 of SEQ ID NO: 4273, which is underlined and in bold, may be mutated (*e.g.*, to a A579) to yield a SaCas9 nickase.

Exemplary SaCas9n sequence

KRNYILGLDIGITSVGYGIIDYETRDVIDAGVRLFKEANVENNEGRRSKRGARRLKRR
 RRHRIQRVKKLLFDYNLLTDHSELSGINPYEARVKGLSQKLSEEEFSAALLHLAKRRG
 VHNVNEVEEDTGNELSTKEQISRNSKALEEKYVAELQLERLKKDGEVRGSINRFKTS
 DYVKEAKQLLKVQKAYHQLDQSFIDTYIDLLETRRITYYEGPGEGSPFGWKDIKEWY
 EMLMGHCTYFPEELRSVKYAYNADLYNALNDLNNLVITRDENEKLEYEKFQIIENV
 FKQKKKPTLKQIAKEILVNEEDIKGYRVTSTGKPEFTNLKVYHDIKDITARKEIENAE
 LLDQIAKILTIYQSSEDIQEELTNLNSELTQEEIEQISNLKGYTGTHNLSLKAINLILDEL
 WHTNDNQIAIFNRLKLVPKKVDLSQQKEIPTTLVDDFILSPVVKRSFIQSIKVINAIKK
 YGLPNDIIIELAREKNSKDAQKMINEMQKRNRQTNERIEEIIIRTTGKENAKYLIEKIKL
 HDMQEGKCLYSLEAIPLEDLLNPNFNYEVDHIIPRSVSFDNSFNKVLVKQEE**A**SKKG
 NRTPFQYLSSSDSKISYETFKKHILNLAGKGRISKTKKEYLLEERDINRFVQKDFIN
 RNLVDTRYATRGLMNLRSYFRVNNLDVKVKSINGGFTSFLRRKWKFKKERNKGY
 KHHAEDALIIANADFIFKEWKKLDKAKKVMENQMFEKQAESMPEIETEQEYKEIFIT
 PHQIKHIKDFKDYKYSHRVDKKPNRELINDTLYSTRKDDKGNTLIVNNLNGLYDKDN
 DKLKLINKSPEKLLMYHHPQTYQKLKLIMEQYGDEKNPLYKYEEETGNLYTKYS
 KKDNGPVIKKIKYYGNKLNHLADITDDYPNSRNKVVKLSLKPYRFDVYLDNGVYKF
 VTVKNLDVIKKENYYEVNSKCYEEAKKLLKISNQAEFIASFYNNDLIKINGELYRVIG
 VNNDLLNRIEVNMIDITYREYLENMNDKRPPRIIKTIASKTQSIKKYSTDILGNLYEVK
 SKKHPQIIKKG (SEQ ID NO: 4274).

Residue A579 of SEQ ID NO: xx, which can be mutated from N579 of SEQ ID NO: 4274 to yield a SaCas9 nickase, is underlined and in bold.

Exemplary SaKKH Cas9

KRNYILGLDIGITSVGYGIIDYETRDVIDAGVRLFKEANVENNEGRRSKRGARRLKRR
 RRHRIQRVKKLLFDYNLLTDHSELSGINPYEARVKGLSQKLSEEEFSAALLHLAKRRG
 VHNVNEVEEDTGNELSTKEQISRNSKALEEKYVAELQLERLKKDGEVRGSINRFKTS
 DYVKEAKQLLKVQKAYHQLDQSFIDTYIDLLETRRITYYEGPGEGSPFGWKDIKEWY
 EMLMGHCTYFPEELRSVKYAYNADLYNALNDLNNLVITRDENEKLEYEKFQIIENV
 FKQKKKPTLKQIAKEILVNEEDIKGYRVTSTGKPEFTNLKVYHDIKDITARKEIENAE
 LLDQIAKILTIYQSSEDIQEELTNLNSELTQEEIEQISNLKGYTGTHNLSLKAINLILDEL
 WHTNDNQIAIFNRLKLVPKKVDLSQQKEIPTTLVDDFILSPVVKRSFIQSIKVINAIKK
 YGLPNDIIIELAREKNSKDAQKMINEMQKRNRQTNERIEEIIIRTTGKENAKYLIEKIKL
 HDMQEGKCLYSLEAIPLEDLLNPNFNYEVDHIIPRSVSFDNSFNKVLVKQEE**A**SKKG

NRTPFQYLSSSDSKISYETFKKHILNLA KGKGRISKTKKEYLLEERDINRFSVQKDFIN
 RNLVDTRYATRGLMNLRSYFRVNNLDVKVKSINGGFTSFLRRKWKFKKERNKGY
 KHHAEDALIIANADFIFKEWKKLDKAKKVMENQMFE EKQAESMPEIETE QEYKEIFIT
 PHQIKHIKDFKDYKYSHRVDKKPNR**K**LINDTLYSTRKDDKGNTLIVNNLNGLYDKD
 NDKLLKLINKSPEKLLMYHHDPQTYQKLKLIMEQYGDEKNPLYKYEEETGNYLTKY
 SKKDNGPVIKKIKYYGNKLNAHLDITDDYPNSRNKVVKLSLKPYPYRFDVYLDNGVYK
 FVTVKNLDVIKKENYYEVNSKCYEEAKKLKISNQAEFIASFY**K**NDLIKINGELYRVI
 GVNNDLLNRIEVNMIDITYREYLENMNDKRPP**H**IKTIASKTQSIKKYSTDILGNLYEV
 KSKKHPQIIKKG (SEQ ID NO: 4275).

Residue A579 of SEQ ID NO: 4275, which can be mutated from N579 of SEQ ID NO: 4275 to yield a SaCas9 nickase, is underlined and in bold. Residues K781, K967, and H1014 of SEQ ID NO: 4275, which can be mutated from E781, N967, and R1014 of SEQ ID NO: 4275 to yield a SaKKH Cas9 are underlined and in italics.

[00183] In some embodiments, the Cas9 domain is a Cas9 domain from *Streptococcus pyogenes* (SpCas9). In some embodiments, the SpCas9 domain is a nuclease active SpCas9, a nuclease inactive SpCas9 (SpCas9d), or a SpCas9 nickase (SpCas9n). In some embodiments, the SpCas9 comprises the amino acid sequence SEQ ID NO: 4276. In some embodiments, the SpCas9 comprises a D9X mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for D. In some embodiments, the SpCas9 comprises a D9A mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the SpCas9 domain, the SpCas9d domain, or the SpCas9n domain can bind to a nucleic acid sequence having a non-canonical PAM. In some embodiments, the SpCas9 domain, the SpCas9d domain, or the SpCas9n domain can bind to a nucleic acid sequence having a NGG, a NGA, or a NGCG PAM sequence. In some embodiments, the SpCas9 domain comprises one or more of a D1134X, a R1334X, and a T1336X mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid. In some embodiments, the SpCas9 domain comprises one or more of a D1134E, R1334Q, and T1336R mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the SpCas9 domain comprises a D1134E, a R1334Q, and a T1336R mutation of SEQ ID NO: 4276, or corresponding mutations in any of the amino acid

sequences provided in SEQ ID NOs: 11-260. In some embodiments, the SpCas9 domain comprises one or more of a D1134X, a R1334X, and a T1336X mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid. In some embodiments, the SpCas9 domain comprises one or more of a D1134V, a R1334Q, and a T1336R mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the SpCas9 domain comprises a D1134V, a R1334Q, and a T1336R mutation of SEQ ID NO: 4276, or corresponding mutations in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the SpCas9 domain comprises one or more of a D1134X, a G1217X, a R1334X, and a T1336X mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid. In some embodiments, the SpCas9 domain comprises one or more of a D1134V, a G1217R, a R1334Q, and a T1336R mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the SpCas9 domain comprises a D1134V, a G1217R, a R1334Q, and a T1336R mutation of SEQ ID NO: 4276, or corresponding mutations in any of the amino acid sequences provided in SEQ ID NOs: 11-260.

[00184] In some embodiments, the Cas9 domain of any of the fusion proteins provided herein comprises an amino acid sequence that is at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of SEQ ID NOs: 4276-4280. In some embodiments, the Cas9 domain of any of the fusion proteins provided herein comprises the amino acid sequence of any one of SEQ ID NOs: 4276-4280. In some embodiments, the Cas9 domain of any of the fusion proteins provided herein consists of the amino acid sequence of any one of SEQ ID NOs: 4276-4280.

Exemplary SpCas9

DKKYSIGLDIGTNSVGVAVITDEYKVPSKKFKVLGNTDRHSIKKNLIGALLFDSGETA
 EATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFLVEEDKKHERH
 PIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADRLIYLALAHMIKFRGHFLIEGDL
 NPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGE
 KKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDQYDDDLNLLAQIGDQYADL
 FLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKY

KEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTF
DNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSRFAW
MTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPHSLLYEYFTVY
NELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDS
VEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERL
KTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRN
FMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIAKKGILQTVKVVDELVK
VMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEIEGKELGSQILKEHPVENTQL
QNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDK
NRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELKAGFIK
RQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKV
REINNYHHAHDAYLNAVVGTAIIKKYPKLESEFVYGDYKVYDVRKMIASEQEIGK
ATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLMS
PQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVV
AKVEKKGKSKKLKSVKELLGITIMERSSEKPNIDFLEAKGYKEVKKDLIIKLPKYSLFE
LENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQ
HKHYLDEIIEQISEFSKRVLADANLDKVLSAYNKHRDKPIREQAENIIHLFTLTNLGA
PAAFKYFDTTIDRKRYTSTKEVLDATLIHQSIITGLYETRIDLSQLGGD (SEQ ID NO:
4276)

Exemplary SpCas9n

DKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIGALLFDSGETA
EATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVVDSFFHRLEESFLVEEDKKHERH
PIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDL
NPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGE
KKNGLFGNLIALSLGLTPNFKSNFDLAEDAQLQSKDQYDDDLNLLAQIGDQYADL
FLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKY
KEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTF
DNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSRFAW
MTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPHSLLYEYFTVY
NELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDS
VEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERL
KTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRN
FMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIAKKGILQTVKVVDELVK

VMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIEELGSELKESQILKEHPVENTQL
 QNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDK
 NRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGELSELDKAGFIK
 RQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKV
 REINNYHHAHDAYLNAVVGTAIIKKYPKLESEFVYGDYKVYDVRKMIKSEQEIGK
 ATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLMS
 PQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVV
 AKVEKGGKSKKLSVKELLGITIMERSSEFEKNPIDFLEAKGYKEVKKDLIIKLPKYSLEF
 LENGKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQ
 HKHYLDEIIEQISEFSKRVILADANLDKVL SAYNKHRDKPIREQAENIIHLFTLTNLGA
 PAAFKYFDTTIDRKRYTSTKEVLDATLIHQSI TGLYETRIDLSQLGGD (SEQ ID NO:
 4277)

Exemplary SpEQR Cas9

DKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIGALLFDSGETA
 EATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVVDSFFHRLEESFLVEEDKKHERH
 PIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDL
 NPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGE
 KKNGLFGNLIALSLGLTPNFKSNFDLAEDAQLSKD TYDDDLNLLAQIGDQYADL
 FLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKY
 KEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTF
 DNGSIPHQIHLGELHAILRRQEDFY PFLKDNREKIEKILTFRIPYYVGPLARGNSRFAW
 MTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPKHSLLYEYFTVY
 NELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDS
 VEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERL
 KTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDFANRN
 FMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIIKKGILQTVKVVDELVK
 VMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIEELGSELKESQILKEHPVENTQL
 QNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDK
 NRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGELSELDKAGFIK
 RQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKV
 REINNYHHAHDAYLNAVVGTAIIKKYPKLESEFVYGDYKVYDVRKMIKSEQEIGK
 ATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLMS
 PQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFESPTVAYSVLVV

AKVEKGKSKKLKSVKELLGITIMERSSSFENPIDFLEAKGYKEVKKDLIIKLPKYSLFE
 LENGKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQ
 HKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIIHLFTLTNLGA
 PAAFKYFDTTIDRKQYRSTKEVLDATLIHQSTGLYETRIDLSQLGGD (SEQ ID NO:
 4278)

Residues E1134, Q1334, and R1336 of SEQ ID NO: 4278, which can be mutated from
 D1134, R1334, and T1336 of SEQ ID NO: 4278 to yield a SpEQR Cas9, are underlined and
 in bold.

Exemplary SpVQR Cas9

DKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLIGALLFDSGETA
 EATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDDSFFHRLEESFLVEEDKKHERH
 PIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADRLIYLALAHMIKFRGHFLIEGDL
 NPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGE
 KKNGLFGNLIALSLGLTPNFKSNFDLAEDAQLQSKDQYDDDLNLLAQIGDQYADL
 FLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKY
 KEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTF
 DNGSIPHQIHLGELHAILRRQEDFYFLKDNREKIEKILTFRIPYYVGPLARGNSRFAW
 MTRKSEETITPWNFEEVVDKGGASAQSFIERMTNFDKNLPNEKVLPHSLLYEYFTVY
 NELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDS
 VEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERL
 KTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRN
 FMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIIKKGILQTVKVVDELVK
 VMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIEIEGKELGSQILKEHPVENTQL
 QNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDK
 NRGKSDNVPSEEVVKMKNYWRQLLNAKLITQRKFDNLTKAERGGSELKAGFIK
 RQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKV
 REINNYHHAHDAYLNAVVGTAIIKYPKLESEFVYGDYKVYDVRKMIKSEQEIGK
 ATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLMS
 PQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFVSPTVAYSVLVV
 AKVEKGKSKKLKSVKELLGITIMERSSSFENPIDFLEAKGYKEVKKDLIIKLPKYSLFE
 LENGKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQ
 HKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIIHLFTLTNLGA

PAAFKYFDTTIDRKQYRSTKEVLDATLIHQSI TGLYETRIDLSQLGGD (SEQ ID NO: 4279)

Residues V1134, Q1334, and R1336 of SEQ ID NO: 4279, which can be mutated from D1134, R1334, and T1336 of SEQ ID NO: 4279 to yield a SpVQR Cas9, are underlined and in bold.

Exemplary SpVRER Cas9

DKKYSIGLAIGTNSVGWAVITDEYKVP SKKFKVLGNTDRHSIKKNLIGALLFDSGETA
EATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDD SFFHRLEESFLVEEDKKHERH
PIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLR LIYLALAHMIKFRGHFLIEGDL
NPDNSDV DKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGE
KKNGLFGNLIALSLGLTPNFKSNFDLAEDA KLQLSKDTYDDDLNLLAQIGDQYADL
FLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKY
KEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTF
DNGSIPHQIHLGELHAILRRQEDFY PFLKDNREKIEKILTFRIPYYVGPLARGNSRFAW
MTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPHSLLYEYFTVY
NELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDS
VEISGVEDRFNASLGT YHDLKIIKDKDFLDNEENEDI EDIVLTLTLFEDREMIEERL
KTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRN
FMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPA IKGILQTVKVVDELVK
VMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIE EGikelGSQILKEHPVENTQL
QNEKLYLYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDK
NRGKSDNVPSEEVVKMKNYWRQLLNAKLITQRKFDNLTKA ERGGLSELDKAGFIK
RQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITL KSKLVSDFRKDFQFYKV
REINNYHHAHDAYLNAVVGTA LIKKYPKLESEFVYGDYKVYDVRKMIAKSEQEIGK
ATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEI VWDKGRDFATVRKVL SM
PQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWD PKKYGGFVSPTVAYSVLVV
AKVEKGKSKKLKSVKELLGITIMERS SFEKNPIDFLEAKGYKEVKKDLIIKLPKYSLFE
LENGRKRMLASARELQKGNELALPSKYVNFLYLASHYEK LKGSPEDNEQKQLFVEQ
HKHYLDEIIEQISEFSKR VILADANLDKVLSAYNKHRDKPIREQAENIIHLFTLTNLGA
PAAFKYFDTTIDRKEYRSTKEVLDATLIHQSI TGLYETRIDLSQLGGD (SEQ ID NO: 4280)

Residues V1134, R1217, Q1334, and R1336 of SEQ ID NO: 4280, which can be mutated from D1134, G1217, R1334, and T1336 of SEQ ID NO: 4280 to yield a SpVRER Cas9, are underlined and in bold.

[00185] The following are exemplary fusion proteins (*e.g.*, base editing proteins) capable of binding to a nucleic acid sequence having a non-canonical (*e.g.*, a non-NGG) PAM sequence:

Exemplary SaBE3 (rAPOBEC1-XTEN-SaCas9n-UGI-NLS)

MSSETGPVAVDPTLRRRIEPHEFEVFFDPREL RKETCLLYEINWGGRHSIWRHTSQNT
 NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRN FVNYSNEAHWPRYPHLW
 VRLYVLELYCII LGLPPCLNLRKQPQLTFFTIALQSCHYQRLPPHILWATGLKSGSET
PGTSESATPESKRNYILGLDIGITSVGYGIIDYETRDVIDAGVRLFKEANVENNEGRRS
KRGARRLKRRRRHRIQRVKLLFDYNLLTDHSELSGINPYEARVKGLSOKLSEEEFS
AALLHLAKRRGVHNVNEVEEDTGNELSTKEQISRNSKALEEKYVAELQLERLKKDG
EVRSINRFKTS DYVKEAKQLLKVOKAYHQLDQSFIDTYIDLLETRRTYYEGPGE GSP
FGWKDIKEWYEMLMGHCTYFPEELRSVKYAYNADLYNALNDLNNLVITRDENEKL
EYYEKFOIENVFKQKKKPTLKQIAKEILVNEEDIKGYRVTSTGKPEFTNLKVYHDIK
DITARKEIENAELLDQIAKILTIYQSSEDIQEELTNLSELTOEEIEQISNLKGYTGTHN
LSLKAINLILDELWHTNDNQIAIFNRLKLVPKKVDLSQOKEIPTTLVDDFILSPVVKRS
FIOSIKVINAIKKYGLPNDIIEELAREKNSKDAQKMINEMQKRNRQTNERIEEIRTTGK
ENAKYLIEKIKLHDMQEGKCLYSLEAIPLEDLLNPNFNYEVDHII PRSVSFDNSFNK V
LVKQEEASKKGNRTPFOYLSSSDSKISYETFKKHILNLA KGKGRISKTKKEYLLEERDI
NRFSVQKDFINRLVDTRYATRGLMNLRSYFRVNNLDVKVKSINGGFTSFLRRKW
KFKKERNKGYKHAEDALIANADFIKWKLDKAKKVMENQMFE EKQAESMPEI
ETEQEYKEIFITPHQIKHIKDFKDYKYSHRVDKKNRELINDTLYSTRKDDKGNTLIV
NNLNGLYDKDNDKLLKLINKSPEKLLMYHHPQTYQKLKLIMEQY GDEKNPLYKY
YEETGNYLTKYSKDN GPVIKKIKYYGNKLN AHLDDITDDYPNSRNKVVKLSLKP YRF
DVYLDNGVYKFTVKNLDVIKKENYEVNSKCYEEAKK LKKISNQAEFIASFYNN D
LIKINGELYRVIGVNNDLNRIEVNMIDITYREYLENMNDKRPPRIIKTIASKTQSIKKY
STDILGNLYEVKSKKHPQIIKKGSGGSTNLSDIEKETGKQLVIQESILMLPEEVEEVIG
NKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSPKK
KRKV (SEQ ID NO: 4281)

Exemplary SaKKH-BE3 (rAPOBEC1-XTEN-SaCas9n-UGI-NLS)

MSSETGPVAVDPTLRRRIEPHEFEVFFDPREL RKETCLLYEINWGGRHSIWRHTSQNT
 NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRN FVNYSNEAHWPRYPHLW
 VRLYVLELYCII LGLPPCLNLRKQPQLTFFTIALQSCHYQRLPPHILWATGLKSGSET
PGTSESATPESKRNYILGLDIGITSVGYGIIDYETRDVIDAGVRLFKEANVENNEGRRS
KRGARRLKRRRRHRIQRVKLLFDYNLLTDHSELSGINPYEARVKGLSOKLSEEEFS
AALLHLAKRRGVHNVNEVEEDTGNELSTKEQISRNSKALEEKYVAELQLERLKKDG
EVRSINRFKTS DYVKEAKQLLKVOKAYHQLDQSFIDTYIDLLETRRTYYEGPGE GSP
FGWKDIKEWYEMLMGHCTYFPEELRSVKYAYNADLYNALNDLNNLVITRDENEKL
EYYEKFOIENVFKQKKKPTLKQIAKEILVNEEDIKGYRVTSTGKPEFTNLKVYHDIK
DITARKEIENAELLDQIAKILTIYQSSEDIQEELTNLSELTOEEIEQISNLKGYTGTHN
LSLKAINLILDELWHTNDNQIAIFNRLKLVPKKVDLSQOKEIPTTLVDDFILSPVVKRS
FIOSIKVINAIKKYGLPNDIIEELAREKNSKDAQKMINEMQKRNRQTNERIEEIRTTGK

ENAKYLIEKIKLHDMQEGKCLYSLEAIPLEDLLNNPFNYEVDHIIPRSVSFDNSFNKVK
LVKQEEASKKGNRTPFOYLSSSDSKISYETFKKHILNLAAGKGRISKTKKEYLLEERDI
NRFVQKDFINRNLVDTRYATRGLMNLRSYFRVNNLDVKVKSINGGFTSFLRRKW
KFKKERNKGYKHHAEDALIANADFIKWKLDKAKKVMENQMFEKQAESMPEI
ETEQEYKEIFITPHQIKHIKDFKDYKYSHRVDKKNRKLINDTL YSTRKDDKGNTLIV
NNLNGLYDKDNDKLLKLINKSPEKLLMYHHPQTYQKLLIMEQYGDENPLYKY
YEETGNYLTKYSKDNNGPVIKKIKYYGNKLNHLADITDDYPNSRNKVVKLSLKPYRF
DVYLDNGVYKFTVKNLDVIKKENYEVNSKCYEEAKKLLKISNOAEFIASFYKND
LIKINGELYRVIGVNNDLLNRIEVNMIDITYREYLENMNDKRPPHIIKTIASKTQSIKKY
STDILGNLYEVKSKKHPQIIKKGSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIG
NKPESDILVHTAYDESTDENVMMLTSDAPEYKPWALVIQDSNGENKIKMLSGGSPKK
KRKV (SEQ ID NO: 4282)

Exemplary EQR-BE3 (rAPOBEC1-XTEN-Cas9n-UGI-NLS)

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRHSIWRHTSQNT
NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNRFVNYSPSNEAHWPRYPHLW
VRLYVLELYCIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLKSGSET
PGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDRHSIKKNI
GALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFL
VEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIK
FRGHFLIEGDLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSR
LENLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDLAEDAQLQSKDITYDDDLN
LAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLK
ALVRQOLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLN
REDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGP
LARGNSRFAWMTRKSEETITPWNFEEVVDKGGASAQSFIERMTNFDKNLPNEKVLPK
HSLLEYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKE
DYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLF
EDREMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLLINGIRDKQSGKTILDFL
KSDGFANRNFMLIHDDSLTFKEDIQKAQVSGQGDLSLHEHIANLAGSPAIIKKGILQTV
KVVDELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIIEGKELGSQILKE
HPVENTQLQNEKLYLYLQNGRDMYVDQELDINRLSDYVDHIVPOSFLKDDSIDN
KVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLS
ELDKAGFIKROLVETROITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFR
KDFQFYK VREINNYHHAHDAYLNAVVGTAALIKKYPKLESEFVYGDYKVVYDVRKMI
AKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFA
TVRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKGFFESPT
VAYSVLVVAKVEKGKSKKLSVKELLGITIMERSSEFEKNPIDFLEAKGYKEVKKDLII
KLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLLKGSPELN
EQKQLFVEQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIIH
LFTLTNLGAPAAFYFDTTIDRKQYRSTKEVLDTLIHQ SITGLYETRIDLSQLGGDSG
GSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMML
TSDAPEYKPWALVIQDSNGENKIKMLSGGSPKKRKV (SEQ ID NO: 4283)

VQR-BE3 (rAPOBEC1-XTEN-Cas9n-UGI-NLS)

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRHSIWRHTSQNT
NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNRFVNYSPSNEAHWPRYPHLW
VRLYVLELYCIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLKSGSET

PGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLI
GALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFL
VEEDKKHERHPIFGNIVDEVAHYEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIK
FRGHFLIEGDLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRR
LENLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDLAEDAQLQSKDITYDDDLNLI
LAQIGDOYADLFLAAKNLSAAILSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLK
ALVRQOLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLN
REDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGP
LARGNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPK
HSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKE
DYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLF
EDREMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFL
KSDGFANRNFMLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIIKKGILQTV
KVVDDELVKVMGRHKPENIVIAMARENQTTQKGOKNSRERMKRIIEGIKELGSQILKE
HPVENTQLQNEKLYLYLQNGRDMYVDQELDINRLSDYVDHIVPOSFLKDDSIDN
KVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLS
ELDKAGFIKROLVETROITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFR
KDFQFYK VREINNYHHAHDAYLNAVVGTAIIKYPKLESEFVYGDYK VYDVRKMI
AKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRLIETNGETGEIVWDKGRDFA
TVRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGF/SPT
VAYSVLVVAKEKVKSKKLKSVKELLGITIMERSSEKPNIDFLEAKGYKEVKKDLII
KLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLLASHYEKLLKGSPEDN
EQKQLFVEQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIHH
LFTLTNLGAPAAFKYFDTTIDRKQYRSTKEVL DATLIHQ SITGLYETRIDLSQLGGDSG
GSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMML
TSDAPEYKPWALVIQDSNGENKIKMLSGGSPKKKRV (SEQ ID NO: 4284)

VRER-BE3 (rAPOBEC1-XTEN-Cas9n-UGI-NLS)

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCLLYEINWGGRHSIWRHTSQNT
NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNPSNEAHWPRYPHLW
VRLYVLELYCII LGLPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLKSGSET
PGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLI
GALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFL
VEEDKKHERHPIFGNIVDEVAHYEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIK
FRGHFLIEGDLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRR
LENLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDLAEDAQLQSKDITYDDDLNLI
LAQIGDOYADLFLAAKNLSAAILSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLK
ALVRQOLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLN
REDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGP
LARGNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPK
HSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKE
DYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLF
EDREMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFL
KSDGFANRNFMLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIIKKGILQTV
KVVDDELVKVMGRHKPENIVIAMARENQTTQKGOKNSRERMKRIIEGIKELGSQILKE
HPVENTQLQNEKLYLYLQNGRDMYVDQELDINRLSDYVDHIVPOSFLKDDSIDN
KVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLS
ELDKAGFIKROLVETROITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFR
KDFQFYK VREINNYHHAHDAYLNAVVGTAIIKYPKLESEFVYGDYK VYDVRKMI

AKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFA
TVRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKGFF/SPT
VAYSVLVVAKVEKGGKSKKLKSVKELLGITIMERSSFENPIDFLEAKGYKEVKKDLII
KLPKYSLFELNGRKRMLASARELQKGNELALPSKYVNFLYLASHYEKLGKSPEDNE
QKQLFVEQHKHYLDEIIEQISEFSKRVILADANLDKVL SAYNKHRDKPIREQAENIIHL
FTLTNLGAPAAFKYFDTTIDRKE~~Y~~RSTKEVLDA~~T~~LHQSITGLYETRIDLSQLGGDSGG
STNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTS
DAPEYKPWALVIQDSNGENKIKMLSGGSPKKR~~K~~V (SEQ ID NO: 4285)

High Fidelity Base Editors

[00186] Some aspects of the disclosure provide Cas9 fusion proteins (*e.g.*, any of the fusion proteins provided herein) comprising a Cas9 domain that has high fidelity. Additional aspects of the disclosure provide Cas9 fusion proteins (*e.g.*, any of the fusion proteins provided herein) comprising a Cas9 domain with decreased electrostatic interactions between the Cas9 domain and a sugar-phosphate backbone of a DNA, as compared to a wild-type Cas9 domain. In some embodiments, a Cas9 domain (*e.g.*, a wild type Cas9 domain) comprises one or more mutations that decreases the association between the Cas9 domain and a sugar-phosphate backbone of a DNA. In some embodiments, any of the Cas9 fusion proteins provided herein comprise one or more of a N497X, a R661X, a Q695X, and/or a Q926X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid. In some embodiments, any of the Cas9 fusion proteins provided herein comprise one or more of a N497A, a R661A, a Q695A, and/or a Q926A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the Cas9 domain comprises a D10A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the Cas9 domain (*e.g.*, of any of the fusion proteins provided herein) comprises the amino acid sequence as set forth in SEQ ID NO: 325. In some embodiments, the fusion protein comprises the amino acid sequence as set forth in SEQ ID NO: 285. Cas9 domains with high fidelity are known in the art and would be apparent to the skilled artisan. For example, Cas9 domains with high fidelity have been described in Kleinstiver, B.P., *et al.* “High-fidelity CRISPR-Cas9 nucleases with no detectable genome-wide off-target effects.” *Nature* 529, 490-495 (2016); and Slaymaker, I.M., *et al.* “Rationally engineered Cas9 nucleases with improved specificity.” *Science* 351, 84-88 (2015); the entire contents of each are incorporated herein by reference.

[00187] It should be appreciated that the base editors provided herein, for example base editor 2 (BE2) or base editor 3 (BE3), may be converted into high fidelity base editors by modifyint the Cas9 domain as described herein to generate high fidelity base editors, for example high fidelity base editor 2 (HF-BE2) or high fidelity base editor 3 (HF-BE3). In some embodiments, base editor 2 (BE2) comprises a deaminase domain, a dCas9, and a UGI domain. In some embodiments, base editor 3 (BE3) comprises a deaminase domain an nCas9 domain and a UGI domain.

Cas9 domain where mutations relative to Cas9 of SEQ ID NO: 10 are shown in bold and underlines

DKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLIGALLFDSGETAEATRLKRTARR
 RYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRK
 KLVSTDKADLRLIYLALAHMIKFRGHFLIEGDLNPDNSVDKLFQQLVQTYNQLFEENPINASGVDK
 AILSARLSKSRLENLIAQLPGEKKNLFGNLIASLGLTPNFKSNFDLAEDAKLQLSKDYYDDLDNLL
 AQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKYK
 EIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTFDNGSIPHQIHLGE
 LHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSRFAWMTRKSEETITPWNFEEVVDKGASA
 QSFIERMTAFDKNLPNEKVLPHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTN
 RKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLE
 DREMIEERLKTYAHLFDDKVMKQLKRRRYTGWGALSRKLINGIRDKQSGKTILDFLKSDFANRNF
ALIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGILQTVKVVDELVKVMGRHKPENIVIE
 ARENQTTQKGQNSRERMKRIEIGKELGSQILKEHPVENTQLQNEKLYLYYLQNGRDMYVDQELDIN
 RLSDYDVDHIVPQSFLKDDSIDNKVLTRSDKNRGSNDVPSEEVVKKMKNYWRQLLNAKLITQRKFD
 NLTKAERGGLSELKAGFIKRQLVETRAITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFR
 KDFQFYKREINNYHHAHDAYLNAVVGTAIIKKYPKLESEFVYGDYKVVYDVRKMIKSEQEIGKATA
 KYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLVSMQVNVKKTVEVQTG
 GFSKESILPKRNSDKLIARKKDWDPKKGFFSPTVAYSVLVVAKVEKGKSKKLKSVKELLGITIMERS
 SFEKNPIDFLEAKGYKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASH
 YEKLGSPEDNEQKQLFVEQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIIH
 LFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQISITGLYETRIDLSQLGGD (SEQ ID NO: 325)

HF-BE3

MSETGPVAVDPTLRRRIEPHEFEVFFDPRELRKETCLLYEINWGRHSIWRHTSQNTNKHVEVNFIEKF
 TTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYHHADPRNRQGLRDLISSGVTI
 QIMTEQESGYCWRNFVNYSNEAHWPRYPHLWVRLYVLELYCIILGLPPCLNILRRKQPQLTFFTIALQ
 SCHYQRLPPHILWATGLKSGSETPGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLG
 NDRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFL
 VEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVSTDKADLRLIYLALAHMIKFRGHFLIEGDLN

PDNSDVDKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRRLLENLIAQLPGEKKNGLFGNLIALS
 LGLTPNFKSNFDLAEDAQLQSKDQYDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITK
 APLSASMIKRYDEHHQDLTLLKALVRQQLPKEYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKM
 DGTEELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGP
 LARGNSRFAWMTRKSEETITPWNFEEVVDKGASQAQSFIERMTAFDKNLPNEKVLPHKSLLYEYFTVYN
 ELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASL
 GTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERLKYAHLFDDKVMKQLKRRRYTGWG
 ALSRKLINGIRDKQSGKTILDFLKSDGFANRNFMALIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGS
 PAIKKGILQTVKVVDELVKVMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIEEGKELGSQILKEH
 PVENTQLQNEKLYLYLQNGRDMYVDQELDINRLSDYVDHIVPQSFLKDDSIDNKVLTRSDKNRGRKS
 DNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDKAGFIKRQLVETRAITKHVAQIL
 DSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKREINNYHHAHDAYLNAVVGTAIHKYKPK
 LESEFVYGDYKVDVRKMIKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIV
 WDKGRDFATVRKVLSPQVNIKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKGFFSPTVA
 YSVLVVAKVEKGGSKKLSVKELLGITIMERSSEKFNPIDFLEAKGYKEVKKDLIHKLPKYSLFELENGR
 KRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQQLFVEQHKHYLDEIIEQISEFSKR
 VILADANLDKVL SAYNKHDKPIREQAENIIHLFTLNLGAPAAFYFDTTIDRKRYTSTKEVLDATLIH
 QSITGLYETRIDLSQLGGD (SEQ ID NO: 285)

Cas9 fusion proteins

[00188] Any of the Cas9 domains (e.g., a nuclease active Cas9 protein, a nuclease-inactive dCas9 protein, or a Cas9 nickase protein) disclosed herein may be fused to a second protein, thus fusion proteins provided herein comprise a Cas9 domain as provided herein and a second protein, or a “fusion partner”. In some embodiments, the second protein is fused to the N-terminus of the Cas9 domain. However, in other embodiments, the second protein is fused to the C-terminus of the Cas9 domain. In some embodiments, the second protein that is fused to the Cas9 domain is a nucleic acid editing domain. In some embodiments, the Cas9 domain and the nucleic acid editing domain are fused via a linker, while in other embodiments the Cas9 domain and the nucleic acid editing domain are fused directly to one another. In some embodiments, the linker comprises (GGGS)_n (SEQ ID NO: 265), (GGGGS)_n (SEQ ID NO: 5), (G)_n, (EAAAK)_n (SEQ ID NO: 6), (GGS)_n, (SGGS)_n (SEQ ID NO: 4288), SGSETPGTSESATPES (SEQ ID NO: 7), or (XP)_n motif, or a combination of any of these, wherein n is independently an integer between 1 and 30, and wherein X is any amino acid. In some embodiments, the linker comprises a (GGS)_n motif, wherein n is 1, 3, or 7. In some embodiments, the linker comprises a (GGS)_n motif, wherein n is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, or 15. In some embodiments, the linker comprises an amino acid sequence of

SGSETPGTSESATPES (SEQ ID NO: 7), also referred to as the XTEN linker in the Examples). The length of the linker can influence the base to be edited, as illustrated in the Examples. For example, a linker of 3-amino-acid long (*e.g.*, (GGS)₁) may give a 2-5, 2-4, 2-3, 3-4 base editing window relative to the PAM sequence, while a 9-amino-acid linker (*e.g.*, (GGS)₃ (SEQ ID NO: 596)) may give a 2-6, 2-5, 2-4, 2-3, 3-6, 3-5, 3-4, 4-6, 4-5, 5-6 base editing window relative to the PAM sequence. A 16-amino-acid linker (*e.g.*, the XTEN linker) may give a 2-7, 2-6, 2-5, 2-4, 2-3, 3-7, 3-6, 3-5, 3-4, 4-7, 4-6, 4-5, 5-7, 5-6, 6-7 base window relative to the PAM sequence with exceptionally strong activity, and a 21-amino-acid linker (*e.g.*, (GGS)₇ (SEQ ID NO: 597)) may give a 3-8, 3-7, 3-6, 3-5, 3-4, 4-8, 4-7, 4-6, 4-5, 5-8, 5-7, 5-6, 6-8, 6-7, 7-8 base editing window relative to the PAM sequence. The novel finding that varying linker length may allow the dCas9 fusion proteins of the disclosure to edit nucleobases different distances from the PAM sequence affords significant clinical importance, since a PAM sequence may be of varying distance to the disease-causing mutation to be corrected in a gene. It is to be understood that the linker lengths described as examples here are not meant to be limiting.

[00189] In some embodiments, the second protein comprises an enzymatic domain. In some embodiments, the enzymatic domain is a nucleic acid editing domain. Such a nucleic acid editing domain may be, without limitation, a nuclease, a nickase, a recombinase, a deaminase, a methyltransferase, a methylase, an acetylase, or an acetyltransferase. Non-limiting exemplary binding domains that may be used in accordance with this disclosure include transcriptional activator domains and transcriptional repressor domains.

Deaminase Domains

[00190] In some embodiments, second protein comprises a nucleic acid editing domain. In some embodiments, the nucleic acid editing domain can catalyze a C to U base change. In some embodiments, the nucleic acid editing domain is a deaminase domain. In some embodiments, the deaminase is a cytidine deaminase or a cytosine deaminase. In some embodiments, the deaminase is an apolipoprotein B mRNA-editing complex (APOBEC) family deaminase. In some embodiments, the deaminase is an APOBEC1 deaminase. In some embodiments, the deaminase is an APOBEC2 deaminase. In some embodiments, the deaminase is an APOBEC3 deaminase. In some embodiments, the deaminase is an APOBEC3A deaminase. In some embodiments, the deaminase is an APOBEC3B deaminase. In some embodiments, the deaminase is an APOBEC3C deaminase. In some embodiments, the deaminase is an APOBEC3D deaminase. In some embodiments, the deaminase is an

APOBEC3E deaminase. In some embodiments, the deaminase is an APOBEC3F deaminase. In some embodiments, the deaminase is an APOBEC3G deaminase. In some embodiments, the deaminase is an APOBEC3H deaminase. In some embodiments, the deaminase is an APOBEC4 deaminase. In some embodiments, the deaminase is an activation-induced deaminase (AID). In some embodiments, the deaminase is a vertebrate deaminase. In some embodiments, the deaminase is an invertebrate deaminase. In some embodiments, the deaminase is a human, chimpanzee, gorilla, monkey, cow, dog, rat, or mouse deaminase. In some embodiments, the deaminase is a human deaminase. In some embodiments, the deaminase is a rat deaminase, *e.g.*, rAPOBEC1. In some embodiments, the deaminase is a *Petromyzon marinus* cytidine deaminase 1 (pmCDA1). In some embodiments, the deaminase is a human APOBEC3G (SEQ ID NO: 275). In some embodiments, the deaminase is a fragment of the human APOBEC3G (SEQ ID NO: 5740). In some embodiments, the deaminase is a human APOBEC3G variant comprising a D316R_D317R mutation (SEQ ID NO: 5739). In some embodiments, the deaminase is a fragment of the human APOBEC3G and comprising mutations corresponding to the D316R_D317R mutations in SEQ ID NO: 275 (SEQ ID NO: 5741).

[00191] In some embodiments, the nucleic acid editing domain is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to the deaminase domain of any one of SEQ ID NOs: 266-284, 607-610, 5724-5736, or 5738-5741. In some embodiments, the nucleic acid editing domain comprises the amino acid sequence of any one of SEQ ID NOs: 266-284, 607-610, 5724-5736, or 5738-5741.

Deaminase Domains that Modulate the Editing Window of Base Editors

[00192] Some aspects of the disclosure are based on the recognition that modulating the deaminase domain catalytic activity of any of the fusion proteins provided herein, for example by making point mutations in the deaminase domain, affect the processivity of the fusion proteins (*e.g.*, base editors). For example, mutations that reduce, but do not eliminate, the catalytic activity of a deaminase domain within a base editing fusion protein can make it less likely that the deaminase domain will catalyze the deamination of a residue adjacent to a target residue, thereby narrowing the deamination window. The ability to narrow the deamination window may prevent unwanted deamination of residues adjacent of specific target residues, which may decrease or prevent off-target effects.

[00193] In some embodiments, any of the fusion proteins provided herein comprise a deaminase domain (*e.g.*, a cytidine deaminase domain) that has reduced catalytic deaminase activity. In some embodiments, any of the fusion proteins provided herein comprise a deaminase domain (*e.g.*, a cytidine deaminase domain) that has a reduced catalytic deaminase activity as compared to an appropriate control. For example, the appropriate control may be the deaminase activity of the deaminase prior to introducing one or more mutations into the deaminase. In other embodiments, the appropriate control may be a wild-type deaminase. In some embodiments, the appropriate control is a wild-type apolipoprotein B mRNA-editing complex (APOBEC) family deaminase. In some embodiments, the appropriate control is an APOBEC1 deaminase, an APOBEC2 deaminase, an APOBEC3A deaminase, an APOBEC3B deaminase, an APOBEC3C deaminase, an APOBEC3D deaminase, an APOBEC3F deaminase, an APOBEC3G deaminase, or an APOBEC3H deaminase. In some embodiments, the appropriate control is an activation induced deaminase (AID). In some embodiments, the appropriate control is a cytidine deaminase 1 from *Petromyzon marinus* (pmCDA1). In some embodiments, the deaminase domain may be a deaminase domain that has at least 1%, at least 5%, at least 15%, at least 20%, at least 25%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 80%, at least 90%, or at least 95% less catalytic deaminase activity as compared to an appropriate control.

[00194] In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising one or more mutations selected from the group consisting of H121X, H122X, R126X, R126X, R118X, W90X, W90X, and R132X of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase, wherein X is any amino acid. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising one or more mutations selected from the group consisting of H121R, H122R, R126A, R126E, R118A, W90A, W90Y, and R132E of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase.

[00195] In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising one or more mutations selected from the group consisting of D316X, D317X, R320X, R320X, R313X, W285X, W285X, R326X of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase, wherein X is any amino acid. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising one or more mutations selected from the group consisting of D316R, D317R, R320A, R320E, R313A, W285A, W285Y, R326E of

hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase.

[00196] In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a H121R and a H122R mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R126A mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R126E mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R118A mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W90A mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W90Y mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R132E mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W90Y and a R126E mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R126E and a R132E mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W90Y and a R132E mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W90Y, R126E, and R132E mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase.

[00197] In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a D316R and a D317R mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R320A mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R320E mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R313A mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W285A mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W285Y mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R326E mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W285Y and a R320E mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a R320E and a R326E mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W285Y and a R326E mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase. In some embodiments, any of the fusion proteins provided herein comprise an APOBEC deaminase comprising a W285Y, R320E, and R326E mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase.

[00198] Some aspects of this disclosure provide fusion proteins comprising (i) a nuclease-inactive Cas9 domain; and (ii) a nucleic acid editing domain. In some embodiments, a

nuclease-inactive Cas9 domain (dCas9), comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to the amino acid sequence of a Cas9 as provided by any one of SEQ ID NOs: 10-263, and comprises mutations that inactivate the nuclease activity of Cas9. Mutations that render the nuclease domains of Cas9 inactive are well-known in the art. For example, the DNA cleavage domain of Cas9 is known to include two subdomains, the HNH nuclease subdomain and the RuvC1 subdomain. The HNH subdomain cleaves the strand complementary to the gRNA, whereas the RuvC1 subdomain cleaves the non-complementary strand. Mutations within these subdomains can silence the nuclease activity of Cas9. For example, the mutations D10A and H840A completely inactivate the nuclease activity of *S. pyogenes* Cas9 (Jinek *et al.*, *Science*. 337:816-821(2012); Qi *et al.*, *Cell*. 28;152(5):1173-83 (2013)). In some embodiments, the dCas9 of this disclosure comprises a D10A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the dCas9 of this disclosure comprises a H840A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the dCas9 of this disclosure comprises both D10A and H840A mutations of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. In some embodiments, the Cas9 further comprises a histidine residue at position 840 of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. The presence of the catalytic residue H840 restores the activity of the Cas9 to cleave the non-edited strand containing a G opposite the targeted C. Restoration of H840 does not result in the cleavage of the target strand containing the C. In some embodiments, the dCas9 comprises an amino acid sequence of SEQ ID NO: 263. It is to be understood that other mutations that inactivate the nuclease domains of Cas9 may also be included in the dCas9 of this disclosure.

[00199] The Cas9 or dCas9 domains comprising the mutations disclosed herein, may be a full-length Cas9, or a fragment thereof. In some embodiments, proteins comprising Cas9, or fragments thereof, are referred to as “Cas9 variants.” A Cas9 variant shares homology to Cas9, or a fragment thereof. For example a Cas9 variant is at least about 70% identical, at least about 80% identical, at least about 90% identical, at least about 95% identical, at least about 96% identical, at least about 97% identical, at least about 98% identical, at least about

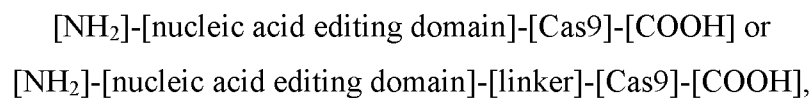
99% identical, at least about 99.5% identical, or at least about 99.9% to wild type Cas9. In some embodiments, the Cas9 variant comprises a fragment of Cas9 (*e.g.*, a gRNA binding domain or a DNA-cleavage domain), such that the fragment is at least about 70% identical, at least about 80% identical, at least about 90% identical, at least about 95% identical, at least about 96% identical, at least about 97% identical, at least about 98% identical, at least about 99% identical, at least about 99.5% identical, or at least about 99.9% identical to the corresponding fragment of wild type Cas9, *e.g.*, a Cas9 comprising the amino acid sequence of SEQ ID NO: 10.

[00200] Any of the Cas9 fusion proteins of this disclosure may further comprise a nucleic acid editing domain (*e.g.*, an enzyme that is capable of modifying nucleic acid, such as a deaminase). In some embodiments, the nucleic acid editing domain is a DNA-editing domain. In some embodiments, the nucleic acid editing domain has deaminase activity. In some embodiments, the nucleic acid editing domain comprises or consists of a deaminase or deaminase domain. In some embodiments, the deaminase is a cytidine deaminase. In some embodiments, the deaminase is an apolipoprotein B mRNA-editing complex (APOBEC) family deaminase. In some embodiments, the deaminase is an APOBEC1 family deaminase. In some embodiments, the deaminase is an activation-induced cytidine deaminase (AID). Some nucleic-acid editing domains as well as Cas9 fusion proteins including such domains are described in detail herein. Additional suitable nucleic acid editing domains will be apparent to the skilled artisan based on this disclosure and knowledge in the field.

[00201] Some aspects of the disclosure provide a fusion protein comprising a Cas9 domain fused to a nucleic acid editing domain, wherein the nucleic acid editing domain is fused to the N-terminus of the Cas9 domain. In some embodiments, the Cas9 domain and the nucleic acid editing domain are fused via a linker. In some embodiments, the linker comprises a (GGGS)_n (SEQ ID NO: 265), a (GGGGS)_n (SEQ ID NO: 5), a (G)_n, an (EAAAK)_n (SEQ ID NO: 6), a (GGS)_n, (SGGS)_n (SEQ ID NO: 4288), an SGSETPGTSESATPES (SEQ ID NO: 7) motif (see, *e.g.*, Guilinger JP, Thompson DB, Liu DR. Fusion of catalytically inactive Cas9 to FokI nuclease improves the specificity of genome modification. *Nat. Biotechnol.* 2014; 32(6): 577-82; the entire contents are incorporated herein by reference), or an (XP)_n motif, or a combination of any of these, wherein n is independently an integer between 1 and 30. In some embodiments, n is independently 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, or 30, or if more than one linker or more than one linker motif is present, any combination thereof. In some embodiments, the linker comprises a (GGS)_n motif,

wherein n is 1,2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15. In some embodiments, the linker comprises a (GGS)_n motif, wherein n is 1, 3, or 7. In some embodiments, the linker comprises the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7). Additional suitable linker motifs and linker configurations will be apparent to those of skill in the art. In some embodiments, suitable linker motifs and configurations include those described in Chen *et al.*, Fusion protein linkers: property, design and functionality. *Adv Drug Deliv Rev.* 2013; 65(10):1357-69, the entire contents of which are incorporated herein by reference.

Additional suitable linker sequences will be apparent to those of skill in the art based on the instant disclosure. In some embodiments, the general architecture of exemplary Cas9 fusion proteins provided herein comprises the structure:

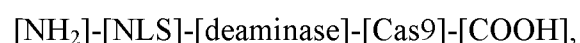


wherein NH₂ is the N-terminus of the fusion protein, and COOH is the C-terminus of the fusion protein.

[00202] The fusion proteins of the present disclosure may comprise one or more additional features. For example, in some embodiments, the fusion protein comprises a nuclear localization sequence (NLS). In some embodiments, the NLS of the fusion protein is localized between the nucleic acid editing domain and the Cas9 domain. In some embodiments, the NLS of the fusion protein is localized C-terminal to the Cas9 domain.

[00203] Other exemplary features that may be present are localization sequences, such as cytoplasmic localization sequences, export sequences, such as nuclear export sequences, or other localization sequences, as well as sequence tags that are useful for solubilization, purification, or detection of the fusion proteins. Suitable protein tags provided herein include, but are not limited to, biotin carboxylase carrier protein (BCCP) tags, myc-tags, calmodulin-tags, FLAG-tags, hemagglutinin (HA)-tags, polyhistidine tags, also referred to as histidine tags or His-tags, maltose binding protein (MBP)-tags, nus-tags, glutathione-S-transferase (GST)-tags, green fluorescent protein (GFP)-tags, thioredoxin-tags, S-tags, Softags (*e.g.*, Softag 1, Softag 3), strep-tags, biotin ligase tags, FAsH tags, V5 tags, and SBP-tags. Additional suitable sequences will be apparent to those of skill in the art. In some embodiments, the fusion protein comprises one or more His tags.

[00204] In some embodiments, the nucleic acid editing domain is a deaminase. For example, in some embodiments, the general architecture of exemplary Cas9 fusion proteins with a deaminase domain comprises the structure:



[NH₂]-[Cas9]-[deaminase]-[COOH],
 [NH₂]-[deaminase]-[Cas9]-[COOH], or
 [NH₂]-[deaminase]-[Cas9]-[NLS]-[COOH]

wherein NLS is a nuclear localization sequence, NH₂ is the N-terminus of the fusion protein, and COOH is the C-terminus of the fusion protein. Nuclear localization sequences are known in the art and would be apparent to the skilled artisan. For example, NLS sequences are described in Plank *et al.*, PCT/EP2000/011690, the contents of which are incorporated herein by reference for their disclosure of exemplary nuclear localization sequences. In some embodiments, a NLS comprises the amino acid sequence PKKKRKV (SEQ ID NO: 741) or MDSLLMNRKFLYQFKNVRWAKGRRETYLC (SEQ ID NO: 742). In some embodiments, a linker is inserted between the Cas9 and the deaminase. In some embodiments, the NLS is located C-terminal of the Cas9 domain. In some embodiments, the NLS is located N-terminal of the Cas9 domain. In some embodiments, the NLS is located between the deaminase and the Cas9 domain. In some embodiments, the NLS is located N-terminal of the deaminase domain. In some embodiments, the NLS is located C-terminal of the deaminase domain.

[00205] One exemplary suitable type of nucleic acid editing domain is a cytidine deaminase, for example, of the APOBEC family. The apolipoprotein B mRNA-editing complex (APOBEC) family of cytidine deaminase enzymes encompasses eleven proteins that serve to initiate mutagenesis in a controlled and beneficial manner.²⁹ One family member, activation-induced cytidine deaminase (AID), is responsible for the maturation of antibodies by converting cytosines in ssDNA to uracils in a transcription-dependent, strand-biased fashion.³⁰ The apolipoprotein B editing complex 3 (APOBEC3) enzyme provides protection to human cells against a certain HIV-1 strain via the deamination of cytosines in reverse-transcribed viral ssDNA.³¹ These proteins all require a Zn²⁺-coordinating motif (His-X-Glu-X₂₃₋₂₆-Pro-Cys-X₂₋₄-Cys; SEQ ID NO: 598) and bound water molecule for catalytic activity. The Glu residue acts to activate the water molecule to a zinc hydroxide for nucleophilic attack in the deamination reaction. Each family member preferentially deaminates at its own particular “hotspot”, ranging from WRC (W is A or T, R is A or G) for hAID, to TTC for hAPOBEC3F.³² A recent crystal structure of the catalytic domain of APOBEC3G revealed a secondary structure comprised of a five-stranded β-sheet core flanked by six α-helices, which is believed to be conserved across the entire family.³³ The active center loops have been shown to be responsible for both ssDNA binding and in determining “hotspot” identity.³⁴

Overexpression of these enzymes has been linked to genomic instability and cancer, thus highlighting the importance of sequence-specific targeting.³⁵

[00206] Some aspects of this disclosure relate to the recognition that the activity of cytidine deaminase enzymes such as APOBEC enzymes can be directed to a specific site in genomic DNA. Without wishing to be bound by any particular theory, advantages of using Cas9 as a recognition agent include (1) the sequence specificity of Cas9 can be easily altered by simply changing the sgRNA sequence; and (2) Cas9 binds to its target sequence by denaturing the dsDNA, resulting in a stretch of DNA that is single-stranded and therefore a viable substrate for the deaminase. It should be understood that other catalytic domains, or catalytic domains from other deaminases, can also be used to generate fusion proteins with Cas9, and that the disclosure is not limited in this regard.

[00207] Some aspects of this disclosure are based on the recognition that Cas9:deaminase fusion proteins can efficiently deaminate nucleotides at positions 3-11 according to the numbering scheme in Figure 3. In view of the results provided herein regarding the nucleotides that can be targeted by Cas9:deaminase fusion proteins, a person of skill in the art will be able to design suitable guide RNAs to target the fusion proteins to a target sequence that comprises a nucleotide to be deaminated.

[00208] In some embodiments, the deaminase domain and the Cas9 domain are fused to each other via a linker. Various linker lengths and flexibilities between the deaminase domain (*e.g.*, AID) and the Cas9 domain can be employed (*e.g.*, ranging from very flexible linkers of the form (GGGGS)_n (SEQ ID NO: 5), (GGS)_n, and (G)_n to more rigid linkers of the form (EAAAK)_n (SEQ ID NO: 6), (SGGS)_n (SEQ ID NO: 4288), SGSETPGTSESATPES (SEQ ID NO: 7) (*see, e.g.*, Guilinger JP, Thompson DB, Liu DR. Fusion of catalytically inactive Cas9 to FokI nuclease improves the specificity of genome modification. *Nat. Biotechnol.* 2014; 32(6): 577-82; the entire contents are incorporated herein by reference) and (XP)_n)³⁶ in order to achieve the optimal length for deaminase activity for the specific application. In some embodiments, the linker comprises a (GGS)_n motif, wherein n is 1, 3, or 7. In some embodiments, the linker comprises a (an SGSETPGTSESATPES (SEQ ID NO: 7) motif.

[00209] Some exemplary suitable nucleic-acid editing domains, *e.g.*, deaminases and deaminase domains, that can be fused to Cas9 domains according to aspects of this disclosure are provided below. It should be understood that, in some embodiments, the active domain of the respective sequence can be used, *e.g.*, the domain without a localizing signal (nuclear localization sequence, without nuclear export signal, cytoplasmic localizing signal).

[00210] Human AID:

MDSLLMNRRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNGC
 HVELLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGPNLSLRIFTAR
 LYFCEDRKAPEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHEN
 SVRLSRQLRRILLPLYEVDDLRDAFRTLGL (SEQ ID NO: 266)

(underline: nuclear localization sequence; double underline: nuclear export signal)

[00211] Mouse AID:

MDSLLMKQKKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSCSLDFGHNRNKSGC
 HVELLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVAEFLRWPNLSLRIFTAR
 LYFCEDRKAPEPEGLRRLHRAGVQIGIMTFKDYFYCWNTFVENRERTFKAWEGLHEN
 SVRLTRQLRRILLPLYEVDDLRDAFRMLGF (SEQ ID NO: 267)

(underline: nuclear localization sequence; double underline: nuclear export signal)

[00212] Dog AID:

MDSLLMKQRKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSFSLDFGHNRNKSGC
 HVELLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSLRIFAAR
 LYFCEDRKAPEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENREKTFKAWEGLHEN
 SVRLSRQLRRILLPLYEVDDLRDAFRTLGL (SEQ ID NO: 268)

(underline: nuclear localization sequence; double underline: nuclear export signal)

[00213] Bovine AID:

MDSLLKKQRQFLYQFKNVRWAKGRHETYLCYVVKRRDSPTSFSLDFGHNRNKAGC
 HVELLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSLRIFTAR
 LYFCDKERKAPEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHE
 NSVRLSRQLRRILLPLYEVDDLRDAFRTLGL (SEQ ID NO: 269)

(underline: nuclear localization sequence; double underline: nuclear export signal)

[00214] Rat AID

MAVGSKPKAALVGPHWERERIWCFLCSTGLGTQQTGQTSRWLRPAATQDPVSPPRS
 LLMKQRKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSFSLDFGYLRNKSGCHVE
 LLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGPNLSLRIFTARLTG
 WGALPAGLMSPARPSDYFYCWNTFVENHERTFKAWEGLHENSSVRLSRRLRRILLPL
YEVDDLRDAFRTLGL (SEQ ID NO: 5725)

(underline: nuclear localization sequence; double underline: nuclear export signal)

[00215] Mouse APOBEC-3:

MGPFCLGCSHRKCYSPIRNLISQETFKFHFKNLGYAKGRKDTFLCYEVTRKDCDSPVS
 LHHGVFKNKDNIAEICFLYWFHDKVLKVLSPREEFKITWYMSWSPCFECAEQIVRFLA

THHNLSLDIFSSRLYNVQDPETQQNLCRLVQEGAQVAAMDLYEFKKCWKKFVDNG
 GRRFRPWKRLLTNFRYQDSKLQEILRPCYIPVSSSSSTLSNICLTKGLPETRFCVEGR
 RMDPLSEEFYSQFYQNRVKHLCYYHRMKPYLCYQLEQFNGQAPLKGCLLSEKGGKQ
HAEILFLDKIRSMELSQVTITCYLTWSPCPNCAWQLAAFKRDRPDLILHIYTSRLYFHWK
 RPFQKGLCSLWQSGILVDVMDLPQFTDCWTNFWNPKRPFWPWKGLEIISRRTQRRLR
 RIKESWGLQDLVNDFGNLQLGPPMS (SEQ ID NO: 270)

(italic: nucleic acid editing domain)

[00216] Rat APOBEC-3:

MGPFCLGCSHRKCYSPIRNLISQETFKFHFKNLRYAIDRKDTFLCYEVTRKDCDSPVS
 LHHGVFKNKDNIAEICFLYWFHDKVLKVLSPREEFKITWYMSWSPCFECAEQVLRFLA
 THHNLSLDIFSSRLYNIRDPENQQNLCRLVQEGAQVAAMDLYEFKKCWKKFVDNGG
 RRFWRPWKLLTNFRYQDSKLQEILRPCYIPVSSSSSTLSNICLTKGLPETRFCVERRR
 VHLLSEEFYSQFYQNRVKHLCYYHGVPYLCYQLEQFNGQAPLKGCLLSEKGGKQH
AEILFLDKIRSMELSQVIITCYLTWSPCPNCAWQLAAFKRDRPDLILHIYTSRLYFHWKR
 PFQKGLCSLWQSGILVDVMDLPQFTDCWTNFWNPKRPFWPWKGLEIISRRTQRRLHR
 IKESWGLQDLVNDFGNLQLGPPMS (SEQ ID NO: 271)

(italic: nucleic acid editing domain)

[00217] Rhesus macaque APOBEC-3G:

MVEPMDPRTFVSNFNRPILSGLNTVWLCCEVKTKDPSGPPLDAKIFQGKVYSKAKY
HPEMRFLRWFHKWRQLHHDQEYKVTWYVSWSPCTRCANSVATFLAKDPKVTLTIFVA
 RLYYFWKPDYQQALRILCQKRGGPHATMKIMNYNEFQDCWNKFVDGRGKPFKPRN
 NLPKHYTLLQATLGELLRHLMDPGTFTSNFNKPVVSGQHETYLCYKVERLHNDT
 WVPLNQHRGFLRNQAPNIHGFPKGRHAELCFLDLIPFWKLDGQQYRVTCFTSWSPCF
 CAQEMAKFISNNEHVSLCIFAARIYDDQGRYQEGLRALHRDGAKIAMMNYSEFEYC
 WDTFVDRQGRPFQPWDGLDEHSQALSGRLRAI (SEQ ID NO: 272)

(italic: nucleic acid editing domain; underline: cytoplasmic localization signal)

[00218] Chimpanzee APOBEC-3G:

MKPHFRNPVERMYQDTFSDNFYNRPILSHRNTVWLCYEVKTKGSPRPPLDAKIFRGQ
VYSKLYHPEMRFFHWF~~SK~~WRKLHRDQEYEV~~W~~YISWSPCTKCTRDVATFLAEDPKV
 TLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKIMNYDEFQHCWSKFVYSQRE
 LFEPWNNLPKYYILLHIMLGEILRHSM DPPTFTSNFNNELWVRGRHETYLCYEVERL
 HNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFLDVIPFWKLDLHQDYRVTCFTS
 WSPCFSCAQEMAKFISNNKHVSLCIFAARIYDDQGRYQEGLRRTLAKAGAKISIMTYSE
 FKHCWDTFVDHQGCPFPQWDGLEEHSQALSGRLRAILQNQGN (SEQ ID NO: 273)

(italic: nucleic acid editing domain; underline: cytoplasmic localization signal)

[00219] Green monkey APOBEC-3G:

MNPQIRNMVEQMEPDIFVYYFNNRPILSGRNTVWLCYEVKTKDPSGPPLDANIFQGK
LYPEAKDHPEMKFLHWFRKWRQLHRDQEYEV TWYVSWSPCTRCANSVATFLAEDPKV
 TLTIFVARLYYFWKPDYQQALRLCQERGGPHATMKIMNYNEFQHCWNEFVDGQG
 KPFKPRKNLPKH Y TLLHATLGELLRHVMDPGTFTSNFNKPKWVSGQRETYLCYKVE
 RSHNDTWVLLNQHRGFLRNQAPDRHGFPKGRHAELCFDLIPFWKLDDQQYRVTCFT
 SWSPCFSCAQKMAKFISNNKHVSLCIFAARIYDDQGRCQEGLRTLHRDGAKIAVMNY
 SEFEYCWDTFVDRQGRPFQPWDGLDEHSQALSGRLRAI (SEQ ID NO: 274)

(italic: nucleic acid editing domain; underline: cytoplasmic localization signal)

[00220] Human APOBEC-3G:

MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTVWLCYEVKTKGPSRPPLDAKIFRGQ
VYSELKYHPEMRFFHWFSKWRKLHRDQEYEV TWYISWSPCTKCTRDMATFLAEDPKV
 TLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKIMNYDEFQHCWSKFVYSQRE
 LFEPWNNLPKYYILLHIMLGEILRHSM DPPTFTFNFNNEPWVRGRHETLYCYEVERM
 HNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFDVIPFWKLDDLDQDYRVTCFTS
 WSPCFSCAQEMAKFISK NKHVSLCIFTARIYDDQGRCQEGLRTLAEAGAKISIMTYSE
 FKHCWDTFVDHQGCPFQPWDGLDEHSQDLSGRLRAILQNQEN (SEQ ID NO: 275)

(italic: nucleic acid editing domain; underline: cytoplasmic localization signal)

[00221] Human APOBEC-3F:

MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTVWLCYEVKTKGPSRPRLDAKIFRGQ
 VYSQPEHHAEMCFLSWFCGNQLPAYKCFQITWVSWTPCPDCVAKLAEFLAHPNVTL
 TISAARLYYYWERDYRRALCRLSQA GARVKIMDDEEFAYCWENFVYSEGQPFMPW
 YKFDDNYAFLHRTLKEILRNPMEAMYPHIFYHFKNLRKAYGRNESWLCFTMEVVK
 HHSPVSWKRGVFRNQVDPETHCHAERCFLSWFCDDILSPNTNYEVTWYTSWSPCPECA
 GEVAEFLARHSNVNLTIFTARLYYFWDTDYQEGLRSLSQEGASVEIMGYKDFKYCW
 ENFVYNDDEPFKPKWGLKYNFLFLDSKLQEILE (SEQ ID NO: 276)

(italic: nucleic acid editing domain)

[00222] Human APOBEC-3B:

MNPQIRNPMERMYRDTFYDNFENEPILYGRSYTWLCYEVKIKRGRSNLLWDTGVFR
 GQVYFKPQYHAEMCFLSWFCGNQLPAYKCFQITWVSWTPCPDCVAKLAEFLSEHPN
 VTLTISAARLYYYWERDYRRALCRLSQA GARVTIMDYEEFAYCWENFVYNEGQQF
 MPWYKFDENYAFLHRTLKEILRYLMDPDTFTFNFNNDPLVLRRTYLYCYEVERLD

NGTWVLMQHMGLFCNEAKNLLCGFYGRHAELRFLDLVPSLQLDPAQIYRVTWFISWS
PCFSWGCAGEVRAFLQENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIMTY
 DEFEYCWDTFVYRQGCPFQPWDGLEEHSQALSGRLRAILQNQGN (SEQ ID NO: 277)
 (italic: nucleic acid editing domain)

[00223] Rat APOBEC-3B:

MQPQGLGPNAGMGPVCLGCSHRPYSPIRNPLKKLYQQTFYFHFKNVRYAWGRKN
 NFLCYEVNGMDCALPVPLRQGVFRKQGHIIHAELCFIYWFHDKVLRVLSPMEEFKVT
 WYMSWSPCSKCAEQVARFLAAHRNLSLAIFSSRLYYLNRPNYQQKLCRLIQEGVH
 VAAMDLPFEFKKCWNKFVDNDGQPFPRPWRMLRINFSFYDCKLQEIFSRMNLLREDVF
 YLQFNNSHRVKPVQNRYYRRKSYLCYQLERANGQEPLKGYLLYKKGEQHVEILFLE
 KMRSMELSQVRITCYLTWSPCPNCARQLAAFKKDHPDLILRIYTSRLYFYWRKKFQK
 GLCTLWRSGLHVDVMDLPQFADCWTNFVNPQRPFPRPWNELEKNSWRIQRRLRIKE
 SWGL (SEQ ID NO: 5729)

[00224] Bovine APOBEC-3B:

DGWEVAFRSGTVLKAGVLGVSMTTEGWAGSGHPGQGACVWTPGTRNTMNLREVL
 FKQQFGNQPRVPAPYYRRKTYLCYQLKQRNDLTLDRGCFRNKKQRHAEIRFIDKINS
 LDLNPSQSYKIICYITWSPCPNCANELVNFITRNNHLKLEIFASRLYFHWIKSFKMGLQ
 DLQONAGISVAVMTHTEFEDCWEQFVDNQSRPFQPWDKLEQYSASIRRLRILTAPI
 (SEQ ID NO: 5730)

[00225] Chimpanzee APOBEC-3B:

MNPQIRNPMEMYQRTFYYNFENEPILYGRSYTWLCYEVKIRRGHSNLLWDTGVFR
 GQMQSPEHHAEMCFLSWFCGNQLSAYKCFQITWFVSWTPCPDCVAKLAKFLAEHP
 NVTLTISAARLYYYWERDYRRALCRLSQAGARVKIMDDEEFAYCWENFVYNEGQPF
 MPWYKFDDNYAFLHRTLKEIIRHLMDDPTFTFNFNNDPLVLRRHQTYLCYEVERLD
 NGTWVLMQHMGLFCNEAKNLLCGFYGRHAELRFLDLVPSLQLDPAQIYRVTWFIS
 WSPCFSWGCAGQVRAFLQENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIM
 TYDEFEYCWDTFVYRQGCPFQPWDGLEEHSQALSGRLRAILQVRASSLCMVPHRPPP
 PPQSPGPCLPLCSEPPLGSLPTGRPAPSLPFLLTASFSPPPASLPPLPSLSLSPGHLPVP
 SFHSLTSCSIQPPCSSRIRETEGWASVSKEGRDLG (SEQ ID NO: 5731)

[00226] Human APOBEC-3C:

MNPQIRNPMKAMYPGTFYFQFKNLWEANDRNETWLCFTVEGIKRRSVVSWKTGVF
 RNQVDSETHCHAERCFLSWFCDDILSPNTKYQVTWYTSWSPCPDCAGEVAEFLARHSN
 VNLTIFTARLYYFQYPCYQEGLRSLSQEGVAVEIMDYEDFKYCWENFVYNDNEPFKP
 WKGLKTNFRLKRRLRESLQ (SEQ ID NO: 278)

(italic: nucleic acid editing domain)

[00227] Gorilla APOBEC3C

MNPQIRNPMKAMYPGTFYFQFKNLWEANDRNETWLCFTVEGIKRRSVVSWKTGVF
RNQVDSETH*CHAERCFLSWFCDDILSPNTNYQVTWYTSWSPCPECAGEVAEFLARHSN*
VNLTIFTARLYYFQD TDYQEGLRSLSQEGVAVKIMDYKDFKYCWENFVYNDDEPFK
PWKGLKYNFRFLKRRLQEILE (SEQ ID NO: 5726)

(italic: nucleic acid editing domain)

[00228] Human APOBEC-3A:

MEASPASGPRHLMDPHIFTSNFFNNGIGRHKTYLCYEVERLDNGTSVKMDQHRGFLH
NQAKNLLCGFYGR*HAEFLDLVPSLQLDPAQIYRVTWFISWSPCFSWGCAGEVRAFLQ*
ENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIMTYDEFKHCWDTFVDHQGC
PFQPWDGLDEHSQALSGRLRAILQNQGN (SEQ ID NO: 279)

(italic: nucleic acid editing domain)

[00229] Rhesus macaque APOBEC-3A:

MDGSPASRPRHLMDPNTFTFNFNNDLSVRGRHQTYLCYEVERLDNGTWVPM DERR
GFLCNKAKNVPCGDY*GCHVELRFLCEVPSWQLDPAQTYRVTWFISWSPCFRRGCAGQ*
VRVFLQENKHVRLRIFAARIYDYDPLYQEALRTL RDAGAQVSIMTYEEFKHCWDTF
VDRQGRPFQPWDGLDEHSQALSGRLRAILQNQGN (SEQ ID NO: 5727)

(italic: nucleic acid editing domain)

[00230] Bovine APOBEC-3A:

MDEYTF TENFNQGWPSKTYLCYEMERLDGDATIPLDEYKGFVRNKGLDQPEKPC*H*
AELYFLGKIHSWNLDNRNQHYRLTCFISWSPCYDCAQKLT TFLKENHHISLHILASRIYTH
NRFGCHQSGLCELQAAGARITIMTFEDFKHCWETFVDHKGKPFQWEGLVNKSQAL
CTELQAILKTQQN (SEQ ID NO: 5728)

(italic: nucleic acid editing domain)

[00231] Human APOBEC-3H:

MALLTAETFRLQFNKRRLRRPYYPRKALLCYQLTPQNGSTPTRGYFENKKK*CHAEI*
CFINEIKSMGLDETQCYQVTCYLTWSPCSSCAWELVDFIKAHDHLNLGIFASRLYYHWC
KPQQKGLRLLCGSQVPVEVMGFPKFADCWENFVDHEKPLSFNPYKMLEELDKNSRA
IKRRLERIKIPGVRAQGRYMDILCDAEV (SEQ ID NO: 280)

(italic: nucleic acid editing domain)

[00232] Rhesus macaque APOBEC-3H:

MALLTAKTFSLQFNKR RVNKPYYPRKALLCYQLTPQNGSTPTRGHLKNKKKDHA*E*
IRFINKIKSMGLDETQCYQVTCYLTWSPCSCAGELVDFIKAHRLNLRIFASRLYYH

WRPNYQEGLLLLCGSQVPVEVMGLPEFTDCWENFVDHKEPPSFNPSEKLEELDKN
 QAIKRRLERIKSRSDVLENGRLSLQLGPVTPSSSIRNSR (SEQ ID NO: 5732)

[00233] Human APOBEC-3D:

MNPQIRNPMERMYRDTFYDNFENEPILYGRSYTWLCYEYVKIKRGRSNLLWDTGVFR
 GPVLPKRQSNHRQEVYFRFEN*HAEMCFLSWFCGNRLPANRRFQITWFWVSWNPCLPCVV*
 KVTKFLAEHPNVTLTISAARLYYYRDRDWRWVLLRLHKAGARVKIMDYEDFAFCW
 ENFVCNEGQPFMPWYKFDDNYASLHRTLKEILRNPMEAMYPHIFYHFKNLLKACG
 RNESWLCFTMEVTKHHSVFRKRGVFRNQVDPETH*CHAERCFLSWFCDDILSPNTNY*
EVTWYTSWSPCECAGEVAEFLARHSNVNLTIFTARLCYFWDDTDYQEGLCSLSQEGAS
 VKIMGYKDFVSCWKNFVYSDDEPFKPKWGLQTNFRLLKRRLREILQ (SEQ ID NO:
 281)

(italic: nucleic acid editing domain)

[00234] Human APOBEC-1:

MTSEKGPSTGDPTLRRRIEPWEFDVFYDPRELKKEACLLYEIKWGMSRKIWRSSGKN
 TTNHVEVNFIEKFTSERDFHPSMSCSITWFLSWSPCWECQAIREFLSRHPGVTLVIYV
 ARLFWHMDQQNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYP
 PLWMMLYALELHCILSLPPCLKISRRWQNHLTFFRLHLQNCHYQTIPPHILLATGLIH
 PSVAWR (SEQ ID NO: 282)

[00235] Mouse APOBEC-1:

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKKEACLLYEINWGGRHSVWRHTSQN
 TSNHVEVNFIEKFTTERRYFRPNTRCSITWFLSWSPCGECSRAITEFLSRHPYVTLFIYIA
 RLYHHTDQRNRQGLRDLISSGVTIQIMTEQEYCYCWRNFVNYPSPNEAYWPRYPHL
 WVKLYVLELYCIIILGLPPCLKILRRKQPQLTFFTITLQTCHYQRIPPHLLWATGLK
 (SEQ ID NO: 283)

[00236] Rat APOBEC-1:

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKKEACLLYEINWGGRHSIWRHTSQNT
 NKHVEVNFIEKFTTERRYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWPYPHLLW
 VRLYVLELYCIIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLK (SEQ
 ID NO: 284)

[00237] Human APOBEC-2:

MAQKEEAAVATEAASQNGEDLENLDDPEKLELIELPPFEIVTGERLPANFFKFQFRN
 VE

YSSGRNKTFLCYVVEAQGGQVQASRGYLEDEHAAAHAEAEAFFNTILPAFDPALR
 YNVTWYVSSSPCAACADRIIKTLSTKNLRLLLILVGRLFMWEEPEIQAALKKLKEAG
 CKLRIMKPQDFEYVWQNFVEQEEGESKAFQPWEDIQENFLYYEEKLADILK (SEQ ID
 NO: 5733)

[00238] Mouse APOBEC-2:

MAQKEEAAEAAAPASQNGDDLENLEDPEKLLKELIDLPPFEIVTGVRLPVNFFKFQFR
 NVEYSSGRNKTFLCYVVEVQSKGGQAQATQGYLEDEHAGAHAEAEAFFNTILPAFDP
 ALKYNVTWYVSSSPCAACADRILKTLSTKNLRLLLILVSRLFMWEEPEVQAALKKLK
 EAGCKLRIMKPQDFEYIWQNFVEQEEGESKAFEPWEDIQENFLYYEEKLADILK
 (SEQ ID NO: 5734)

[00239] Rat APOBEC-2:

MAQKEEAAEAAAPASQNGDDLENLEDPEKLLKELIDLPPFEIVTGVRLPVNFFKFQFR
 NVEYSSGRNKTFLCYVVEAQSKGGQVQATQGYLEDEHAGAHAEAEAFFNTILPAFDP
 ALKYNVTWYVSSSPCAACADRILKTLSTKNLRLLLILVSRLFMWEEPEVQAALKKLK
 EAGCKLRIMKPQDFEYLWQNFVEQEEGESKAFEPWEDIQENFLYYEEKLADILK
 (SEQ ID NO: 5735)

[00240] Bovine APOBEC-2:

MAQKEEAAAAAEPASQNGEEVENLEDPEKLLKELIELPPFEIVTGERLPAHYFKFQFRN
 VE
 YSSGRNKTFLCYVVEAQSKGGQVQASRGYLEDEHATNHAEAEAFFNSIMPTFDPALR
 YMVTWYVSSSPCAACADRIVKTLNKTKNLRLLLILVGRLFMWEEPEIQAALRKLKEA
 GCRLRIMKPQDFEYIWQNFVEQEEGESKAFEPWEDIQENFLYYEEKLADILK (SEQ
 ID NO: 5736)

[00241] Petromyzon marinus CDA1 (pmCDA1)

MTDAEYVRIHEKLDIYTFKKQFFNNKKSVSHRCYVLFELKRRGERRACFWGYAVNK
 PQSGTERGIHAEIFSIRKVEEYLRDNPQGFTINWYSSWSPCADCAEKILEWYNQELRG
 NGHTLKIWACKLYYEKNARNQIGLWNLRDNGVGLNVMVSEHYQCCRKIFIQSSHNQ
 LNENRWLEKTLKRAEKRRSELSIMIQVKILHTTKSPAV (SEQ ID NO: 5738)

[00242] Human APOBEC3G D316R_D317R

MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTVWLCYEVKTKGPSRPLDAKIFRGQ
 VYSELKYHPEMRFFHWFSKWRKLHRDQEYEV TWYISWSPCTKCTRDMATFLAEDP
 KVTLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKIMNYDEFQHCWSKFFVYSQ
 RELFEPWNNLPKYIILLHIMLGEILRHSMDPPTFTFNFNNEPWRGRHETYLCYEVEER

MHNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFLDVIPFWKLDLDQDYRVTC
FTSWSPCFSCAQEMAKFISKKNKHVSLCIFTARIYRRQGRCQEGLRTLAEAGAKISIMT
YSEFKHCWDTFVDHQGCPFQPWDGLDEHSQDLSGRLRAILQNQEN (SEQ ID NO:
5739)

[00243] Human APOBEC3G chain A

MDPPTFTFNFNNEPWVRGRHETYLCYEVMHNDTWVLLNQRRGFLCNQAPHKHG
FLEGRHAELCFLDVIPFWKLDLDQDYRVTCFTSWSPCFSCAQEMAKFISKKNKHVSLCI
FTARIYDDQGRCQEGLRTLAEAGAKISIMTYSEFKHCWDTFVDHQGCPFQPWDGLD
EHSQDLSGRLRAILQ (SEQ ID NO: 5740)

[00244] Human APOBEC3G chain A D120R_D121R

MDPPTFTFNFNNEPWVRGRHETYLCYEVMHNDTWVLLNQRRGFLCNQAPHKHG
FLEGRHAELCFLDVIPFWKLDLDQDYRVTCFTSWSPCFSCAQEMAKFISKKNKHVSLCI
FTARIYRRQGRCQEGLRTLAEAGAKISIMTYSEFKHCWDTFVDHQGCPFQPWDGLDE
HSQDLSGRLRAILQ (SEQ ID NO: 5741)

[00245] In some embodiments, fusion proteins as provided herein comprise the full-length amino acid of a nucleic acid editing enzyme, *e.g.*, one of the sequences provided above. In other embodiments, however, fusion proteins as provided herein do not comprise a full-length sequence of a nucleic acid editing enzyme, but only a fragment thereof. For example, in some embodiments, a fusion protein provided herein comprises a Cas9 domain and a fragment of a nucleic acid editing enzyme, *e.g.*, wherein the fragment comprises a nucleic acid editing domain. Exemplary amino acid sequences of nucleic acid editing domains are shown in the sequences above as italicized letters, and additional suitable sequences of such domains will be apparent to those of skill in the art.

[00246] Additional suitable nucleic-acid editing enzyme sequences, *e.g.*, deaminase enzyme and domain sequences, that can be used according to aspects of this invention, *e.g.*, that can be fused to a nuclease-inactive Cas9 domain, will be apparent to those of skill in the art based on this disclosure. In some embodiments, such additional enzyme sequences include deaminase enzyme or deaminase domain sequences that are at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% similar to the sequences provided herein. Additional suitable Cas9 domains, variants, and sequences will also be apparent to those of skill in the art. Examples of such additional suitable Cas9 domains include, but are not limited to, D10A,

D10A/D839A/H840A, and D10A/D839A/H840A/N863A mutant domains (See, *e.g.*, Prashant *et al.*, CAS9 transcriptional activators for target specificity screening and paired nickases for cooperative genome engineering. *Nature Biotechnology*. 2013; 31(9): 833-838 the entire contents of which are incorporated herein by reference). In some embodiments, the Cas9 comprises a histidine residue at position 840 of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260. The presence of the catalytic residue H840 restores the activity of the Cas9 to cleave the non-edited strand containing a G opposite the targeted C. Restoration of H840 does not result in the cleavage of the target strand containing the C.

[00247] Additional suitable strategies for generating fusion proteins comprising a Cas9 domain and a deaminase domain will be apparent to those of skill in the art based on this disclosure in combination with the general knowledge in the art. Suitable strategies for generating fusion proteins according to aspects of this disclosure using linkers or without the use of linkers will also be apparent to those of skill in the art in view of the instant disclosure and the knowledge in the art. For example, Gilbert *et al.*, CRISPR-mediated modular RNA-guided regulation of transcription in eukaryotes. *Cell*. **2013**; 154(2):442-51, showed that C-terminal fusions of Cas9 with VP64 using 2 NLS's as a linker (SPKKKRKVEAS, SEQ ID NO: 599), can be employed for transcriptional activation. Mali *et al.*, CAS9 transcriptional activators for target specificity screening and paired nickases for cooperative genome engineering. *Nat Biotechnol*. **2013**; 31(9):833-8, reported that C-terminal fusions with VP64 without linker can be employed for transcriptional activation. And Maeder *et al.*, CRISPR RNA-guided activation of endogenous human genes. *Nat Methods*. **2013**; 10: 977-979, reported that C-terminal fusions with VP64 using a Gly₄Ser (SEQ ID NO: 5) linker can be used as transcriptional activators. Recently, dCas9-FokI nuclease fusions have successfully been generated and exhibit improved enzymatic specificity as compared to the parental Cas9 enzyme (In Guilinger JP, Thompson DB, Liu DR. Fusion of catalytically inactive Cas9 to FokI nuclease improves the specificity of genome modification. *Nat. Biotechnol*. 2014; 32(6): 577-82, and in Tsai SQ, Wyvekens N, Khayter C, Foden JA, Thapar V, Reyon D, Goodwin MJ, Aryee MJ, Joung JK. Dimeric CRISPR RNA-guided FokI nucleases for highly specific genome editing. *Nat Biotechnol*. 2014; 32(6):569-76. PMID: 24770325 a SGSETPGTSESATPES (SEQ ID NO: 7) or a GGGGS (SEQ ID NO: 5) linker was used in FokI-dCas9 fusion proteins, respectively).

[00248] Some aspects of this disclosure provide fusion proteins comprising (i) a Cas9 enzyme or domain (*e.g.*, a first protein); and (ii) a nucleic acid-editing enzyme or domain

(e.g., a second protein). In some aspects, the fusion proteins provided herein further include (iii) a programmable DNA-binding protein, for example, a zinc-finger domain, a TALE, or a second Cas9 protein (e.g., a third protein). Without wishing to be bound by any particular theory, fusing a programmable DNA-binding protein (e.g., a second Cas9 protein) to a fusion protein comprising (i) a Cas9 enzyme or domain (e.g., a first protein); and (ii) a nucleic acid-editing enzyme or domain (e.g., a second protein) may be useful for improving specificity of the fusion protein to a target nucleic acid sequence, or for improving specificity or binding affinity of the fusion protein to bind target nucleic acid sequence that does not contain a canonical PAM (NGG) sequence. In some embodiments, the third protein is a Cas9 protein (e.g., a second Cas9 protein). In some embodiments, the third protein is any of the Cas9 proteins provided herein. In some embodiments, the third protein is fused to the fusion protein N-terminal to the Cas9 protein (e.g., the first protein). In some embodiments, the third protein is fused to the fusion protein C-terminal to the Cas9 protein (e.g., the first protein). In some embodiments, the Cas9 domain (e.g., the first protein) and the third protein (e.g., a second Cas9 protein) are fused via a linker (e.g., a second linker). In some embodiments, the linker comprises a (GGGGS)_n (SEQ ID NO: 5), a (G)_n, an (EAAAK)_n (SEQ ID NO: 6), a (GGG)_n, (SGGS)_n (SEQ ID NO: 4288), an SGSETPGTSESATPES (SEQ ID NO: 7), or an (XP)_n motif, or a combination of any of these, wherein n is independently an integer between 1 and 30. In some embodiments, the general architecture of exemplary Cas9 fusion proteins provided herein comprises the structure:

[NH2]-[nucleic acid-editing enzyme or domain]-[Cas9]-[third protein]-[COOH];
 [NH2]-[third protein]-[Cas9]-[nucleic acid-editing enzyme or domain]-[COOH];
 [NH2]-[Cas9]-[nucleic acid-editing enzyme or domain]-[third protein]-[COOH];
 [NH2]-[third protein]-[nucleic acid-editing enzyme or domain]-[Cas9]-[COOH];
 [NH2]-[UGI]-[nucleic acid-editing enzyme or domain]-[Cas9]-[third protein]-
 [COOH];
 [NH2]-[UGI]-[third protein]-[Cas9]-[nucleic acid-editing enzyme or domain]-
 [COOH];
 [NH2]-[UGI]-[Cas9]-[nucleic acid-editing enzyme or domain]-[third protein]-
 [COOH];
 [NH2]-[UGI]-[third protein]-[nucleic acid-editing enzyme or domain]-[Cas9]-
 [COOH];
 [NH2]-[nucleic acid-editing enzyme or domain]-[Cas9]-[third protein]-[UGI]-
 [COOH];

[NH2]-[third protein]-[Cas9]-[nucleic acid-editing enzyme or domain]-[UGI]-[COOH];

[NH2]-[Cas9]-[nucleic acid-editing enzyme or domain]-[third protein]-[UGI]-[COOH]; or

[NH2]-[third protein]-[nucleic acid-editing enzyme or domain]-[Cas9]-[UGI]-[COOH];

wherein NH2 is the N-terminus of the fusion protein, and COOH is the C-terminus of the fusion protein. In some embodiments, the “[”-“]” used in the general architecture above indicates the presence of an optional linker sequence. In other examples, the general architecture of exemplary Cas9 fusion proteins provided herein comprises the structure:

[NH2]-[nucleic acid-editing enzyme or domain]-[Cas9]-[second Cas9 protein]-[COOH];

[NH2]-[second Cas9 protein]-[Cas9]-[nucleic acid-editing enzyme or domain]-[COOH];

[NH2]-[Cas9]-[nucleic acid-editing enzyme or domain]-[second Cas9 protein]-[COOH];

[NH2]-[second Cas9 protein]-[nucleic acid-editing enzyme or domain]-[Cas9]-[COOH];

[NH2]-[UGI]-[nucleic acid-editing enzyme or domain]-[Cas9]-[second Cas9 protein]-[COOH];

[NH2]-[UGI]-[second Cas9 protein]-[Cas9]-[nucleic acid-editing enzyme or domain]-[COOH];

[NH2]-[UGI]-[Cas9]-[nucleic acid-editing enzyme or domain]-[second Cas9 protein]-[COOH];

[NH2]-[UGI]-[second Cas9 protein]-[nucleic acid-editing enzyme or domain]-[Cas9]-[COOH];

[NH2]-[nucleic acid-editing enzyme or domain]-[Cas9]-[second Cas9 protein]-[UGI]-[COOH];

[NH2]-[second Cas9 protein]-[Cas9]-[nucleic acid-editing enzyme or domain]-[UGI]-[COOH];

[NH2]-[Cas9]-[nucleic acid-editing enzyme or domain]-[second Cas9 protein]-[UGI]-[COOH]; or

[NH2]-[second Cas9 protein]-[nucleic acid-editing enzyme or domain]-[Cas9]-[UGI]-[COOH];

wherein NH₂ is the N-terminus of the fusion protein, and COOH is the C-terminus of the fusion protein. In some embodiments, the “[” used in the general architecture above indicates the presence of an optional linker sequence. In some embodiments, the second Cas9 is a dCas9 protein. In some examples, the general architecture of exemplary Cas9 fusion proteins provided herein comprises a structure as shown in Figure 3. It should be appreciated that any of the proteins provided in any of the general architectures of exemplary Cas9 fusion proteins may be connected by one or more of the linkers provided herein. In some embodiments, the linkers are the same. In some embodiments, the linkers are different. In some embodiments, one or more of the proteins provided in any of the general architectures of exemplary Cas9 fusion proteins are not fused via a linker. In some embodiments, the fusion proteins further comprise a nuclear targeting sequence, for example a nuclear localization sequence. In some embodiments, fusion proteins provided herein further comprise a nuclear localization sequence (NLS). In some embodiments, the NLS is fused to the N-terminus of the fusion protein. In some embodiments, the NLS is fused to the C-terminus of the fusion protein. In some embodiments, the NLS is fused to the N-terminus of the third protein. In some embodiments, the NLS is fused to the C-terminus of the third protein. In some embodiments, the NLS is fused to the N-terminus of the Cas9 protein. In some embodiments, the NLS is fused to the C-terminus of the Cas9 protein. In some embodiments, the NLS is fused to the N-terminus of the nucleic acid-editing enzyme or domain. In some embodiments, the NLS is fused to the C-terminus of the nucleic acid-editing enzyme or domain. In some embodiments, the NLS is fused to the N-terminus of the UGI protein. In some embodiments, the NLS is fused to the C-terminus of the UGI protein. In some embodiments, the NLS is fused to the fusion protein via one or more linkers. In some embodiments, the NLS is fused to the fusion protein without a linker

Uracil glycosylase inhibitor fusion proteins

[00249] Some aspects of the disclosure relate to fusion proteins that comprise a uracil glycosylase inhibitor (UGI) domain. In some embodiments, any of the fusion proteins provided herein that comprise a Cas9 domain (*e.g.*, a nuclease active Cas9 domain, a nuclease inactive dCas9 domain, or a Cas9 nickase) may be further fused to a UGI domain either directly or via a linker. Some aspects of this disclosure provide deaminase-dCas9 fusion proteins, deaminase-nuclease active Cas9 fusion proteins and deaminase-Cas9 nickase fusion proteins with increased nucleobase editing efficiency. Without wishing to be bound by any particular theory, cellular DNA-repair response to the presence of U:G heteroduplex DNA

may be responsible for the decrease in nucleobase editing efficiency in cells. For example, uracil DNA glycosylase (UDG) catalyzes removal of U from DNA in cells, which may initiate base excision repair, with reversion of the U:G pair to a C:G pair as the most common outcome. As demonstrated in the Examples below, Uracil DNA Glycosylase Inhibitor (UGI) may inhibit human UDG activity. Thus, this disclosure contemplates a fusion protein comprising dCas9-nucleic acid editing domain further fused to a UGI domain. This disclosure also contemplates a fusion protein comprising a Cas9 nickase-nucleic acid editing domain further fused to a UGI domain. It should be understood that the use of a UGI domain may increase the editing efficiency of a nucleic acid editing domain that is capable of catalyzing a C to U change. For example, fusion proteins comprising a UGI domain may be more efficient in deaminating C residues. In some embodiments, the fusion protein comprises the structure:

[deaminase]-[optional linker sequence]-[dCas9]-[optional linker sequence]-[UGI];
 [deaminase]-[optional linker sequence]-[UGI]-[optional linker sequence]-[dCas9];
 [UGI]-[optional linker sequence]-[deaminase]-[optional linker sequence]-[dCas9];
 [UGI]-[optional linker sequence]-[dCas9]-[optional linker sequence]-[deaminase];
 [dCas9]-[optional linker sequence]-[deaminase]-[optional linker sequence]-[UGI]; or
 [dCas9]-[optional linker sequence]-[UGI]-[optional linker sequence]-[deaminase].

In other embodiments, the fusion protein comprises the structure:

[deaminase]-[optional linker sequence]-[Cas9 nickase]-[optional linker sequence]-
 [UGI];
 [deaminase]-[optional linker sequence]-[UGI]-[optional linker sequence]-[Cas9
 nickase];
 [UGI]-[optional linker sequence]-[deaminase]-[optional linker sequence]-[Cas9
 nickase];
 [UGI]-[optional linker sequence]-[Cas9 nickase]-[optional linker sequence]-
 [deaminase];
 [Cas9 nickase]-[optional linker sequence]-[deaminase]-[optional linker sequence]-
 [UGI]; or
 [Cas9 nickase]-[optional linker sequence]-[UGI]-[optional linker sequence]-
 [deaminase].

[00250] In some embodiments, the fusion proteins provided herein do not comprise a linker sequence. In some embodiments, one or both of the optional linker sequences are present.

[00251] In some embodiments, the “-” used in the general architecture above indicates the presence of an optional linker sequence. In some embodiments, the fusion proteins comprising a UGI further comprise a nuclear targeting sequence, for example a nuclear localization sequence. In some embodiments, fusion proteins provided herein further comprise a nuclear localization sequence (NLS). In some embodiments, the NLS is fused to the N-terminus of the fusion protein. In some embodiments, the NLS is fused to the C-terminus of the fusion protein. In some embodiments, the NLS is fused to the N-terminus of the UGI protein. In some embodiments, the NLS is fused to the C-terminus of the UGI protein. In some embodiments, the NLS is fused to the N-terminus of the Cas9 protein. In some embodiments, the NLS is fused to the C-terminus of the Cas9 protein. In some embodiments, the NLS is fused to the N-terminus of the deaminase. In some embodiments, the NLS is fused to the C-terminus of the deaminase. In some embodiments, the NLS is fused to the N-terminus of the second Cas9. In some embodiments, the NLS is fused to the C-terminus of the second Cas9. In some embodiments, the NLS is fused to the fusion protein via one or more linkers. In some embodiments, the NLS is fused to the fusion protein without a linker. In some embodiments, the NLS comprises an amino acid sequence of any one of the NLS sequences provided or referenced herein. In some embodiments, the NLS comprises an amino acid sequence as set forth in SEQ ID NO: 741 or SEQ ID NO: 742.

[00252] In some embodiments, a UGI domain comprises a wild-type UGI or a UGI as set forth in SEQ ID NO: 600. In some embodiments, the UGI proteins provided herein include fragments of UGI and proteins homologous to a UGI or a UGI fragment. For example, in some embodiments, a UGI domain comprises a fragment of the amino acid sequence set forth in SEQ ID NO: 600. In some embodiments, a UGI fragment comprises an amino acid sequence that comprises at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% of the amino acid sequence as set forth in SEQ ID NO: 600. In some embodiments, a UGI comprises an amino acid sequence homologous to the amino acid sequence set forth in SEQ ID NO: 600 or an amino acid sequence homologous to a fragment of the amino acid sequence set forth in SEQ ID NO: 600. In some embodiments, proteins comprising UGI or fragments of UGI or homologs of UGI or UGI fragments are referred to as “UGI variants.” A UGI variant shares homology to UGI, or a fragment thereof. For example a UGI variant is at least 70% identical, at least 75% identical, at least 80% identical, at least 85% identical, at least 90% identical, at least 95% identical, at least 96% identical, at least 97% identical, at least 98% identical, at least 99% identical, at least 99.5% identical, or

at least 99.9% identical to a wild type UGI or a UGI as set forth in SEQ ID NO: 600. In some embodiments, the UGI variant comprises a fragment of UGI, such that the fragment is at least 70% identical, at least 80% identical, at least 90% identical, at least 95% identical, at least 96% identical, at least 97% identical, at least 98% identical, at least 99% identical, at least 99.5% identical, or at least 99.9% to the corresponding fragment of wild-type UGI or a UGI as set forth in SEQ ID NO: 600. In some embodiments, the UGI comprises the following amino acid sequence:

>sp|P14739|UNGI_BPPB2 Uracil-DNA glycosylase inhibitor
MTNLSDIIEKETGKQLVIQESILMLPEEVVEEVIGNKPESDILVHTAYDESTDENVMLLT
SDAPEYKPWALVIQDSNGENKIKML (SEQ ID NO: 600)

[00253] Suitable UGI protein and nucleotide sequences are provided herein and additional suitable UGI sequences are known to those in the art, and include, for example, those published in Wang et al., Uracil-DNA glycosylase inhibitor gene of bacteriophage PBS2 encodes a binding protein specific for uracil-DNA glycosylase. *J. Biol. Chem.* 264:1163-1171(1989); Lundquist et al., Site-directed mutagenesis and characterization of uracil-DNA glycosylase inhibitor protein. Role of specific carboxylic amino acids in complex formation with *Escherichia coli* uracil-DNA glycosylase. *J. Biol. Chem.* 272:21408-21419(1997); Ravishankar et al., X-ray analysis of a complex of *Escherichia coli* uracil DNA glycosylase (EcUDG) with a proteinaceous inhibitor. The structure elucidation of a prokaryotic UDG. *Nucleic Acids Res.* 26:4880-4887(1998); and Putnam et al., Protein mimicry of DNA from crystal structures of the uracil-DNA glycosylase inhibitor protein and its complex with *Escherichia coli* uracil-DNA glycosylase. *J. Mol. Biol.* 287:331-346(1999), the entire contents of each are incorporated herein by reference.

[00254] It should be appreciated that additional proteins may be uracil glycosylase inhibitors. For example, other proteins that are capable of inhibiting (*e.g.*, sterically blocking) a uracil-DNA glycosylase base-excision repair enzyme are within the scope of this disclosure. Additionally, any proteins that block or inhibit base-excision repair are also within the scope of this disclosure. In some embodiments, a protein that binds DNA is used. In another embodiment, a substitute for UGI is used. In some embodiments, a uracil glycosylase inhibitor is a protein that binds single-stranded DNA. For example, a uracil glycosylase inhibitor may be an *Erwinia tasmaniensis* single-stranded binding protein. In some embodiments, the single-stranded binding protein comprises the amino acid sequence (SEQ ID NO: 322). In some embodiments, a uracil glycosylase inhibitor is a protein that

binds uracil. In some embodiments, a uracil glycosylase inhibitor is a protein that binds uracil in DNA. In some embodiments, a uracil glycosylase inhibitor is a catalytically inactive uracil DNA-glycosylase protein. In some embodiments, a uracil glycosylase inhibitor is a catalytically inactive uracil DNA-glycosylase protein that does not excise uracil from the DNA. For example, a uracil glycosylase inhibitor is a UdgX. In some embodiments, the UdgX comprises the amino acid sequence (SEQ ID NO: 323). As another example, a uracil glycosylase inhibitor is a catalytically inactive UDG. In some embodiments, a catalytically inactive UDG comprises the amino acid sequence (SEQ ID NO: 324). It should be appreciated that other uracil glycosylase inhibitors would be apparent to the skilled artisan and are within the scope of this disclosure. In some embodiments, a uracil glycosylase inhibitor is a protein that is homologous to any one of SEQ ID NOs: 322-324.. In some embodiments, a uracil glycosylase inhibitor is a protein that is at least 50% identical, at least 55% identical, at least 60% identical, at least 65% identical, at least 70% identical, at least 75% identical, at least 80% identical, at least 85% identical, at least 90% identical, at least 95% identical, at least 96% identical, at least 98% identical, at least 99% identical, or at least 99.5% identical to any one of SEQ ID NOs: 322-324.

Erwinia tasmaniensis SSB (thermostable single-stranded DNA binding protein)

MASRGVNVKVLVGNLQDPEVRYMPNGGAVANITLATSESWRDKQTGETKEKTEW
 HRVVLFGKLAEVAGEYLRKGSQVYIEGALQTRKWTDQAGVEKYTTEVVVNVGGT
 MQMLGGRSQGGASAGGQNGGSNNGWGQPQQPQGGNQFSGGAQQQARPQQQPQ
 QNNAPANNEPPIDFDDDDIP (SEQ ID NO: 322)

UdgX (binds to Uracil in DNA but does not excise)

MAGAQDFVPHTADLAELAAAAGECRGCGLYRDATQAVFGAGGRSARIMMIGEQPG
 DKEDLAGLPFVGPAGRLLDRALEAADIDRDALYVTNAVKHFKFTRAAGGKRRIHKT
 PSRTEVVACRPWLIAEMTSVEPDVVLLGATAAKALLGNDFRVTQHRGEVLHVDDV
 PGDPALVATVHPSSLLRGPKEERESAFAGLVDDLVAADVVP (SEQ ID NO: 323)

UDG (catalytically inactive human UDG, binds to Uracil in DNA but does not excise)

MIGQKTLYSFFSPARKRHAPSPEPAVQGTGVAGVPEESGDAAAIPAKKAPAGQEEP
 GTPPSSPLSAEQLDRIQRNKAALLRLAARNVPVGFGEWKKHLSGEFGKPYFIKLM
 GFVAEERKHVTVYPPPHQVFTWTQMCDIKDVKVVILGQEPYHGPNQAHGLCFVSR
 PVPPPPSLENIYKELSTDIEDFVHPGHGDLGWAQGVLLLNAVLTVRAHQANSHKE

RGWEQFTDAVVSWLNQNSNGLVFLWGSYAQKKGSAIDRKRHHVLQTAHPSPLSV
YRGFFGCRHFSTNELLQKSGKKPIDWKEL (SEQ ID NO: 324)

[00255] In some embodiments, the nucleic acid editing domain is a deaminase domain. In some embodiments, the deaminase is a cytosine deaminase or a cytidine deaminase. In some embodiments, the deaminase is an apolipoprotein B mRNA-editing complex (APOBEC) family deaminase. In some embodiments, the deaminase is an APOBEC1 deaminase. In some embodiments, the deaminase is an APOBEC2 deaminase. In some embodiments, the deaminase is an APOBEC3 deaminase. In some embodiments, the deaminase is an APOBEC3A deaminase. In some embodiments, the deaminase is an APOBEC3B deaminase. In some embodiments, the deaminase is an APOBEC3C deaminase. In some embodiments, the deaminase is an APOBEC3D deaminase. In some embodiments, the deaminase is an APOBEC3E deaminase. In some embodiments, the deaminase is an APOBEC3F deaminase. In some embodiments, the deaminase is an APOBEC3G deaminase. In some embodiments, the deaminase is an APOBEC3H deaminase. In some embodiments, the deaminase is an APOBEC4 deaminase. In some embodiments, the deaminase is an activation-induced deaminase (AID). In some embodiments, the deaminase is a rat APOBEC1 (SEQ ID NO: 282). In some embodiments, the deaminase is a human APOBEC1 (SEQ ID NO: 284). In some embodiments, the deaminase is a *Petromyzon marinus* cytidine deaminase 1 (pmCDA1). In some embodiments, the deaminase is a human APOBEC3G (SEQ ID NO: 275). In some embodiments, the deaminase is a fragment of the human APOBEC3G (SEQ ID NO: 5740). In some embodiments, the deaminase is a human APOBEC3G variant comprising a D316R_D317R mutation (SEQ ID NO: 5739). In some embodiments, the deaminase is a fragment of the human APOBEC3G and comprising mutations corresponding to the D316R_D317R mutations in SEQ ID NO: 275 (SEQ ID NO: 5741).

[00256] In some embodiments, the linker comprises a (GGGS)_n (SEQ ID NO: 265), (GGGG)_n (SEQ ID NO: 5), a (G)_n, an (EAAAK)_n (SEQ ID NO: 6), a (GGS)_n, an SGSETPGTSESATPES (SEQ ID NO: 7), or an (XP)_n motif, or a combination of any of these, wherein n is independently an integer between 1 and 30.

[00257] Suitable UGI protein and nucleotide sequences are provided herein and additional suitable UGI sequences are known to those in the art, and include, for example, those published in Wang *et al.*, Uracil-DNA glycosylase inhibitor gene of bacteriophage PBS2 encodes a binding protein specific for uracil-DNA glycosylase. *J. Biol. Chem.* 264:1163-

1171(1989); Lundquist *et al.*, Site-directed mutagenesis and characterization of uracil-DNA glycosylase inhibitor protein. Role of specific carboxylic amino acids in complex formation with Escherichia coli uracil-DNA glycosylase. *J. Biol. Chem.* 272:21408-21419(1997); Ravishankar *et al.*, X-ray analysis of a complex of Escherichia coli uracil DNA glycosylase (EcUDG) with a proteinaceous inhibitor. The structure elucidation of a prokaryotic UDG. *Nucleic Acids Res.* 26:4880-4887(1998); and Putnam *et al.*, Protein mimicry of DNA from crystal structures of the uracil-DNA glycosylase inhibitor protein and its complex with Escherichia coli uracil-DNA glycosylase. *J. Mol. Biol.* 287:331-346(1999), the entire contents of which are incorporated herein by reference. In some embodiments, the optional linker comprises a (GGS)_n motif, wherein n is 1, 2, 3, 4, 5, 6, 7, 8, 9, 19, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20. In some embodiments, the optional linker comprises a (GGS)_n motif, wherein n is 1, 3, or 7. In some embodiments, the optional linker comprises the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7), which is also referred to as the XTEN linker in the Examples.

[00258] In some embodiments, a Cas9 nickase may further facilitate the removal of a base on the non-edited strand in an organism whose genome is edited *in vivo*. The Cas9 nickase, as described herein, may comprise a D10A mutation in SEQ ID NO: 10, or a corresponding mutation in any of SEQ ID NOs: 11-260. In some embodiments, the Cas9 nickase of this disclosure may comprise a histidine at mutation 840 of SEQ ID NO: 10, or a corresponding residue in any of SEQ ID NOs: 11-260. Such fusion proteins comprising the Cas9 nickase, can cleave a single strand of the target DNA sequence, *e.g.*, the strand that is not being edited. Without wishing to be bound by any particular theory, this cleavage may inhibit mis-match repair mechanisms that reverse a C to U edit made by the deaminase.

Cas9 complexes with guide RNAs

[00259] Some aspects of this disclosure provide complexes comprising any of the fusion proteins provided herein, and a guide RNA bound to a Cas9 domain (*e.g.*, a dCas9, a nuclease active Cas9, or a Cas9 nickase) of fusion protein.

[00260] In some embodiments, the guide RNA is from 15-100 nucleotides long and comprises a sequence of at least 10 contiguous nucleotides that is complementary to a target sequence. In some embodiments, the guide RNA is 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, or 50 nucleotides long. In some embodiments, the guide RNA comprises a sequence of 15, 16,

17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, or 40 contiguous nucleotides that is complementary to a target sequence. In some embodiments, the target sequence is a DNA sequence. In some embodiments, the target sequence is a sequence in the genome of a mammal. In some embodiments, the target sequence is a sequence in the genome of a human. In some embodiments, the 3' end of the target sequence is immediately adjacent to a canonical PAM sequence (NGG). In some embodiments, the guide RNA is complementary to a sequence associated with a disease or disorder. In some embodiments, the guide RNA is complementary to a sequence associated with a disease or disorder having a mutation in a gene selected from the genes disclosed in any one of Tables 1-3. In some embodiments, the guide RNA comprises a nucleotide sequence of any one of the guide sequences provided in Table 2 or Table 3. Exemplary sequences in the human genome that may be targeted by the complexes of this disclosure are provided herein in Tables 1-3.

Methods of using Cas9 fusion proteins

[00261] Some aspects of this disclosure provide methods of using the Cas9 proteins, fusion proteins, or complexes provided herein. For example, some aspects of this disclosure provide methods comprising contacting a DNA molecule (a) with any of the the Cas9 proteins or fusion proteins provided herein, and with at least one guide RNA, wherein the guide RNA is about 15-100 nucleotides long and comprises a sequence of at least 10 contiguous nucleotides that is complementary to a target sequence; or (b) with a Cas9 protein, a Cas9 fusion protein, or a Cas9 protein or fusion protein complex with at least one gRNA as provided herein. In some embodiments, the 3' end of the target sequence is not immediately adjacent to a canonical PAM sequence (NGG). In some embodiments, the 3' end of the target sequence is immediately adjacent to an AGC, GAG, TTT, GTG, or CAA sequence.

[00262] In some embodiments, the target DNA sequence comprises a sequence associated with a disease or disorder. In some embodiments, the target DNA sequence comprises a point mutation associated with a disease or disorder. In some embodiments, the activity of the Cas9 protein, the Cas9 fusion protein, or the complex results in a correction of the point mutation. In some embodiments, the target DNA sequence comprises a T→C point mutation associated with a disease or disorder, and wherein the deamination of the mutant C base results in a sequence that is not associated with a disease or disorder. In some embodiments, the target DNA sequence encodes a protein and wherein the point mutation is in a codon and results in a change in the amino acid encoded by the mutant codon as compared to the wild-type codon. In some embodiments, the deamination of the mutant C results in a change of the amino acid

encoded by the mutant codon. In some embodiments, the deamination of the mutant C results in the codon encoding the wild-type amino acid. In some embodiments, the contacting is *in vivo* in a subject. In some embodiments, the subject has or has been diagnosed with a disease or disorder. In some embodiments, the disease or disorder is cystic fibrosis, phenylketonuria, epidermolytic hyperkeratosis (EHK), Charcot-Marie-Toot disease type 4J, neuroblastoma (NB), von Willebrand disease (vWD), myotonia congenital, hereditary renal amyloidosis, dilated cardiomyopathy (DCM), hereditary lymphedema, familial Alzheimer's disease, HIV, Prion disease, chronic infantile neurologic cutaneous articular syndrome (CINCA), desmin-related myopathy (DRM), a neoplastic disease associated with a mutant PI3KCA protein, a mutant CTNNB1 protein, a mutant HRAS protein, or a mutant p53 protein.

[00263] Some embodiments provide methods for using the Cas9 DNA editing fusion proteins provided herein. In some embodiments, the fusion protein is used to introduce a point mutation into a nucleic acid by deaminating a target nucleobase, *e.g.*, a C residue. In some embodiments, the deamination of the target nucleobase results in the correction of a genetic defect, *e.g.*, in the correction of a point mutation that leads to a loss of function in a gene product. In some embodiments, the genetic defect is associated with a disease or disorder, *e.g.*, a lysosomal storage disorder or a metabolic disease, such as, for example, type I diabetes. In some embodiments, the methods provided herein are used to introduce a deactivating point mutation into a gene or allele that encodes a gene product that is associated with a disease or disorder. For example, in some embodiments, methods are provided herein that employ a Cas9 DNA editing fusion protein to introduce a deactivating point mutation into an oncogene (*e.g.*, in the treatment of a proliferative disease). A deactivating mutation may, in some embodiments, generate a premature stop codon in a coding sequence, which results in the expression of a truncated gene product, *e.g.*, a truncated protein lacking the function of the full-length protein.

[00264] In some embodiments, the purpose of the methods provide herein is to restore the function of a dysfunctional gene via genome editing. The Cas9 deaminase fusion proteins provided herein can be validated for gene editing-based human therapeutics *in vitro*, *e.g.*, by correcting a disease-associated mutation in human cell culture. It will be understood by the skilled artisan that the fusion proteins provided herein, *e.g.*, the fusion proteins comprising a Cas9 domain and a nucleic acid deaminase domain can be used to correct any single point T → C or A → G mutation. In the first case, deamination of the mutant C back to U corrects the mutation, and in the latter case, deamination of the C that is base-paired with the mutant G, followed by a round of replication, corrects the mutation.

[00265] An exemplary disease-relevant mutation that can be corrected by the provided fusion proteins *in vitro* or *in vivo* is the H1047R (A3140G) polymorphism in the PI3KCA protein. The phosphoinositide-3-kinase, catalytic alpha subunit (PI3KCA) protein acts to phosphorylate the 3-OH group of the inositol ring of phosphatidylinositol. The PI3KCA gene has been found to be mutated in many different carcinomas, and thus it is considered to be a potent oncogene.³⁷ In fact, the A3140G mutation is present in several NCI-60 cancer cell lines, such as, for example, the HCT116, SKOV3, and T47D cell lines, which are readily available from the American Type Culture Collection (ATCC).³⁸

[00266] In some embodiments, a cell carrying a mutation to be corrected, *e.g.*, a cell carrying a point mutation, *e.g.*, an A3140G point mutation in exon 20 of the PI3KCA gene, resulting in a H1047R substitution in the PI3KCA protein, is contacted with an expression construct encoding a Cas9 deaminase fusion protein and an appropriately designed sgRNA targeting the fusion protein to the respective mutation site in the encoding PI3KCA gene. Control experiments can be performed where the sgRNAs are designed to target the fusion enzymes to non-C residues that are within the PI3KCA gene. Genomic DNA of the treated cells can be extracted, and the relevant sequence of the PI3KCA genes PCR amplified and sequenced to assess the activities of the fusion proteins in human cell culture.

[00267] It will be understood that the example of correcting point mutations in PI3KCA is provided for illustration purposes and is not meant to limit the instant disclosure. The skilled artisan will understand that the instantly disclosed DNA-editing fusion proteins can be used to correct other point mutations and mutations associated with other cancers and with diseases other than cancer including other proliferative diseases.

[00268] The successful correction of point mutations in disease-associated genes and alleles opens up new strategies for gene correction with applications in therapeutics and basic research. Site-specific single-base modification systems like the disclosed fusions of Cas9 and deaminase enzymes or domains also have applications in “reverse” gene therapy, where certain gene functions are purposely suppressed or abolished. In these cases, site-specifically mutating Trp (TGG), Gln (CAA and CAG), or Arg (CGA) residues to premature stop codons (TAA, TAG, TGA) can be used to abolish protein function *in vitro*, *ex vivo*, or *in vivo*.

[00269] The instant disclosure provides methods for the treatment of a subject diagnosed with a disease associated with or caused by a point mutation that can be corrected by a Cas9 DNA editing fusion protein provided herein. For example, in some embodiments, a method is provided that comprises administering to a subject having such a disease, *e.g.*, a cancer associated with a PI3KCA point mutation as described above, an effective amount of a Cas9

deaminase fusion protein that corrects the point mutation or introduces a deactivating mutation into the disease-associated gene. In some embodiments, the disease is a proliferative disease. In some embodiments, the disease is a genetic disease. In some embodiments, the disease is a neoplastic disease. In some embodiments, the disease is a metabolic disease. In some embodiments, the disease is a lysosomal storage disease. Other diseases that can be treated by correcting a point mutation or introducing a deactivating mutation into a disease-associated gene will be known to those of skill in the art, and the disclosure is not limited in this respect.

[00270] The instant disclosure provides methods for the treatment of additional diseases or disorders, *e.g.*, diseases or disorders that are associated or caused by a point mutation that can be corrected by deaminase-mediated gene editing. Some such diseases are described herein, and additional suitable diseases that can be treated with the strategies and fusion proteins provided herein will be apparent to those of skill in the art based on the instant disclosure. Exemplary suitable diseases and disorders are listed below. It will be understood that the numbering of the specific positions or residues in the respective sequences depends on the particular protein and numbering scheme used. Numbering might be different, *e.g.*, in precursors of a mature protein and the mature protein itself, and differences in sequences from species to species may affect numbering. One of skill in the art will be able to identify the respective residue in any homologous protein and in the respective encoding nucleic acid by methods well known in the art, *e.g.*, by sequence alignment and determination of homologous residues. Exemplary suitable diseases and disorders include, without limitation, cystic fibrosis (see, *e.g.*, Schwank *et al.*, Functional repair of CFTR by CRISPR/Cas9 in intestinal stem cell organoids of cystic fibrosis patients. *Cell stem cell*. **2013**; 13: 653-658; and Wu *et al.*, Correction of a genetic disease in mouse via use of CRISPR-Cas9. *Cell stem cell*. **2013**; 13: 659-662, neither of which uses a deaminase fusion protein to correct the genetic defect); phenylketonuria – *e.g.*, phenylalanine to serine mutation at position 835 (mouse) or 240 (human) or a homologous residue in phenylalanine hydroxylase gene (T>C mutation) – see, *e.g.*, McDonald *et al.*, *Genomics*. **1997**; 39:402-405; Bernard-Soulier syndrome (BSS) – *e.g.*, phenylalanine to serine mutation at position 55 or a homologous residue, or cysteine to arginine at residue 24 or a homologous residue in the platelet membrane glycoprotein IX (T>C mutation) – see, *e.g.*, Noris *et al.*, *British Journal of Haematology*. **1997**; 97: 312-320, and Ali *et al.*, *Hematol*. **2014**; 93: 381-384; epidermolytic hyperkeratosis (EHK) – *e.g.*, leucine to proline mutation at position 160 or 161 (if counting the initiator methionine) or a homologous residue in keratin 1 (T>C mutation) – see, *e.g.*,

Chipev *et al.*, *Cell*. **1992**; 70: 821-828, see also accession number P04264 in the UNIPROT database at [www\[dot\]uniprot\[dot\]org](http://www.uniprot.org); chronic obstructive pulmonary disease (COPD) – *e.g.*, leucine to proline mutation at position 54 or 55 (if counting the initiator methionine) or a homologous residue in the processed form of α_1 -antitrypsin or residue 78 in the unprocessed form or a homologous residue (T>C mutation) – see, *e.g.*, Poller *et al.*, *Genomics*. **1993**; 17: 740-743, see also accession number P01011 in the UNIPROT database; Charcot-Marie-Toot disease type 4J – *e.g.*, isoleucine to threonine mutation at position 41 or a homologous residue in FIG4 (T>C mutation) – see, *e.g.*, Lenk *et al.*, *PLoS Genetics*. 2011; 7: e1002104; neuroblastoma (NB) – *e.g.*, leucine to proline mutation at position 197 or a homologous residue in Caspase-9 (T>C mutation) – see, *e.g.*, Kundu *et al.*, *3 Biotech*. **2013**, **3**:225-234; von Willebrand disease (vWD) – *e.g.*, cysteine to arginine mutation at position 509 or a homologous residue in the processed form of von Willebrand factor, or at position 1272 or a homologous residue in the unprocessed form of von Willebrand factor (T>C mutation) – see, *e.g.*, Lavergne *et al.*, *Br. J. Haematol*. **1992**, see also accession number P04275 in the UNIPROT database; 82: 66-72; myotonia congenital – *e.g.*, cysteine to arginine mutation at position 277 or a homologous residue in the muscle chloride channel gene CLCN1 (T>C mutation) – see, *e.g.*, Weinberger *et al.*, *The J. of Physiology*. **2012**; 590: 3449-3464; hereditary renal amyloidosis – *e.g.*, stop codon to arginine mutation at position 78 or a homologous residue in the processed form of apolipoprotein AII or at position 101 or a homologous residue in the unprocessed form (T>C mutation) – see, *e.g.*, Yazaki *et al.*, *Kidney Int*. **2003**; 64: 11-16; dilated cardiomyopathy (DCM) – *e.g.*, tryptophan to Arginine mutation at position 148 or a homologous residue in the FOXD4 gene (T>C mutation), see, *e.g.*, Minoretti *et al.*, *Int. J. of Mol. Med*. **2007**; 19: 369-372; hereditary lymphedema – *e.g.*, histidine to arginine mutation at position 1035 or a homologous residue in VEGFR3 tyrosine kinase (A>G mutation), see, *e.g.*, Irrthum *et al.*, *Am. J. Hum. Genet*. **2000**; 67: 295-301; familial Alzheimer's disease – *e.g.*, isoleucine to valine mutation at position 143 or a homologous residue in presenilin1 (A>G mutation), see, *e.g.*, Gallo *et al.*, *J. Alzheimer's disease*. **2011**; 25: 425-431; Prion disease – *e.g.*, methionine to valine mutation at position 129 or a homologous residue in prion protein (A>G mutation) – see, *e.g.*, Lewis *et al.*, *J. of General Virology*. **2006**; 87: 2443-2449; chronic infantile neurologic cutaneous articular syndrome (CINCA) – *e.g.*, Tyrosine to Cysteine mutation at position 570 or a homologous residue in cryopyrin (A>G mutation) – see, *e.g.*, Fujisawa *et al.* *Blood*. **2007**; 109: 2903-2911; and desmin-related myopathy (DRM) – *e.g.*, arginine to glycine mutation at position 120 or a homologous residue in $\alpha\beta$ crystallin (A>G mutation) – see, *e.g.*, Kumar *et al.*, *J.*

Biol. Chem. **1999**; 274: 24137-24141. The entire contents of all references and database entries is incorporated herein by reference.

[00271] The instant disclosure provides lists of genes comprising pathogenic T>C or A>G mutations. Provided herein, are the names of these genes, their respective SEQ ID NOs, their gene IDs, and sequences flanking the mutation site. (Tables 2 and 3). In some instances, the gRNA sequences that can be used to correct the mutations in these genes are disclosed (Tables 2 and 3).

[00272] In some embodiments, a Cas9-deaminase fusion protein recognizes canonical PAMs and therefore can correct the pathogenic T>C or A>G mutations with canonical PAMs, e.g., NGG (listed in Tables 2 and 3, SEQ ID NOs: 2540-2702 and 5084-5260), respectively, in the flanking sequences. For example, the Cas9 proteins that recognize canonical PAMs comprise an amino acid sequence that is at least 90% identical to the amino acid sequence of *Streptococcus pyogenes* Cas9 as provided by SEQ ID NO: 10, or to a fragment thereof comprising the RuvC and HNH domains of SEQ ID NO: 10.

[00273] It will be apparent to those of skill in the art that in order to target a Cas9:nucleic acid editing enzyme/domain fusion protein as disclosed herein to a target site, e.g., a site comprising a point mutation to be edited, it is typically necessary to co-express the Cas9:nucleic acid editing enzyme/domain fusion protein together with a guide RNA, e.g., an sgRNA. As explained in more detail elsewhere herein, a guide RNA typically comprises a tracrRNA framework allowing for Cas9 binding, and a guide sequence, which confers sequence specificity to the Cas9:nucleic acid editing enzyme/domain fusion protein. In some embodiments, the guide RNA comprises a structure 5'-[guide sequence]-
guuuuagagcuagaaaagcaaguuaaaauaaaggcuaguccguuaucaacuugaaaaaguggcaccgagucggugcuuuuu-3' (SEQ ID NO: 601), wherein the guide sequence comprises a sequence that is complementary to the target sequence. The guide sequence is typically 20 nucleotides long. The sequences of suitable guide RNAs for targeting Cas9:nucleic acid editing enzyme/domain fusion proteins to specific genomic target sites will be apparent to those of skill in the art based on the instant disclosure. Such suitable guide RNA sequences typically comprise guide sequences that are complementary to a nucleic sequence within 50 nucleotides upstream or downstream of the target nucleotide to be edited. Some exemplary guide RNA sequences suitable for targeting Cas9:nucleic acid editing enzyme/domain fusion proteins to specific target sequences are provided below.

Base Editor Efficiency

[00274] Some aspects of the disclosure are based on the recognition that any of the base editors provided herein are capable of modifying a specific nucleotide base without generating a significant proportion of indels. An “indel”, as used herein, refers to the insertion or deletion of a nucleotide base within a nucleic acid. Such insertions or deletions can lead to frame shift mutations within a coding region of a gene. In some embodiments, it is desirable to generate base editors that efficiently modify (*e.g.* mutate or deaminate) a specific nucleotide within a nucleic acid, without generating a large number of insertions or deletions (*i.e.*, indels) in the nucleic acid. In certain embodiments, any of the base editors provided herein are capable of generating a greater proportion of intended modifications (*e.g.*, point mutations or deaminations) versus indels. In some embodiments, the base editors provided herein are capable of generating a ratio of intended point mutations to indels that is greater than 1:1. In some embodiments, the base editors provided herein are capable of generating a ratio of intended point mutations to indels that is at least 1.5:1, at least 2:1, at least 2.5:1, at least 3:1, at least 3.5:1, at least 4:1, at least 4.5:1, at least 5:1, at least 5.5:1, at least 6:1, at least 6.5:1, at least 7:1, at least 7.5:1, at least 8:1, at least 10:1, at least 12:1, at least 15:1, at least 20:1, at least 25:1, at least 30:1, at least 40:1, at least 50:1, at least 100:1, at least 200:1, at least 300:1, at least 400:1, at least 500:1, at least 600:1, at least 700:1, at least 800:1, at least 900:1, or at least 1000:1, or more. The number of intended mutations and indels may be determined using any suitable method, for example the methods used in the below Examples.

[00275] In some embodiments, the base editors provided herein are capable of limiting formation of indels in a region of a nucleic acid. In some embodiments, the region is at a nucleotide targeted by a base editor or a region within 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides of a nucleotide targeted by a base editor. In some embodiments, any of the base editors provided herein are capable of limiting the formation of indels at a region of a nucleic acid to less than 1%, less than 1.5%, less than 2%, less than 2.5%, less than 3%, less than 3.5%, less than 4%, less than 4.5%, less than 5%, less than 6%, less than 7%, less than 8%, less than 9%, less than 10%, less than 12%, less than 15%, or less than 20%. The number of indels formed at a nucleic acid region may depend on the amount of time a nucleic acid (*e.g.*, a nucleic acid within the genome of a cell) is exposed to a base editor. In some embodiments, an number or proportion of indels is determined after at least 1 hour, at least 2 hours, at least 6 hours, at least 12 hours, at least 24 hours, at least 36 hours, at least 48 hours, at least 3 days, at least 4 days, at least 5 days, at least 7 days, at least 10 days, or at least 14 days of exposing a nucleic acid (*e.g.*, a nucleic acid within the genome of a cell) to a base editor.

[00276] Some aspects of the disclosure are based on the recognition that any of the base editors provided herein are capable of efficiently generating an intended mutation, such as a point mutation, in a nucleic acid (*e.g.* a nucleic acid within a genome of a subject) without generating a significant number of unintended mutations, such as unintended point mutations. In some embodiments, a intended mutation is a mutation that is generated by a specific base editor bound to a gRNA, specifically designed to generate the intended mutation. In some embodiments, the intended mutation is a mutation associated with a disease or disorder. In some embodiments, the intended mutation is a cytosine (C) to thymine (T) point mutation associated with a disease or disorder. In some embodiments, the intended mutation is a guanine (G) to adenine (A) point mutation associated with a disease or disorder. In some embodiments, the intended mutation is a cytosine (C) to thymine (T) point mutation within the coding region of a gene. In some embodiments, the intended mutation is a guanine (G) to adenine (A) point mutation within the coding region of a gene. In some embodiments, the intended mutation is a point mutation that generates a stop codon, for example, a premature stop codon within the coding region of a gene. In some embodiments, the intended mutation is a mutation that eliminates a stop codon. In some embodiments, the intended mutation is a mutation that alters the splicing of a gene. In some embodiments, the intended mutation is a mutation that alters the regulatory sequence of a gene (*e.g.*, a gene promotor or gene repressor). In some embodiments, any of the base editors provided herein are capable of generating a ratio of intended mutations to unintended mutations (*e.g.*, intended point mutations:unintended point mutations) that is greater than 1:1. In some embodiments, any of the base editors provided herein are capable of generating a ratio of intended mutations to unintended mutations (*e.g.*, intended point mutations:unintended point mutations) that is at least 1.5:1, at least 2:1, at least 2.5:1, at least 3:1, at least 3.5:1, at least 4:1, at least 4.5:1, at least 5:1, at least 5.5:1, at least 6:1, at least 6.5:1, at least 7:1, at least 7.5:1, at least 8:1, at least 10:1, at least 12:1, at least 15:1, at least 20:1, at least 25:1, at least 30:1, at least 40:1, at least 50:1, at least 100:1, at least 150:1, at least 200:1, at least 250:1, at least 500:1, or at least 1000:1, or more. It should be appreciated that the characteristics of the base editors described in the “*Base Editor Efficiency*” section, herein, may be applied to any of the fusion proteins, or methods of using the fusion proteins provided herein.

Methods for Editing Nucleic Acids

[00277] Some aspects of the disclosure provide methods for editing a nucleic acid. In some embodiments, the method is a method for editing a nucleobase of a nucleic acid (*e.g.*, a

base pair of a double-stranded DNA sequence). In some embodiments, the method comprises the steps of: a) contacting a target region of a nucleic acid (*e.g.*, a double-stranded DNA sequence) with a complex comprising a base editor (*e.g.*, a Cas9 domain fused to a cytidine deaminase domain) and a guide nucleic acid (*e.g.*, gRNA), wherein the target region comprises a targeted nucleobase pair, b) inducing strand separation of said target region, c) converting a first nucleobase of said target nucleobase pair in a single strand of the target region to a second nucleobase, and d) cutting no more than one strand of said target region, where a third nucleobase complementary to the first nucleobase base is replaced by a fourth nucleobase complementary to the second nucleobase; and the method results in less than 20% indel formation in the nucleic acid. It should be appreciated that in some embodiments, step b is omitted. In some embodiments, the first nucleobase is a cytosine. In some embodiments, the second nucleobase is a deaminated cytosine, or a uracil. In some embodiments, the third nucleobase is a guanine. In some embodiments, the fourth nucleobase is an adenine. In some embodiments, the first nucleobase is a cytosine, the second nucleobase is a deaminated cytosine, or a uracil, the third nucleobase is a guanine, and the fourth nucleobase is an adenine. In some embodiments, the method results in less than 19%, 18%, 16%, 14%, 12%, 10%, 8%, 6%, 4%, 2%, 1%, 0.5%, 0.2%, or less than 0.1% indel formation. In some embodiments, the method further comprises replacing the second nucleobase with a fifth nucleobase that is complementary to the fourth nucleobase, thereby generating an intended edited base pair (*e.g.*, C:G → T:A). In some embodiments, the fifth nucleobase is a thymine. In some embodiments, at least 5% of the intended basepairs are edited. In some embodiments, at least 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, or 50% of the intended basepairs are edited.

[00278] In some embodiments, the ratio of intended products to unintended products in the target nucleotide is at least 2:1, 5:1, 10:1, 20:1, 30:1, 40:1, 50:1, 60:1, 70:1, 80:1, 90:1, 100:1, or 200:1, or more. In some embodiments, the ratio of intended point mutation to indel formation is greater than 1:1, 10:1, 50:1, 100:1, 500:1, or 1000:1, or more. In some embodiments, the cut single strand (nicked strand) is hybridized to the guide nucleic acid. In some embodiments, the cut single strand is opposite to the strand comprising the first nucleobase. In some embodiments, the base editor comprises a Cas9 domain. In some embodiments, the first base is cytosine, and the second base is not a G, C, A, or T. In some embodiments, the second base is uracil. In some embodiments, the first base is cytosine. In some embodiments, the second base is not a G, C, A, or T. In some embodiments, the second base is uracil. In some embodiments, the base editor inhibits base excision repair of the

edited strand. In some embodiments, the base editor protects or binds the non-edited strand. In some embodiments, the base editor comprises UGI activity. In some embodiments, the base editor comprises nickase activity. In some embodiments, the intended edited basepair is upstream of a PAM site. In some embodiments, the intended edited base pair is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides upstream of the PAM site. In some embodiments, the intended edited basepair is downstream of a PAM site. In some embodiments, the intended edited base pair is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides downstream stream of the PAM site. In some embodiments, the method does not require a canonical (*e.g.*, NGG) PAM site. In some embodiments, the nucleobase editor comprises a linker. In some embodiments, the linker is 1-25 amino acids in length. In some embodiments, the linker is 5-20 amino acids in length. In some embodiments, linker is 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 amino acids in length. In some embodiments, the target region comprises a target window, wherein the target window comprises the target nucleobase pair. In some embodiments, the target window comprises 1-10 nucleotides. In some embodiments, the target window is 1-9, 1-8, 1-7, 1-6, 1-5, 1-4, 1-3, 1-2, or 1 nucleotides in length. In some embodiments, the target window is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides in length. In some embodiments, the intended edited base pair is within the target window. In some embodiments, the target window comprises the intended edited base pair. In some embodiments, the method is performed using any of the base editors provided herein. In some embodiments, a target window is a deamination window

[00279] In some embodiments, the disclosure provides methods for editing a nucleotide. In some embodiments, the disclosure provides a method for editing a nucleobase pair of a double-stranded DNA sequence. In some embodiments, the method comprises a) contacting a target region of the double-stranded DNA sequence with a complex comprising a base editor and a guide nucleic acid (*e.g.*, gRNA), where the target region comprises a target nucleobase pair, b) inducing strand separation of said target region, c) converting a first nucleobase of said target nucleobase pair in a single strand of the target region to a second nucleobase, d) cutting no more than one strand of said target region, wherein a third nucleobase complementary to the first nucleobase base is replaced by a fourth nucleobase complementary to the second nucleobase, and the second nucleobase is replaced with a fifth nucleobase that is complementary to the fourth nucleobase, thereby generating an intended edited basepair, wherein the efficiency of generating the intended edited basepair is at least 5%. It should be appreciated that in some embodiments, step b is omitted. In some

embodiments, at least 5% of the intended basepairs are edited. In some embodiments, at least 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, or 50% of the intended basepairs are edited. In some embodiments, the method causes less than 19%, 18%, 16%, 14%, 12%, 10%, 8%, 6%, 4%, 2%, 1%, 0.5%, 0.2%, or less than 0.1% indel formation. In some embodiments, the ratio of intended product to unintended products at the target nucleotide is at least 2:1, 5:1, 10:1, 20:1, 30:1, 40:1, 50:1, 60:1, 70:1, 80:1, 90:1, 100:1, or 200:1, or more. In some embodiments, the ratio of intended point mutation to indel formation is greater than 1:1, 10:1, 50:1, 100:1, 500:1, or 1000:1, or more. In some embodiments, the cut single strand is hybridized to the guide nucleic acid. In some embodiments, the cut single strand is opposite to the strand comprising the first nucleobase. In some embodiments, the first base is cytosine. In some embodiments, the second nucleobase is not G, C, A, or T. In some embodiments, the second base is uracil. In some embodiments, the base editor inhibits base excision repair of the edited strand. In some embodiments, the base editor protects or binds the non-edited strand. In some embodiments, the nucleobase editor comprises UGI activity. In some embodiments, the nucleobase edit comprises nickase activity. In some embodiments, the intended edited basepair is upstream of a PAM site. In some embodiments, the intended edited base pair is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides upstream of the PAM site. In some embodiments, the intended edited basepair is downstream of a PAM site. In some embodiments, the intended edited base pair is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides downstream stream of the PAM site. In some embodiments, the method does not require a canonical (e.g., NGG) PAM site. In some embodiments, the nucleobase editor comprises a linker. In some embodiments, the linker is 1-25 amino acids in length. In some embodiments, the linker is 5-20 amino acids in length. In some embodiments, the linker is 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 amino acids in length. In some embodiments, the target region comprises a target window, wherein the target window comprises the target nucleobase pair. In some embodiments, the target window comprises 1-10 nucleotides. In some embodiments, the target window is 1-9, 1-8, 1-7, 1-6, 1-5, 1-4, 1-3, 1-2, or 1 nucleotides in length. In some embodiments, the target window is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides in length. In some embodiments, the intended edited base pair occurs within the target window. In some embodiments, the target window comprises the intended edited base pair. In some embodiments, the nucleobase editor is any one of the base editors provided herein.

[00280]

Kits, vectors, cells

[00281] Some aspects of this disclosure provide kits comprising a nucleic acid construct, comprising (a) a nucleotide sequence encoding a Cas9 protein or a Cas9 fusion protein as provided herein; and (b) a heterologous promoter that drives expression of the sequence of (a). In some embodiments, the kit further comprises an expression construct encoding a guide RNA backbone, wherein the construct comprises a cloning site positioned to allow the cloning of a nucleic acid sequence identical or complementary to a target sequence into the guide RNA backbone.

[00282] Some aspects of this disclosure provide polynucleotides encoding a Cas9 protein or a fusion protein as provided herein. Some aspects of this disclosure provide vectors comprising such polynucleotides. In some embodiments, the vector comprises a heterologous promoter driving expression of polynucleotide.

[00283] Some aspects of this disclosure provide cells comprising a Cas9 protein, a fusion protein, a nucleic acid molecule encoding the fusion protein, a complex comprising the Cas9 protein and the gRNA, and/or a vector as provided herein.

[00284] The description of exemplary embodiments of the reporter systems above is provided for illustration purposes only and not meant to be limiting. Additional reporter systems, e.g., variations of the exemplary systems described in detail above, are also embraced by this disclosure.

EXAMPLES

EXAMPLE 1: Cas9 Deaminase Fusion Proteins

[00285] A number of Cas9:Deaminase fusion proteins were generated and deaminase activity of the generated fusions was characterized. The following deaminases were tested:

Human AID (hAID):

MDSLLMNRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNKC
 HVLLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGNPYLSLRIFTAR
 LYFCEDRKAEPGLRRLHRAGVQIAIMTFKDYFCWNTFVENHERTFKAWEWGLHEN
 SVRLSRQLRRILLPLYEVDDLRFDAFRTLGLLD (SEQ ID NO: 607)

Human AID-DC (hAID-DC, truncated version of hAID with 7-fold increased activity):

MDSLLMNRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNKC
 HVLLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGNPYLSLRIFTAR
 LYFCEDRKAEPGLRRLHRAGVQIAIMTFKDYFCWNTFVENHERTFKAWEWGLHEN

SVRLSRQLRRILL (SEQ ID NO: 608)

Rat APOBEC1 (rAPOBEC1):

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRHSIWRHTSQNT
 NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWPYPHPLW
 VRLYVLELYCIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLK (SEQ
 ID NO: 284)

Human APOBEC1 (hAPOBEC1)

MTSEKGPSTGDPTLRRRIEPWEFDVVFYDPRELKREACCLLYEIKWGMRSRKIWRSSGKN
 TTNHVEVNFIEKFTSERDFHPSMSCSITWFLSWSPCWECQAIREFLSRHPGVTLVIYV
 ARLFWHMDQQRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYP
 PLWMMLYALELHCILSLPPCLKISRRWQNHLLTFFRLHLQNCHYQTIPPHILLATGLIH
 PSVAWR (SEQ ID NO: 5724)

Petromyzon marinus (Lamprey) CDA1 (pmCDA1):

MTDAEYVRIHEKLDIYTFKKQFFNKKSVSHRCYVLFELKRRGERRACFWGYAVNK
 PQSGTERGIIHAEIFSIRKVEEYLRDNPQGFTINWYSSWSPCADCAEKILEWYNQELRG
 NGHTLKIWACKLYYEKNARNQIGLWNLRDNGVGLNVMVSEHYQCCRKIFIQSSHNQ
 LNENRWLEKTLKRAEKRRSELSIMIQVKILHTTKSPAV (SEQ ID NO: 609)

Human APOBEC3G (hAPOBEC3G):

MELKYHPEMRFFHWFSKWRKLRDQEYEV TWYISWSPCTKCTRDMATFLAEDPKV
 TLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKIMNYDEFQHCWSKFFVYSQRE
 LFEPWNNLPKYYILLHIMLGEILRHSMDPPTFTFNFNNEPWVRGRHETLYCYEVERM
 HNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFLDVIPFWKLDLDQDYRVTCTF
 SWSPCFSCAQEMAKFISKNKHVSLCIFTARIYDDQGRCEGLRTLAEAGAKISIMTYS
 EFKHCWDTFVDHQGCPFQPWDGLDEHSQDLSGRLRAILQNQEN (SEQ ID NO: 610)

[00286] Deaminase Activity on ssDNA. A USER (Uracil-Specific Excision Reagent)

Enzyme-based assay for deamination was employed to test the activity of various deaminases on single-stranded DNA (ssDNA) substrates. USER Enzyme was obtained from New England Biolabs. An ssDNA substrate was provided with a target cytosine residue at different positions. Deamination of the ssDNA cytosine target residue results in conversion of the target cytosine to a uracil. The USER Enzyme excises the uracil base and cleaves the ssDNA backbone at that position, cutting the ssDNA substrate into two shorter fragments of DNA. In some assays, the ssDNA substrate is labeled on one end with a dye, *e.g.*, with a 5' Cy3 label (the * in the scheme below). Upon deamination, excision, and cleavage of the strand, the substrate can be subjected to electrophoresis, and the substrate and any fragment released from it can be visualized by detecting the label. Where Cy5 is images, only the fragment with the label will be visible via imaging.

[00287] In one USER Enzyme assay, ssDNA substrates were used that matched the target sequences of the various deaminases tested. Expression cassettes encoding the deaminases

tested were inserted into a CMV backbone plasmid that has been used previously in the lab (Addgene plasmid 52970). The deaminase proteins were expressed using a TNT Quick Coupled Transcription/Translation System (Promega) according to the manufacturers recommendations. After 90 min of incubation, 5 mL of lysate was incubated with 5' Cy3-labeled ssDNA substrate and 1 unit of USER Enzyme (NEB) for 3 hours. The DNA was resolved on a 10% TBE PAGE gel and the DNA was imaged using Cy-dye imaging. A schematic representation of the USER Enzyme assay is shown in Figure 41.

[00288] **Figure 1** shows the deaminase activity of the tested deaminases on ssDNA substrates, such as Doench 1, Doench 2, G7' and VEGF Target 2. The rAPOBEC1 enzyme exhibited a substantial amount of deamination on the single-stranded DNA substrate with a canonical NGG PAM, but not with a negative control non-canonical NNN PAM.

Cas9 fusion proteins with APOBEC family deaminases were generated. The following fusion architectures were constructed and tested on ssDNA:

rAPOBEC1-GGG-*dCas9* primary sequence

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELRKETCLLYEINWGGRHSIWRHTSQNT
NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWPYRPHLW
VRLYVLELYCIILGLPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLKGGSD
KKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIGALLFDSGETAE
ATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFHRLVESFLVEEDKKHERHP
IFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDLN
PDNSVDKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGEK
KNGLFGNLIASLGLTPNFKSNFDLAEDAKLQLSKD TYDDDLDNLLAQIGDQYADLF
LAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKYK
EIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTF
DNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSRFA
WMTRKSEETITPWNFEEVVDKGASAQSFIERMNTNFDKNLPNEKVLPHKSLLEYEFT
VYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFD
SVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEE
RLKTYAHLFDDKVMKQLKRRRYTGWRLSRKLINGIRDKQSGKTILDFLKSDFANR
NFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGILQTVKVVDELV
KVMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIE EGIKELGSQILKEHPVENTQ
LQNEKLYLYLQNGRDMYVDQELDINRLSDYDVDAIVPQSFLKDDSIDNKVLTRSDK
NRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGELSELDKAGFIK
RQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKVR
EINNYHHAHDAYLNAVVG TALIKKYPKLESEFVYGDYKVYDVRKMIKSEQEIGKAT
AKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVL SMP
QVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKGFFDSPTVAYSVLVVAK
VEKGKSKKLKSVKELGITIMERSSEFEKNPIDFLEAKGYKEVKKDLIILPKYSLFELE
NGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQHK
HYLDEIIEQISEFSKRVLADANLDKVL SAYNKH RDKPIREQAENIIHLFTLTNLGAPA
AFKYFDTTIDRKRYTSTKEVLDATLIHQ SITGLYETRIDLSQLGGD (SEQ ID NO: 611)

rAPOBEC1-(GGG)₃-dCas9 primary sequence

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELRKETCLLYEINWGGRHSIWRHTSQNT
NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWPYPHPLW
VRLYVLELYCIIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPAPHILWATGLKGGSG
GSGGSMDKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIGALLF
DSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRLEESFLVEED
KKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGH
FLIEGDLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIA
AQLPGEKKNGLFGNLIASLGLTPNFKSNFDLAEDAQLQSKDQYDDDLNLLAQIG
DQYADLFLAAKNLSAAILSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQ
QLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKNREDLL
RKQRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLAR
GNSRFAWMTRKSEETITPWNFEEVVDKGGASAQSFIERMNTNFDKNLPNEKVLPHKSL
LYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFK
KIECFDSVEISGVEDRFNASLGTYHDLKIKDKDFLDNEENEDILEDIVLTLTLFEDR
EMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSD
GFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKILQTVKV
VDELVKVMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIEEGIKELGSQILKEHP
VENTQLQNEKLYLYLQNGRDMYVDQELDINRLSDYDVAIVPQSFLKDDSIDNKVL
TRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDK
AGFIKQQLVETROITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQ
FYKVINNYHHAHDAYLNAVVGTAIKKYPKLESEFVYGDYKVYDVRKMIKSEQE
IGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKV
LSMPQVNIVKKTQVQGGFSKESILPKRNSDKLIARKKDWDPKKGFFDSPTVAYSVL
VVAKVEKGGKSKKLSVKELLGITIMERSSEFEKNPIDFLEAKGYKEVKKDLIHKLPKYSL
FELENGRKRMLASAGELQKGNELALPSKYVNFYLASHYEKLGSPEDNEQKQLFV
EQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHDKPIREQAENIHLFTLNL
GAPAAFKYFDTTIDRKRYTSTKEVL DATLIHQ SITGLYETRIDLSQLGGD (SEQ ID
NO: 612)

dCas9-GGG-rAPOBEC1

DKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIGALLFDSGETA
EATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRLEESFLVEEDKKHERH
PIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDL
NPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAAQLPGE
KKNGLFGNLIASLGLTPNFKSNFDLAEDAQLQSKDQYDDDLNLLAQIGDQYADL
FLAAKNLSAAILSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKY
KEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKNREDLLRKQRT
FDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSR
AWMTRKSEETITPWNFEEVVDKGGASAQSFIERMNTNFDKNLPNEKVLPHKSLLYEYF
TVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECF
DSVEISGVEDRFNASLGTYHDLKIKDKDFLDNEENEDILEDIVLTLTLFEDREMIE
ERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFAN
RNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKILQTVKVVDEL
VKVMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIEEGIKELGSQILKEHPVENT
QLQNEKLYLYLQNGRDMYVDQELDINRLSDYDVAIVPQSFLKDDSIDNKVLTRSD
KNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDKAGFI

KRQLVETROITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKV
REINNYHHAHDAYLNAVVGTAIIKKYPKLESEFVYGDYKVYDVRKMIKSEQEIGKA
TAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLSP
QVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVVAK
VEKGKSKKLKSVKELGITIMERSSEKPNIDFLEAKGYKEVKKDLIILPKYSLFELE
NGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGKSPEDNEQKQLFVEQHK
HYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIHLFTLTNLGAPA
AFKYFDTTIDRKRYTSTKEVLDATLIHQ SITGLYETRIDLSQLGGDGGSMMSSETGPVA
VDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRHSIWRHTSQNTNKHVEVNF
EKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYHHADPR
NRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYS SPSNEAHWPRYPHLWVRLYVLEL
YCII LGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLK (SEQ ID NO: 613)

dCas9-GGG₃-rAPOBEC1

DKKYSIGLAIGTNSVGVAVITDEYKVPSKFKVLGNTDRHSIKKNLIGALLFDSGETA
EATRLKRTARRRYTRRKNRICYLQEIFS NEMAKVDDSFHRLEESFLVEEDKKHERH
PIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDL
NPDNSVDKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGE
KKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLNLLAQIGDQYADL
FLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKY
KEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKNREDLLRKQRT
FDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYVYVGPLARGNSRF
AWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPKHSLLEYEF
TVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECF
DSVEISGVEDRFNASLGTYHDLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIE
ERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDFAN
RNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKILQTVKVVDEL
VKVMGRHKPENIVEMARENQTTQKGQKNSRERMKRIE EGIKELGSQILKEHPVENT
QLQNEKLYLYLQNGRDMYVDQELDINRLSDYDVAIVPQSFLKDDSIDNKVLTRSD
KNRGKSDNVPSEEVVKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDKAGFI
KRQLVETROITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKV
REINNYHHAHDAYLNAVVGTAIIKKYPKLESEFVYGDYKVYDVRKMIKSEQEIGKA
TAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLSP
QVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVVAK
VEKGKSKKLKSVKELGITIMERSSEKPNIDFLEAKGYKEVKKDLIILPKYSLFELE
NGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGKSPEDNEQKQLFVEQHK
HYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIHLFTLTNLGAPA
AFKYFDTTIDRKRYTSTKEVLDATLIHQ SITGLYETRIDLSQLGGDGGSGSGGSMSS
ETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRHSIWRHTSQNTNKH
VEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYH
HADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYS SPSNEAHWPRYPHLWVRL
YVLEL YCII LGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLK (SEQ ID
 NO: 614)

rAPOBEC1-XTEN-*dCas9* primary sequence

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRHSIWRHTSQNT
NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYS SPSNEAHWPRYPHLW

VRLYVLELYCIILGLPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLKSGSE
TPGTSESATPESDKKYSIGLAIGTNSVGVAVITDEYKVPSSKFKVLGNTDRHSIKKNLI
 GALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSDFFHRLEESF
 LVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIK
 FRGHFLIEGDLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRR
 LENLIAQLPGEKKNGLFGNLIALSGLTPNFKSNFDLAEDAKLQLSKDTYDDDLN
 LAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLK
 ALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLN
 REDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTRIPYYV
 GPLARGNSRFAWMTRKSEETITPWNFEVVVDKGASAQSFIERMTNFDKNLPNEKVL
 PKHSLLEYEFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLK
 EDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIKDKDFLDNEENEDILEDIVLTLT
 LFEDREMIEERLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILD
 FLKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGI
 QTVKVVDELVKVMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIEEGIKELGSQI
 LKEHPVENTQLQNEKLYLYLQNGRDMYVDQELDINRLSDYDVDAIVPOSFLKDDSI
 DNKVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGL
 SELDKAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSKLVSDF
 RKDFQFYKREINNYHHAHDAYLNAVVGTALIKKYPKLESEFVYGDYKVYDVRKMIA
 KSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFA
 TVRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPT
 VAYSVLVVAKVEKGKSKKLKSVKELLGITIMERSSSFEKNPIDFLEAKGYKEVKKDLIK
 LPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLKGSPEDNEQ
 KQLFVEQHKKHYLDEIIEQISEFSKRVILADANLDKVLSAYNKHRDKPIREQAENIIHLF
 TLNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQSITGLYETRIDLSQLGGD (SEQ
 ID NO: 615)

[00289] **Figure 2** shows that the N-terminal deaminase fusions showed significant activity on the single stranded DNA substrates. For this reason, only the N-terminal architecture was chosen for further experiments.

[00290] **Figure 3** illustrates double stranded DNA substrate binding by deaminase-dCas9:sgRNA complexes. A number of double stranded deaminase substrate sequences were generated. The sequences are provided below. The structures according to **Figure 3** are identified in these sequences (36bp: underlined, sgRNA target sequence: bold; PAM: boxed; 21bp: italicized). All substrates were labeled with a 5'-Cy3 label:

- 2: GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGTCCCGCGGATTTATTTATTTAA**TGG**ATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 616)
- 3: GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCTTCCCGCGGATTTATTTATT**TA**TGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 617)
- 4: GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCTTCCCGCGGATTTATTTAT**TA**TGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 618)
- 5: GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCATTCCCGCGGATTTATTTA**TT**TGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 619)
- 6: GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCTATTCCCGCGGATTTATTT**AT**TGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 620)

7:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCTTATTCCGCGGATTTATT
 TATGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 621)
 8:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCATTATTCCGCGGATTTAT
 TTGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 622)
 9:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCTTATTATTCCGCGGATTTA
 TTGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 623)
 10:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCATTATATTCCGCGGATTT
 ATGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 624)
 11:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCTTATTATATTCCGCGGATT
 TATGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 625)
 12:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCTTATTATATTCCGCGGAT
 TTGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 626)
 13:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCATTATTATATTCCGCGGA
 TTGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 627)
 14:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCTTATTATTATATTCCGCGG
 ATGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 628)
 15:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCATTATTATTATTACCGCG
 GATGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 629)
 18:GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCATTATTATTATTATTACC
 GCTGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 630)
 “-“ :
 GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGTAATATTAATTTATTTATTTAA
 TGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 631)
 8U:GTAGGTAGTTAGGATGAATGGAAGGTTGGTGTAGATTATTATCUGCGGATTTA
 TGGATGACCTCTGGATCCATGGACAT-3' (SEQ ID NO: 632)

*In all substrates except for “8U”, the top strand in Figure 3 is the complement of the sequence specified here. In the case of “8U”, there is a “G” opposite the U.

[00291] **Figure 4** shows the results of a double stranded DNA Deamination Assay. The fusions were expressed and purified with an N-terminal His6 tag via both Ni-NTA and sepharose chromatography. In order to assess deamination on dsDNA substrates, the various dsDNA substrates shown on the previous slide were incubated at a 1:8 dsDNA:fusion protein ratio and incubated at 37 °C for 2 hours. Once the dCas9 portion of the fusion binds to the DNA it blocks access of the USER enzyme to the DNA. Therefore, the fusion proteins were denatured following the incubation and the dsDNA was purified on a spin column, followed by incubation for 45 min with the USER Enzyme and resolution of the resulting DNA substrate and substrate fragments on a 10% TBE-urea gel.

[00292] **Figure 5** demonstrates that Cas9 fusions can target positions 3-11 of double-stranded DNA target sequences (numbered according to the schematic in Figure 3). Upper Gel: 1 μM rAPOBEC1-GGS-dCas9, 125 nM dsDNA, 1 eq sgRNA. Mid Gel: 1 μM

rAPOBEC1-(GGG)₃-dCas9, 125 nM dsDNA, 1 eq sgRNA. Lower Gel: 1.85 μM rAPOBEC1-XTEN-dCas9, 125 nM dsDNA, 1 eq sgRNA. Based on the data from these gels, positions 3-11 (according to the numbering in Figure 3) are sufficiently exposed to the activity of the deaminase to be targeted by the fusion proteins tested. Access of the deaminase to other positions is most likely blocked by the dCas9 protein.

[00293] The data further indicates that a linker of only 3 amino acids (GGG) is not optimal for allowing the deaminase to access the single stranded portion of the DNA. The 9 amino acid linker [(GGG)₃] (SEQ ID NO: 596) and the more structured 16 amino acid linker (XTEN) allow for more efficient deamination.

[00294] **Figure 6** demonstrates that the correct guide RNA, *e.g.*, the correct sgRNA, is required for deaminase activity. The gel shows that fusing the deaminase to dCas9, the deaminase enzyme becomes sequence specific (*e.g.*, using the fusion with an eGFP sgRNA results in no deamination), and also confers the capacity to the deaminase to deaminate dsDNA. The native substrate of the deaminase enzyme is ssDNA, and no deamination occurred when no sgRNA was added. This is consistent with reported knowledge that APOBEC deaminase by itself does not deaminate dsDNA. The data indicates that Cas9 opens the double-stranded DNA helix within a short window, exposing single-stranded DNA that is then accessible to the APOBEC deaminase for cytidine deamination. The sgRNA sequences used are provided below. sequences (36bp: underlined, sgRNA target sequence: bold; PAM: boxed; 21bp: italicized)

DNA sequence 8:

5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTATAGCCATTATTCCGCGGATTTATT
TTGGATGACCTCTGGATCCATGGAC-3' (SEQ ID NO: 633)

Correct sgRNA sequence (partial 3' sequence):

5'-**AUUAUCCGCGGAUUUAUUUGUUUUAGAGCUAG**...-3' (SEQ ID NO: 634)

eGFP sgRNA sequence (partial 3'-sequence):

5'-**CGUAGGCCAGGGUGGUCACGGUUUUAGAGCUAG**...-3' (SEQ ID NO: 635)

EXAMPLE 2: Deamination of DNA target sequence

[00295] **Exemplary deamination targets.** The dCas9:deaminase fusion proteins described herein can be delivered to a cell *in vitro* or *ex vivo* or to a subject *in vivo* and can be used to effect C to T or G to A transitions when the target nucleotide is in positions 3-11 with respect to a PAM. Exemplary deamination targets include, without limitation, the following: CCR5

truncations: any of the codons encoding **Q93**, **Q102**, **Q186**, **R225**, **W86**, or **Q261** of CCR5 can be deaminated to generate a STOP codon, which results in a nonfunctional truncation of CCR5 with applications in HIV treatment. APOE4 mutations: mutant codons encoding **C11R** and **C57R** mutant APOE4 proteins can be deaminated to revert to the wild-type amino acid with applications in Alzheimer's treatment. eGFP truncations: any of the codons encoding **Q158**, **Q184**, **Q185** can be deaminated to generate a STOP codon, or the codon encoding **M1** can be deaminated to encode I, all of which result in loss of eGFP fluorescence, with applications in reporter systems. eGFP restoration: a mutant codon encoding **T65A** or **Y66C** mutant GFP, which does not exhibit substantial fluorescence, can be deaminated to restore the wild-type amino acid and confer fluorescence. PIK3CA mutation: a mutant codon encoding **K111E** mutant PIK3CA can be deaminated to restore the wild-type amino acid residue with applications in cancer. CTNNB1 mutation: a mutant codon encoding **T41A** mutant CTNNB1 can be deaminated to restore the wild-type amino acid residue with applications in cancer. HRAS mutation: a mutant codon encoding **Q61R** mutant HRAS can be deaminated to restore the wild-type amino acid residue with applications in cancer. P53 mutations: any of the mutant codons encoding **Y163C**, **Y236C**, or **N239D** mutant p53 can be deaminated to encode the wild type amino acid sequence with applications in cancer.

The feasibility of deaminating these target sequences in double-stranded DNA is demonstrated in **Figures 7 and 8**. **Figure 7** illustrates the mechanism of target DNA binding of *in vivo* target sequences by deaminase-dCas9:sgRNA complexes.

[00296] **Figure 8** shows successful deamination of exemplary disease-associated target sequences. Upper Gel: CCR5 Q93: coding strand target in pos. 10 (potential off-targets at positions 2, **5**, 6, 8, 9); CCR5 Q102: coding strand target in pos. **9** (potential off-targets at positions 1, 12, 14); CCR5 Q186: coding strand target in pos. 9 (potential off-targets at positions 1, **5**, 15); CCR5 R225: coding strand target in pos. **6** (no potential off-targets); eGFP Q158: coding strand target in pos. **5** (potential off-targets at positions 1, 13, 16); eGFP Q184 /185: coding strand target in pos. **4** and **7** (potential off-targets at positions 3, 12, 14, 15, 16, 17, 18); eGFP M1: template strand target in pos. 12 (potential off-targets at positions 2, **3**, 7, 9, 11) (targets positions 7 and 9 to small degree); eGFP T65A: template strand target in pos. 7 (potential off-targets at positions 1, **8**, 17); PIK3CA K111E: template strand target in pos. 2 (potential off-targets at positions **5**, **8**, 10, 16, 17); PIK3CA K111E: template strand target in pos. 13 (potential off-targets at positions 11, 16, 19) **X**. Lower Gel: CCR5 W86: template strand target in pos. 2 and 3 (potential off-targets at positions 1, 13) **X**; APOE4 C11R: coding strand target in pos. 11 (potential off-targets at positions 7, 13, 16, 17); APOE4 C57R: coding

strand target in pos. 5) (potential off-targets at positions 7, 8, 12); eGFP Y66C: template strand target in pos. 11 (potential off-targets at positions 1, 4, 6, 8, 9, 16); eGFP Y66C: template strand target in pos. 3 (potential off-targets at positions 1, 8, 17); CCR5 Q261: coding strand target in pos. 10 (potential off-targets at positions 3, 5, 6, 9, 18); CTNNB1 T41A: template strand target in pos. 7 (potential off-targets at positions 1, 13, 15, 16) X; HRAS Q61R: template strand target in pos. 6 (potential off-targets at positions 1, 2, 4, 5, 9, 10, 13); p53 Y163C: template strand target in pos. 6 (potential off-targets at positions 2, 13, 14); p53 Y236C: template strand target in pos. 8 (potential off-targets at positions 2, 4); p53 N239D: template strand target in pos. 4 (potential off-targets at positions 6, 8). Exemplary DNA sequences of disease targets are provided below (PAMs (5'-NGG-3') and target positions are boxed):

CCR5 Q93: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTAACTATGCTGCCGCC

CAGTGGGACTT**TGG**AAATACAATGTGTCAACTCTT-3' (SEQ ID NO: 636)

CCR5 Q102: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTAAAATACAATGTGT

CAACTCTTGACA**GGG**CTCTATTTTATAGGCTTCTTC-3' (SEQ ID NO: 637)

CCR5 Q186: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTATTTTCCATACAGT

CAGTATCAATTC**TGG**AAGAATTTCCAGACATTAAAG-3' (SEQ ID NO: 638)

CCR5 R225: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTAGCTTCGGTGT**C**GA

AATGAGAAGAAG**AGG**CACAGGGCTGTGAGGCTTATC-3' (SEQ ID NO: 639)

CCR5 W86: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTAGTGAGC**CC**AGAAGG

GGACAGTAAGA**AGG**AAAAACAGGTCAGAGATGGCC-3' (SEQ ID NO: 640)

CCR5 Q261: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTATCCTGAACACCTT

CCAGGAATTC**T**TGG**CCTGAATAATTGCAGTAGCTC**-3' (SEQ ID NO: 641)

APOE4 C11R: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTAGACATGGAGGAC

GTG**C**GCGGCC**CGCC****TGG**TGCAGTACCGCGGCGAGGTGC-3' (SEQ ID NO: 642)

APOE4 C57R: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTACTGCAGAAG**C**GC

CTGGCAGTGTACC**AGG**CCGGGGCCCGCGAGGGCGCCG-3' (SEQ ID NO: 643)

eGFP Q158: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTAGCCGACAAG**C**AGA

AGAACGGCATCA**AGG**TGAACTTCAAGATCCGCCACA-3' (SEQ ID NO: 644)

eGFP Q184/185: 5'-Cy3-GTAGGTAGTTAGGATGAATGGAAGGTTGGTAACTACTAC**C**

AG**C**AGAACACCCCAT**CGG**CGACGGCCCCGTGCTGCTGCC-3' (SEQ ID NO: 645)

eGFP M1: 5'-Cy3-

GTAGGTAGTTAGGATGAATGGAAGGTTGGTACCTCGCCCTTGCTCA

CATCTCGAGT**CGG**CCGCCAGTGTGATGGATATCT-3' (SEQ ID NO: 646)

eGFP T65A: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTACACGCGTAGGCA
GGGTGGTCACGAGGGTGGGCCAGGGCACGGGCAGC-3' (SEQ ID NO: 647)
eGFP Y66C: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTAAAGCACTGCACTC
CGAGGTCAGGGTGGTCACGAGGGTTGGCCAGGGCA-3' (SEQ ID NO: 648)
eGFP Y66C: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTACACTCCGAGGTC
AGGGTGGTCACGAGGGTTGGCCAGGGCACGGGCAGG-3' (SEQ ID NO: 649)
PIK3CA K111E: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTAGGATCTTTC
TTCACGGTTGCCTACTTGGTTCAATTACTTTTAAAAATGG-3' (SEQ ID NO: 650)
PIK3CA K111E: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTATTCTCGATTG
AGGATCTTTCTTCACGGTTGCTACTGGTTCAATTACT-3' (SEQ ID NO: 651)
CTNNB1 T41A: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTAAGGAGCTGTGG
CAGTGGCACCAGAATGGATTCCAGAGTCCAGGTAAGAC-3' (SEQ ID NO: 652)
HRAS Q61R: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTAGTACTCCTCCCGG
CCGGCGGTATCCAGGATGTCCAACAGGCACGTCTCC-3' (SEQ ID NO: 653)
p53 Y163C: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTATGACTGCTTGCAG
ATGGCCATGGCGCGGACGCGGGTGCCGGGCGGGGGT-3' (SEQ ID NO: 654)
p53 Y236C: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTACTGTTACACATGC
AGTTGTAGTGGAATGGTGGTACAGTCAGAGCCAACCT-3' (SEQ ID NO: 655)
p53 N239D: 5'-Cy3-
GTAGGTAGTTAGGATGAATGGAAGGTTGGTAGGAACTGTCACAC
ATGTAGTTGTAGTGGATGGTGGTACAGTCAGAGCCA-3' (SEQ ID NO: 656)

EXAMPLE 3: Uracil Glycosylase Inhibitor Fusion Improves Deamination Efficiency

[00297] Direct programmable nucleobase editing efficiencies in mammalian cells by dCas9:deaminase fusion proteins can be improved significantly by fusing a uracil glycosylase inhibitor (UGI) to the dCas9:deaminase fusion protein.

[00298] Figure 9 shows *in vitro* C→T editing efficiencies in human HEK293 cells using rAPOBEC1-XTEN-dCas9:

rAPOBEC1-XTEN-dCas9-NLS primary sequence

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKTCCLLYEINWGGRHSIWRHTSQNT
NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWPYPHFLWV
RLYVLELYCIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPAPHILWATGLKSGSETP
GTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIG
ALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFLV

EEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLLIYLALAHMIKFR
 GHFLIEGDLNPDNSVDKLFQQLVQTYNQLFEENPINASGVDAKAILSARLSKSRRL
 NLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLNLLA
 QIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALV
 RQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKNRED
 LLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFKLDNREKIEKILTRIPYYVGPLAR
 GNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPKHSLI
 YEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFK
 KIECFDSVEISGVEDRFNASLGTYHDLKIIKDKDFLDNEENEDILEDIVLTLTLFEDRE
 MIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDG
 FANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGILQTVKVVD
 ELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEDIKELGSQILKEHPVE
 NTQLQNEKLYLYYLQNGRDMYVDQELDINRLSDYDVAIVPQSFLKDDSIDNKVLT
 RSDKNRGKSDNPSEEVVKMKKNYWRQLLNAKLITQRKFDNLTKAERGGSELK
 AGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQ
 FYKVRINNYHHAHDAYLNAVVGTAIHKYPKLESEFVYGDYKVDVVRKMIKSEQ
 EIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKV
 LMPQVNVKKTETVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSV
 LVVAKVEKGGKSKKLKSVKELLGITIMERSSSFENPIDFLEAKGYKEVKKDLIILPKYS
 LFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLF
 VEQHKHYLDEIIEQISEFSKRVIADANLDKVL SAYNKHRDKPIREQAENIHLFTLTNL
 GAPAAFKYFDTTIDRKRYTSTKEVLDTLIHQISITGLYETRIDLSQLGGDSSGGSPKKKR
 KV (SEQ ID NO: 657)

Protospacer sequences were as follows:

- EMX1: 5'-GAGTC₅**C**₆GAGC₁₀AGAAGAAGAA**GGG**-3' (SEQ ID NO: 293)
- FANCF: 5'-GGAATC₆**C**₇**C**₈TTC₁₁TGCAGCACC**TGG**-3' (SEQ ID NO: 294)
- HEK293 site 2: 5'-GAAC₄**AC**₆AAAGC₁₁ATAGACTGC**GGG**-3' (SEQ ID NO: 295)
- HEK293 site 3: 5'-GGC₃**C**₄**C**₅AGAC₉TGAGCACGTGA**TGG**-3' (SEQ ID NO: 296)
- HEK293 site 4: 5'-GGC₃**AC**₅TGC₈GGC₁₁TGGAGGTGG**GGG**-3' (SEQ ID NO: 297)
- RNF2: 5'-GTC₃ATC₆TTAGTCATTACCTG**AGG**-3' (SEQ ID NO: 298)

*PAMs are boxed, C residues within target window (positions 3-11) are numbered and bolded.

[00299] **Figure 10** demonstrates that C→T editing efficiencies on the same protospacer sequences in HEK293T cells are greatly enhanced when a UGI domain is fused to the rAPOBEC1:dCas9 fusion protein.

rAPOBEC1-XTEN-dCas9-UGI-NLS primary sequence

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRHSIWRHTSQNT
 NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSPTSNEAHWPRYPHLWV
 RLYVLELYCIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLKSGSETP

GTSESATPESDKKYSIGLAIGTNSVGWAVITDEYK VPSKKFKVLGNTDRHSIKKNLIG
ALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRLEESFLV
EEDKKHERHPIFGNIVDEVAYHEK YPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFR
GHFLIEGDLNPDNSDVDFKLFQQLVQTYNQLFEENPINASGVDAKAILSARLSKSRRL
NLIAQLPGEKKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLDNLLA
QIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALV
RQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKNRED
LLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTRIPYYVGPLAR
GNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPKHSL
YEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFK
KIECFDSVEISGVEDRFNASLGTYHDLKIIKDKDFLDNEENEDILEDIVLTLTFEDRE
MIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKS DG
FANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGILQTVK VVD
ELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEDIKELGSQILKEHPVE
NTQLQNEKLYLYYLQNGRDMYVDQELDINRLSDYDVAIVPQSFLKDDSIDNKVLT
RSKDNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGSEL
AGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLTKSKLVSDFRKDFQ
FYK VREINNYHHAHDAYLNAVVG TALIKKYPKLESEFVYGDYK VYDVRKMIAKSEQ
EIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKV
LSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSV
LVVAKVEK GKSKKLSVKELGITIMERS SFEKNPIDFLEAKGYKEVKKDLIIKLPKYS
LFELENGRKRMLASAGELQKGNELALPSKYVNFYLYLASHYEK LKGSPEDNEQKQLF
VEQHKKHYLDEIIEQISEFSKRVLADANLDK VLSAYNKHRDKPIREQAENIIHLFTLTNL
GAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQ SITGLYETRIDLSQLGGD**SGGSTNLSD**
IEKETGKQLVIOESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAP
EYKPWALVIOQDSNGENKIKMLS**SGGSPKKRKV** (SEQ ID NO: 658)

[00300] The percentages in Figures 9 and 10 are shown from sequencing both strands of the target sequence. Because only one of the strands is a substrate for deamination, the maximum possible deamination value in this assay is 50%. Accordingly, the deamination efficiency is double the percentages shown in the tables. *E.g.*, a value of 50% relates to deamination of 100% of double-stranded target sequences.

When a uracil glycosylase inhibitor (UGI) was fused to the dCas9:deaminase fusion protein (*e.g.*, rAPOBEC1-XTEN-dCas9-[UGI]-NLS), a significant increase in editing efficiency in cells was observed. This result indicates that in mammalian cells, the DNA repair machinery that cuts out the uracil base in a U:G base pair is a rate-limiting process in DNA editing. Tethering UGI to the dVas9:deaminase fusion proteins greatly increases editing yields.

[00301] Without UGI, typical editing efficiencies in human cells were in the ~2-14% yield range (Figure 9 and Figure 10, “XTEN” entries). With UGI (Figure 10, “UGI” entries) the editing was observed in the ~6-40% range. Using a UGI fusion is thus more efficient than the current alternative method of correcting point mutations via HDR, which also creates an

excess of indels in addition to correcting the point mutation. No indels resulting from treatment with the cas9:deaminase:UGI fusions were observed.

EXAMPLE 4: Direct, programmable conversion of a target nucleotide in genomic DNA without double-stranded DNA cleavage

[00302] Current genome-editing technologies introduce double-stranded DNA breaks at a target locus of interest as the first step to gene correction.^{39,40} Although most genetic diseases arise from mutation of a single nucleobase to a different nucleobase, current approaches to revert such changes are very inefficient and typically induce an abundance of random insertions and deletions (indels) at the target locus as a consequence of the cellular response to double-stranded DNA breaks.^{39,40} Reported herein is the development of nucleobase editing, a new strategy for genome editing that enables the direct conversion of one target nucleobase into another in a programmable manner, without requiring double-stranded DNA backbone cleavage. Fusions of CRISPR/Cas9 were engineered and the cytidine deaminase enzyme APOBEC1 that retain the ability to be programmed with a guide RNA, do not induce double-stranded DNA breaks, and mediate the direct conversion of cytidine to uracil, thereby effecting a C→T (or G→A) substitution following DNA replication, DNA repair, or transcription if the template strand is targeted. The resulting “nucleobase editors” convert cytidines within a window of approximately five nucleotides, and can efficiently correct a variety of point mutations relevant to human disease *in vitro*. In four transformed human and murine cell lines, second- and third-generation nucleobase editors that fuse uracil glycosylase inhibitor (UGI), and that use a Cas9 nickase targeting the non-edited strand, respectively, can overcome the cellular DNA repair response to nucleobase editing, resulting in permanent correction of up to 37% or (~15-75%) of total cellular DNA in human cells with minimal (typically ≤ 1%) indel formation. In contrast, canonical Cas9-mediated HDR on the same targets yielded an average of 0.7% correction with 4% indel formation. Nucleobase editors were used to revert two oncogenic *p53* mutations into wild-type alleles in human breast cancer and lymphoma cells, and to convert an Alzheimer’s Disease associated Arg codon in *ApoE4* into a non-disease-associated Cys codon in mouse astrocytes. Base editing expands the scope and efficiency of genome editing of point mutations.

[00303] The clustered regularly interspaced short palindromic repeat (CRISPR) system is a prokaryotic adaptive immune system that has been adapted to mediate genome engineering in a variety of organisms and cell lines.⁴¹ CRISPR/Cas9 protein-RNA complexes localize to a

target DNA sequence through base pairing with a guide RNA, and natively create a DNA double-stranded break (DSB) at the locus specified by the guide RNA. In response to DSBs, endogenous DNA repair processes mostly result in random insertions or deletions (indels) at the site of DNA cleavage through non-homologous end joining (NHEJ). In the presence of a homologous DNA template, the DNA surrounding the cleavage site can be replaced through homology-directed repair (HDR). When simple disruption of a disease-associated gene is sufficient (for example, to treat some gain-of-function diseases), targeted DNA cleavage followed by indel formation can be effective. For most known genetic diseases, however, correction of a point mutation in the target locus, rather than stochastic disruption of the gene, is needed to address or study the underlying cause of the disease.⁶⁸

[00304] Motivated by this need, researchers have invested intense effort to increase the efficiency of HDR and suppress NHEJ. For example, a small-molecule inhibitor of ligase IV, an essential enzyme in the NHEJ pathway, has been shown to increase HDR efficiency.^{42,43} However, this strategy is challenging in post-mitotic cells, which typically down-regulate HDR, and its therapeutic relevance is limited by the potential risks of inhibiting ligase IV in non-target cells. Enhanced HDR efficiency can also be achieved by the timed delivery of Cas9-guide RNA complexes into chemically synchronized cells, as HDR efficiency is highly cell-cycle dependent.⁴⁴ Such an approach, however, is limited to research applications in cell culture since synchronizing cells is highly disruptive. Despite these developments, current strategies to replace point mutations using HDR in most contexts are very inefficient (typically ~0.1 to 5%),^{42,43,45,46,75} especially in unmodified, non-dividing cells. In addition, HDR competes with NHEJ during the resolution of double-stranded breaks, and indels are generally more abundant outcomes than gene replacement. These observations highlight the need to develop alternative approaches to install specific modifications in genomic DNA that do not rely on creating double-stranded DNA breaks. A small-molecule inhibitor of ligase IV, an essential enzyme in the NHEJ pathway, has been shown to increase HDR efficiency.^{42,43} However, this strategy is challenging in post-mitotic cells, which typically down-regulate HDR, and its therapeutic relevance is limited by the potential risks of inhibiting ligase IV in non-target cells. Enhanced HDR efficiency can also be achieved by the timed delivery of Cas9-guide RNA complexes into chemically synchronized cells, as HDR efficiency is highly cell-cycle dependent.⁴⁴ Such an approach, however, is limited to research applications in cell culture since synchronizing cells is highly disruptive. In some cases, it is possible to design HDR templates such that the product of successful HDR contains mutations in the PAM sequence and therefore is no longer a substrate for subsequent Cas9 modification, increasing

the overall yield of HDR products,⁷⁵ although such an approach imposes constraints on the product sequences. Recently, this strategy has been coupled to the use of ssDNA donors that are complementary to the non-target strand and high-efficiency ribonucleoprotein (RNP) delivery to substantially increase the efficiency of HDR, but even in these cases the ratio of HDR to NHEJ outcomes is relatively low (< 2).⁸³

[00305] It was envisioned that direct catalysis of the conversion of one nucleobase to another at a programmable target locus without requiring DNA backbone cleavage could increase the efficiency of gene correction relative to HDR without introducing undesired random indels at the locus of interest. Catalytically dead Cas9 (dCas9), which contains Asp10Ala and His840Ala mutations that inactivate its nuclease activity, retains its ability to bind DNA in a guide RNA-programmed manner but does not cleave the DNA backbone.^{16,47} In principle, conjugation of dCas9 with an enzymatic or chemical catalyst that mediates the direct conversion of one nucleobase to another could enable RNA-programmed nucleobase editing. The deamination of cytosine (C) is catalyzed by cytidine deaminases²⁹ and results in uracil (U), which has the base pairing properties of thymine (T). dCas9 was fused to cytidine deaminase enzymes in order to test their ability to convert C to U at a guide RNA-specified DNA locus. Most known cytidine deaminases operate on RNA, and the few examples that are known to accept DNA require single-stranded DNA.⁴⁸ Recent studies on the dCas9-target DNA complex reveal that at least nine nucleotides of the displaced DNA strand are unpaired upon formation of the Cas9:guide RNA:DNA “R-loop” complex.¹² Indeed, in the structure of the Cas9 R-loop complex the first 11 nucleotides of the protospacer on the displaced DNA strand are disordered, suggesting that their movement is not highly restricted.⁷⁶ It has also been speculated that Cas9 nickase-induced mutations at cytosines in the non-template strand might arise from their accessibility by cellular cytidine deaminase enzymes.⁷⁷ Recent studies on the dCas9-target DNA complex have revealed that at least 26 bases on the non-template strand are unpaired when Cas9 binds to its target DNA sequence.⁴⁹ It was reasoned that a subset of this stretch of single-stranded DNA in the R-loop might serve as a substrate for a dCas9-tethered cytidine deaminase to effect direct, programmable conversion of C to U in DNA (Figure 11A).

[00306] Four different cytidine deaminase enzymes (hAID, hAPOBEC3G, rAPOBEC1, and pmCDA1) were expressed in a mammalian cell lysate-derived *in vitro* transcription-translation system and evaluated for ssDNA deamination. Of the four enzymes, rAPOBEC1 showed the highest deaminase activity under the tested conditions and was chosen for dCas9 fusion experiments (Figure 36A). Although appending rAPOBEC1 to the C-terminus of

dCas9 abolishes deaminase activity, fusion to the N-terminus of dCas9 preserves deaminase activity on ssDNA at a level comparable to that of the unfused enzyme. Four rAPOBEC1-dCas9 fusions were expressed and purified with linkers of different length and composition (Figure 36B), and evaluated each fusion for single guide RNA (sgRNA)-programmed dsDNA deamination *in vitro* (Figures 11A to 11C and Figures 15A to 15D).

Efficient, sequence-specific, sgRNA-dependent C to U conversion was observed *in vitro* (Figures 11A to 11C). Conversion efficiency was greatest using rAPOBEC1-dCas9 linkers over nine amino acids in length. The number of positions susceptible to deamination (the deamination “activity window”) increases with linker length was extended from three to 21 amino acids (Figures 36C to 36F15A to 15D). The 16-residue XTEN linker⁵⁰ was found to offer a promising balance between these two characteristics, with an efficient deamination window of approximately five nucleotides, from positions 4 to 8 within the protospacer, counting the end distal to the protospacer-adjacent motif (PAM) as position 1. The rAPOBEC1-XTEN-dCas9 protein served as the first-generation nucleobase editor (NBE1).

[00307] Elected were seven mutations relevant to human disease that in theory could be corrected by C to T nucleobase editing, synthesized double-stranded DNA 80-mers of the corresponding sequences, and assessed the ability of NBE1 to correct these mutations *in vitro* (Figures 16A to 16B). NBE1 yielded products consistent with efficient editing of the target C, or of at least one C within the activity window when multiple Cs were present, in six of these seven targets *in vitro*, with an average apparent editing efficiency of 44% (Figures 16A to 16B). In the three cases in which multiple Cs were present within the deamination window, evidence of deamination of some or all of these cytosines was observed. In only one of the seven cases tested were substantial yields of edited product observed (Figures 16A to 16B). Although the preferred sequence context for APOBEC1 substrates is reported to be CC or TC,⁵¹ it was anticipated that the increased effective molarity of the deaminase and its single-stranded DNA substrate mediated by dCas9 binding to the target locus may relax this restriction. To illuminate the sequence context generality of NBE1, its ability to edit a 60-mer double-stranded DNA oligonucleotide containing a single fixed C at position 7 within the protospacer was assayed, as well as all 36 singly mutated variants in which protospacer bases 1-6 and 8-13 were individually varied to each of the other three bases. Each of these 37 sequences were treated with 1.9 μ M NBE1, 1.9 μ M of the corresponding sgRNA, and 125 nM DNA for 2 h, similar to standard conditions for *in vitro* Cas9 assays⁵². High-throughput DNA sequencing (HTS) revealed 50 to 80% C to U conversion of targeted strands (25 to 40% of total sequence reads arising from both DNA strands, one of which is not a substrate for

NBE1) (Figure 12A). The nucleotides surrounding the target C had little effect on editing efficiency was independent of sequence context unless the base immediately 5' of the target C is a G, in which case editing efficiency was substantially lower (Figures 12A to 12B). NBE1 activity *in vitro* was assessed on all four NC motifs at positions 1 through 8 within the protospacer (Figures 12A to 12B). In general NBE1 activity on substrates was observed to follow the order $TC \geq CC \geq AC > GC$, with maximum editing efficiency achieved when the target C is at or near position 7. In addition, it was observed that the nucleobase editor is highly processive, and will efficiently convert most of all Cs to Us on the same DNA strand within the 5-base activity window (Figure 17).

[00308] While BE1 efficiently processes substrates in a test tube, in cells a tree of possible DNA repair outcomes determines the fate of the initial U:G product of base editing (Figure 29A). To test the effectiveness of nucleobase editing in human cells, NBE1 codon usage was optimized for mammalian expression, appended a C-terminal nuclear localization sequence (NLS),⁵³ and assayed its ability to convert C to T in human cells on 14Cs in six well-studied target sites throughout the human genome (Figure 37A).⁵⁴ The editable Cs were confirmed within each protospacer *in vitro* by incubating NBE1 with synthetic 80-mers that correspond to the six different genomic sites, followed by HTS (Figures 13A to 13C, Figure 29B and Figure 25). Next, HEK293T cells were transfected with plasmids encoding NBE1 and one of the six target sgRNAs, allowed three days for nucleobase editing to occur, extracted genomic DNA from the cells, and analyzed the loci by HTS. Although C to T editing in cells at the target locus was observed for all six cases, the efficiency of nucleobase editing was 1.1% to 6.3% or 0.8%-7.7% of total DNA sequences (corresponding to 2.2% to 12.6% of targeted strands), a 6.3-fold to 37-fold or 5-fold to 36-fold decrease in efficiency compared to that of *in vitro* nucleobase editing (Figures 13A to 13C, Figure 29B and Figure 25). It was observed that some base editing outside of the typical window of positions 4 to 8 when the substrate C is preceded by a T, which we attribute to the unusually high activity of APOBEC1 for TC substrates.⁴⁸

[00309] It was asked whether the cellular DNA repair response to the presence of U:G heteroduplex DNA was responsible for the large decrease in nucleobase editing efficiency in cells (Figure 29A). Uracil DNA glycosylase (UDG) catalyzes removal of U from DNA in cells and initiates base excision repair (BER), with reversion of the U:G pair to a C:G pair as the most common outcome (Figure 29A).⁵⁵ Uracil DNA glycosylase inhibitor (UGI), an 83-residue protein from *B. subtilis* bacteriophage PBS1, potently blocks human UDG activity ($IC_{50} = 12$ pM).⁵⁶ UGI was fused to the C-terminus of NBE1 to create the second-generation

nucleobase editor NBE2 and repeated editing assays on all six genomic loci. Editing efficiencies in human cells were on average 3-fold higher with NBE2 than with NBE1, resulting in gene conversion efficiencies of up to 22.8% of total DNA sequenced (up to 45.6% of targeted strands) (Figures 13A to 13C and Figure 29B). To test base editing in human cells, BE1 codon usage was optimized for mammalian expression and appended a C-terminal nuclear localization sequence (NLS).⁵³

[00310] Similar editing efficiencies were observed when a separate plasmid overexpressing UGI was co-transfected with NBE1 (Figures 18A to 18H). However, while the direct fusion of UGI to NBE1 resulted in no significant increase in C to T mutations at monitored non-targeted genomic locations, overexpression of unfused UGI detectably increased the frequency of C to T mutations elsewhere in the genome (Figures 18A to 18H). The generality of NBE2-mediated nucleobase editing was confirmed by assessing editing efficiencies on the same six genomic targets in U2OS cells, and observed similar results with those in HEK293T cells (Figure 19). Importantly, NBE2 typically did not result in any detectable indels (Figure 13C and Figure 29C), consistent with the known mechanistic dependence of NHEJ on double-stranded DNA breaks.^{57, 78} Together, these results indicate that conjugating UGI to NBE1 can greatly increase the efficiency of nucleobase editing in human cells.

[00311] The permanence of nucleobase editing in human cells was confirmed by monitoring editing efficiencies over multiple cell divisions in HEK293T cells at two of the tested genomic loci. Genomic DNA was harvested at two time points: three days after transfection with plasmids expressing NBE2 and appropriate sgRNAs, and after passaging the cells and growing them for four additional days (approximately five subsequent cell divisions). No significant change in editing efficiency was observed between the non-passaged cells (editing observed in 4.6% to 6.6% of targeted strands for three different target Cs) and passaged cells (editing observed in 4.6% to 6.4% of targeted strands for the same three target Cs), confirming that the nucleobase edits became permanent following cell division (Figure 20). Indels will on rare occasion arise from the processing of U:G lesions by cellular repair processes, which involve single-strand break intermediates that are known to lead to indels.⁸⁴ Given that several hundred endogenous U:G lesions are generated every day per human cell from spontaneous cytidine deaminase,⁸⁵ it was anticipated that the total indel frequency from U:G lesion repair is unlikely to increase from BE1 or BE2 activity at a single target locus.

[00312] To further increase the efficiency of nucleobase editing in cells, it was anticipated that nicking the non-edited strand may result in a smaller fraction of edited Us being removed

by the cell, since eukaryotic mismatch repair machinery uses strand discontinuity to direct DNA repair to any broken strand of a mismatched duplex (Figure 29A).^{58, 79, 80} The catalytic His residue was restored at position 840 in the Cas9 HNH domain,^{47,59} resulting in the third-generation nucleobase editor NBE3 that nicks the non-edited strand containing a G opposite the targeted C, but does not cleave the target strand containing the C. Because NBE3 still contains the Asp10Ala mutation in Cas9, it does not induce double-stranded DNA cleavage. This strategy of nicking the non-edited strand augmented nucleobase editing efficiency in human cells by an additional 1.4- to 4.8-fold relative to NBE2, resulting in up to 36.3% of total DNA sequences containing the targeted C to T conversion on the same six human genomic targets in HEK293T cells (Figures 13A to 13C and Figure 29B). Importantly, only a small frequency of indels, averaging 0.8% (ranging from 0.2% to 1.6% for the six different loci), was observed from NBE3 treatment (Figure 13C, Figure 29C, and Figure 34). In contrast, when cells were treated with wild-type Cas9, sgRNA, and a single-stranded DNA donor template to mediate HDR at three of these loci C to T conversion efficiencies averaging only 0.7% were observed, with much higher relative indel formation averaging 3.9% (Figures 13A to 13C and Figure 29C). The ratio of allele conversion to NHEJ outcomes averaged >1,000 for BE2, 23 for BE3, and 0.17 for wild-type Cas9 (Fig. 3c). We confirmed the permanence of base editing in human cells by monitoring editing efficiencies over multiple cell divisions in HEK293T cells at the HEK293 site 3 and 4 genomic loci (Figure 38). These results collectively establish that nucleobase editing can effect much more efficient targeted single-base editing in human cells than Cas9-mediated HDR, and with much less (NBE3) or no (NBE2) indel formation.

[00313] Next, the off-target activity of NBE1, NBE2, and NBE3 in human cells was evaluated. The off-target activities of Cas9, dCas9, and Cas9 nickase have been extensively studied (Figures 23 to 24 and 31 to 33).^{54,60-62} Because the sequence preference of rAPOBEC1 has been shown to be independent of DNA bases more than one base from the target C,⁶³ consistent with the sequence context independence observed in Figures 12A to 12B, it was assumed that potential off-target activity of nucleobase editors arises from off-target Cas9 binding. Since only a fraction of Cas9 off-target sites will have a C within the active window for nucleobase editing, off-target nucleobase editing sites should be a subset of the off-target sites of canonical Cas9 variants. For each of the six sites studied, the top ten known Cas9 off-target loci in human cells that were previously determined using the GUIDE-seq method were sequenced (Figures 23 to 27 and 31 to 33).^{54, 61} Detectable off-target nucleobase editing at only a subset (16/34, 47% for NBE1 and NBE2, and 17/34, 50% for

NBE3) of known dCas9 off-target loci was observed. In all cases, the off-target base-editing substrates contained a C within the five-base target window. In general, off-target C to T conversion paralleled off-target Cas9 nuclease-mediated genome modification frequencies (Figures 23 to 27). Also monitored were C to T conversions at 2,500 distinct cytosines surrounding the six on-target and 34 off-target loci tested, representing a total of 14,700,000 sequence reads derived from approximately 1.8×10^6 cells, and observed no detectable increase in C to T conversions at any of these other sites upon NBE1, NBE2, or NBE3 treatment compared to that of untreated cells (Figure 28). Taken together, these findings suggest that off-target substrates of nucleobase editors include a subset of Cas9 off-target substrates, and that nucleobase editors in human cells do not induce untargeted C to T conversion throughout the genome at levels that can be detected by the methods used here. No substantial change was observed in editing efficiency between non-passaged HEK293T cells (editing observed in 1.8% to 2.6% of sequenced strands for the three target Cs with BE2, and 6.2% to 14.3% with BE3) and cells that had undergone approximately five cell divisions after base editing (editing observed in 1.9% to 2.3% of sequenced strands for the same target Cs with BE2, and 6.4% to 14.5% with BE3), confirming that base edits in these cells are durable (Extended Data Fig. 6).

[00314] Finally, the potential of nucleobase editing to correct three disease-relevant mutations in mammalian cells was tested. The apolipoprotein E gene variant *APOE4* encodes two Arg residues at amino acid positions 112 and 158, and is the largest and most common genetic risk factor for late-onset Alzheimer's disease.⁶⁴ ApoE variants with Cys residues in positions 112 or 158, including *APOE2* (Cys112/Cys158), *APOE3* (Cys112/Arg158), and *APOE3'* (Arg112/Cys158) have been shown⁶⁵ or are presumed⁸¹ to confer substantially lower Alzheimer's disease risk than *APOE4*. Encouraged by the ability of NBE1 to convert *APOE4* to *APOE3'* *in vitro* (Figures 16A to 16B), this conversion was attempted in immortalized mouse astrocytes in which the endogenous murine *APOE* gene has been replaced by human *APOE4* (Taconic). DNA encoding NBE3 and an appropriate sgRNA was delivered into these astrocytes by nucleofection (nucleofection efficiency of 25%), extracted genomic DNA from all treated cells two days later, and measured editing efficiency by HTS. Conversion of Arg158 to Cys158 was observed in 58-75% of total DNA sequencing reads (44% of nucleofected astrocytes) (Figures 14A to 14C and Figures 30A). Also observed was 36-50% editing of total DNA at the third position of codon 158 and 38-55% editing of total DNA at the first position of Leu159, as expected since all three of these Cs are within the active nucleobase editing window. However, neither of the other two C→T conversions results in a

change in the amino acid sequence of the ApoE3' protein since both TGC and TGT encode Cys, and both CTG and TTG encode Leu. From > 1,500,000 sequencing reads derived from 1×10^6 cells evidence of 1.7% indels at the targeted locus following NBE3 treatment was observed (Figure 35). In contrast, identical treatment of astrocytes with wt Cas9 and donor ssDNA resulted in 0.1-0.3% APOE4 correction and 26-40% indels at the targeted locus, efficiencies consistent with previous reports of single-base correction using Cas9 and HDR^{45,75} (Figure 30A and Figure 40A). Astrocytes treated identically but with an sgRNA targeting the VEGFA locus displayed no evidence of APOE4 base editing (Figure 34 and Figure 40A). These results demonstrate how nucleobase editors can effect precise, single-amino acid changes in the coding sequence of a protein as the major product of editing, even when their processivity results in more than one nucleotide change in genomic DNA. The off-target activities of Cas9, dCas9, and Cas9 nickase have been extensively studied.^{54, 60-62} In general, off-target C to T conversions by BE1, BE2, and BE3 paralleled off-target Cas9 nuclease-mediated genome modification frequencies.

[00315] The dominant-negative p53 mutations Tyr163Cys and Asn239Asp are strongly associated with several types of cancer.⁶⁶⁻⁶⁷ Both of these mutations can be corrected by a C to T conversion on the template strand (Figures 16A to 16B). A human breast cancer cell line homozygous for the p53 Tyr163Cys mutation (HCC1954 cells) was nucleofected with DNA encoding NBE3 and an sgRNA programmed to correct Tyr163Cys. Because the nucleofection efficiency of HCC1954 cells was < 10%, a plasmid expressing IRFP was co-nucleofected into these cells to enable isolation of nucleofected cells by fluorescence-activated cell sorting two days after treatment. HTS of genomic DNA revealed correction of the Tyr163Cys mutation in 7.6% of nucleofected HCC1954 cells (Figure 30B and Figure 40A to 40B). Also nucleofected was a human lymphoma cell line that is heterozygous for p53 Asn239Asp (ST486 cells) with DNA encoding NBE2 and an sgRNA programmed to correct Asn239Asp with 92% nucleofection efficiency). Correction of the Asn239Asp mutation was observed in 11% of treated ST486 cells (12% of nucleofected ST486 cells). Consistent with the findings in HEK cells, no indels were observed from the treatment of ST486 cells with NBE2, and 0.6% indel formation from the treatment of HCC1954 cells with NBE3. No other DNA changes within at least 50 base pairs of both sides of the protospacer were detected at frequencies above that of untreated controls out of > 2,000,000 sequencing reads derived from 2×10^5 cells (Figures 14A to 14C, Figure 30B and Table 1). These results collectively represent the conversion of three disease-associated alleles in genomic DNA into their wild-

type forms with an efficiency and lack of other genome modification events that is, to our knowledge, not currently achievable using other methods.

[00316] To illuminate the potential relevance of nucleobase editors to address human genetic diseases, the NCBI ClinVar database⁶⁸ was searched for known genetic diseases that could in principle be corrected by this approach. ClinVar was filtered by first examining only single nucleotide polymorphisms (SNPs), then removing any nonpathogenic variants. Out of the 24,670 pathogenic SNPs, 3,956 are caused by either a T to C, or an A to G, substitution. This list was further filtered to only include variants with a nearby NGG PAM that would position the SNP within the deamination activity window, resulting in 1,089 clinically relevant pathogenic gene variants that could in principle be corrected by the nucleobase editors described here (Figure 21 and Table 1). To illuminate the potential relevance of base editors to address human genetic diseases, the NCBI ClinVar database⁶⁸ was searched for known genetic diseases that could in principle be corrected by this approach. ClinVar was filtered by first examining only single nucleotide polymorphisms (SNPs), then removing any non-pathogenic variants. Out of the 24,670 pathogenic SNPs, 3,956 are caused by either a T to C, or an A to G, substitution. This list was further filtered to only include variants with a nearby NGG PAM that would position the SNP within the deamination activity window, resulting in 911 clinically relevant pathogenic gene variants that could in principle be corrected by the base editors described here. Of these, 284 contain only one C within the base editing activity window. A detailed list of these pathogenic mutations can be found in Table 1.

[00317] Table 1. List of 911 base-editable gene variants associated with human disease with an NGG PAM (SEQ ID NOs: 747 to 1868 appear from top to bottom below, respectively). The “Y” in the protospacer and PAM sequences indicates the base to be edited, e.g., C. (SEQ ID NOs: 747 to 1868 appear from top to bottom below, respectively)

dbSNP #	Genotype	Protospacer and PAM sequence(s)	Associated genetic disease
755445790	NM_000391.3(TPP1):c.887-10A>G	TTTTTTTTTTTTTTTTTTGAGG	Ceroid lipofuscinosis, neuronal, 2
113994167	NM_000018.3(ACADVL):c.848T>C (p.Val283Ala)	TTGYGGTGGAGAGGGGCTTCGG, TTGYGGTGGAGAGGGGCTTCGGG	Very long chain acyl-CoA dehydrogenase deficiency
119470018	NM_024996.5(GFM1):c.521A>G (p.Asn174Ser)	TTGYTAATAAAAGTTAGAAACGG	Combined oxidative phosphorylation deficiency 1
115650537	NM_000426.3(LAMA2):c.8282T>C (p.Ile2761Thr)	TTGAYAGGGAGCAAGCAGTTCGG, TGAYAGGGAGCAAGCAGTTCGGG	Merosin deficient congenital muscular dystrophy
587777752	NM_014946.3(SPAST):c.1688-	TTCYGTAAAACATAAAAGTCAGG	Spastic paraplegia 4, autosomal dominant

794726821	NM_001165963.1(SCN1A):c.4055T>C (p.Leu1352Pro)	TTCYGGTTTGTCTTATATTCTGG	Severe myoclonic epilepsy in infancy
397514745	NM_001130089.1(KARS):c.517T>C (p.Tyr173His)	CTTCYATGATCTTCGAGGAGAGG, TTCYATGATCTTCGAGGAGAGG G	Deafness, autosomal recessive 89
376960358	NM_001202.3(BMP4):c.362A>G (p.His121Arg)	TTCGTGGYGGAAAGCTCCTCACGG	Microphthalmia syndromic 6
606231280	NM_001287223.1(SCN11A):c.1142T>C (p.Ile381Thr)	CTTCAYTGTGGTCATTTTCCTGG, TTCAYTGTGGTCATTTTCCTGG G	Episodic pain syndrome, familial, 3
387906735	m.608A>G	TTCAGYGTATTGCTTTGAGGAGG	Cardiomyopathy with or without skeletal myopathy
199474663	m.3260A>G	TAAAGTTYATGCGATTACCGGG	
104894962	NM_003413.3(ZIC3):c.1213A>G (p.Lys405Glu)	TGTGTTYGCGCAGGGAGCTCGGG, ATGTGTTYGCGCAGGGAGCTCG G	Heterotaxy, visceral, X-linked
796053181	NM_021007.2(SCN2A):c.1271T>C (p.Val424Ala)	TGTGGYGGCCATGGCCTATGAGG	not provided
267606788	NM_000129.3(F13A1):c.728T>C (p.Met243Thr)	TGTGAYGGACAGAGCACAAATGG	Factor xiii, a subunit, deficiency of
397514503	NM_003863.3(DPM2):c.68A>G (p.Tyr23Cys)	TGTAGYAGGTGAAGATGATCAGG	Congenital disorder of glycosylation type Iu
104893973	NM_000416.2(IFNGR1):c.260T>C (p.Ile87Thr)	TGTAATAYTTCTGATCATGTTGG	Disseminated atypical mycobacterial infection, Mycobacterium tuberculosis, susceptibility to
121908466	NM_005682.6(ADGRG1):c.263A>G (p.Tyr88Cys)	TGGYAGAGGCCCTGGGGTCAGG	Polymicrogyria, bilateral frontoparietal
147952488	NM_002437.4(MPV17):c.186+2T> C	TGGYAAGTTCTCCCTCAACAGG	Navajo neurohepatopathy
121909537	NM_001145.4(ANG):c.121A>G (p.Lys41Glu)	TGGTTYGGCATCATAGTGCTGGG, GTGGTTYGGCATCATAGTGCTG G	Amyotrophic lateral sclerosis type 9
121918489	NM_000141.4(FGFR2):c.1018T>C (p.Tyr340His)	TGGGGAAYATACGTGCTTGCGGG, GGGGAAYATACGTGCTTGCGGGG	Crouzon syndrome
121434463	m.12320A>G	GAGTYGCACCAAAATTTTGGGG, GGAGTYGCACCAAAATTTTGGG, TGGAGTYGCACCAAAATTTTGG G	Mitochondrial myopathy
121908046	NM_000403.3(GALE):c.101A>G (p.Asn34Ser)	TGGAAGYTATCGATGACCACAGG	UDPglucose-4-epimerase deficiency
431905512	NM_003764.3(STX11):c.173T>C (p.Leu58Pro)	TGCYGGTGGCCGACGTGAAGCGG	Hemophagocytic lymphohistiocytosis, familial, 4
121917905	NM_000124.3(ERCC6):c.2960T>C (p.Leu987Pro)	TGCYAAAAGACCCAAAACAAAGG	Cerebro-oculo-facio-skeletal syndrome
121918500	NM_000141.4(FGFR2):c.874A>G (p.Lys292Glu)	TGCTYATCCACTGGATGTGGGG, GTGCTYATCCACTGGATGTGGG, CGTGCTYATCCACTGGATGTG G	Crouzon syndrome
60431989	NM_000053.3(ATP7B):c.3443T>C (p.Ile1148Thr)	TGCTGAYTGGAACCGTGAGTGG	Wilson disease
78950939	NM_000250.1(MPO):c.518A>G (p.Tyr173Cys)	GTGCGGYATTTGCTCTGCTCCGG, TGCGGYATTTGCTCTGCTCCGG G	Myeloperoxidase deficiency
115677373	NM_201631.3(TGM5):c.763T>C (p.Trp255Arg)	TGCGGAGYGGACGGGCAGCGTGG	Peeling skin syndrome, acral type
5030804	NM_000551.3(VHL):c.233A>G (p.Asn78Ser)	GCGAYTGAGAAGATGACCTGGG, TGCGAYTGAGAAGATGACCTG G	Von Hippel-Lindau syndrome
397508328	NM_000492.3(CFTR):c.1A>G (p.Met1Val)	GCAAYGGTCTCTCGGGCGCTGGG, TGCAAYGGTCTCTCGGGCGCTGGG , CTGCAAYGGTCTCTCGGGCGCTGG	Cystic fibrosis

137853299	NM_000362.4(TIMP3):c.572A>G (p.Tyr191Cys)	TGCAGYAGCCGCCCTTCTGCCGG	Sorsby fundus dystrophy
121908549	NM_000334.4(SCN4A):c.3478A>G (p.Ile1160Val)	TGAYGGAGGGGATGGCGCCTAGG	
121909337	NM_001451.2(FOXF1):c.1138T>C (p.Ter380Arg)	TGATGYGAGGCTGCCGCCGACAGG	Alveolar capillary dysplasia with misalignment of pulmonary veins
281875320	NM_005359.5(SMAD4):c.1500A>G (p.Ile500Met)	TGAGYATGCATAAGCGACGAAGG	Myhre syndrome
730880132	NM_170707.3(LMNA):c.710T>C (p.Phe237Ser)	TGAGTYTGAGAGCCGGCTGGCGG	Primary dilated cardiomyopathy
281875322	NM_005359.5(SMAD4):c.1498A>G (p.Ile500Val)	TGAGTAYGCATAAGCGACGAAGG	Hereditary cancer-predisposing syndrome, Myhre syndrome
72556283	NM_000531.5(OTC):c.527A>G (p.Tyr176Cys)	TGAGGYAATCAGCCAGGATCTGG	not provided
74315311	NM_020435.3(GJC2):c.857T>C (p.Met286Thr)	TGAGAYGGCCACCTGGGCTTGG, GAGAYGGCCACCTGGGCTTGGG	Leukodystrophy, hypomyelinating, 2
121912495	NM_170707.3(LMNA):c.1139T>C (p.Leu380Ser)	TCTYGGAGGGCGAGGAGGAGAGG	Congenital muscular dystrophy, LMNA-related
128620184	NM_000061.2(BTK):c.1288A>G (p.Lys430Glu)	TCTYGATGGCCACGTCGTA CTGG	X-linked agammaglobulinemia
118192252	NM_004519.3(KCNQ3):c.1403A>G (p.Asn468Ser)	TCITTAAYTGTTAAGCCAACAGG	Benign familial neonatal seizures 2, not specified
121909142	NM_001300.5(KLF6):c.190T>C (p.Trp64Arg)	TCTGYGGACCAAAATCATTCTGG	
104895503	NM_001127255.1(NLRP7):c.2738A>G (p.Asn913Ser)	TCTGGYTGATACTCAAGTCCAGG	Hydatidiform mole
587783035	NM_000038.5(APC):c.1744- 2A>G	TCCYAGTAAGAAACAGAATATGG	Familial adenomatous polyposis 1
72556289	NM_000531.5(OTC):c.541- 2A>G	TCCYAAAAGGCACGGGATGAAGG	not provided
28937313	NM_005502.3(ABCA1):c.2804A>G (p.Asn935Ser)	TCCAYTGTGGCCAGGAAGGAGG, CGTCCAYTGTGGCCAGGAAGG	Tangier disease
143246552	NM_001003811.1(TEX11):c.511A>G (p.Met171Val)	TCCAYGGTCAAGTCAGCCTCAGG, CCAYGGTCAAGTCAGCCTCAGG	Spermatogenic failure, X-linked, 2
587776451	NM_002049.3(GATA1):c.2T>C (p.Met1Thr)	CTCCAYGGAGTTCCTGGCCCTGG, TCCAYGGAGTTCCTGGCCCTGGG, CCAYGGAGTTCCTGGCCCTGGG	GATA-1-related thrombocytopenia with dyserythropoiesis
121908403	NM_021102.3(SPINT2):c.488A>G (p.Tyr163Cys)	TCCAYAGATGAAGTTATTGCGAGG	Diarrhea 3, secretory sodium, congenital, syndromic
281874738	NM_000495.4(COL4A5):c.438+2T >C	CTCCAGYAAGTTATAAAAATTGG, TCCAGYAAGTTATAAAAATTGG G	Alport syndrome, X-linked recessive
730880279	NM_030653.3(DDX11):c.2271+2T >C	TCCAGGYGCGGGCGTCATGCTGG, CCAGGYGCGGGCGTCATGCTGGG	Warsaw breakage syndrome
28940272	NM_017890.4(VPS13B):c.8978A>G (p.Asn2993Ser)	TCAYTGATAAGCAGGGCCAGGG, TTCAYTGATAAGCAGGGCCAGG	Cohen syndrome, not specified
137852375	NM_000132.3(F8):c.5372T>C (p.Met1791Thr)	TCAYGGTGTAGTTAAGGACAGTGG	Hereditary factor VIII deficiency disease
11567847	NM_021961.5(TEAD1):c.1261T>C (p.Tyr?His)	TCATATTYACAGGCTTGTA AAGG	
786203989	NM_016069.9(PAM16):c.226A>G (p.Asn76Asp)	CATAGTYCTGCAGAGGAGAGGGG, TCATAGTYCTGCAGAGGAGAGGGG	Chondrodysplasia, megarbane-dagher-melki type
587776437	NC_012920.1:m.9478T>C	TCAGAAGYTTTTTCTTCGCAGG	Leigh disease
121912474	NM_000424.3(KRT5):c.20T>C (p.Val7Ala)	TCAAGTYGTCCTTCCGGAGCGG, CAAGTYGTCCTTCCGGAGCGGG, AAGTYGTCCTTCCGGAGCGGGG, AGTYGTCCTTCCGGAGCGGGG	Epidermolysis bullosa simplex, Koebner type
104886461	NM_020533.2(MCOLN1):c.406- 2A>G	TACYGTGGGCAGAGAAGGGGAGG, AGGTACYGTGGGCAGAGAAGGGG, CAGGTACYGTGGGCAGAGAAGGG	Ganglioside sialidase deficiency

104894275	NM_000317.2(PTS):c.155A>G (p.Asn52Ser)	TAAATGTGCCCATGGCCATTTGG	6-pyruvoyl-tetrahydropterin synthase deficiency
587777562	NM_015599.2(PGM3):c.737A>G (p.Asn246Ser)	TAAATGAYTGAGTTTGCCTTGG	Immunodeficiency 23
121964906	NM_000027.3(AGA):c.916T>C (p.Cys306Arg)	GTTATAYTGCCAATGTGACTGG	Aspartylglycosaminuria
28941769	NM_000356.3(TCOF1):c.149A>G (p.Tyr50Cys)	GTGTGTAYAGATGTCCAGAAGGG	Treacher collins syndrome 1
121434464	m.12297T>C	GTCYTAGGCCCAAAAATTTTGG	Cardiomyopathy, mitochondrial
121908407	NM_054027.4(ANKH):c.143T>C (p.Met48Thr)	GTCGAGAYGCTGGCCAGCTACGG, TCGAGAYGCTGGCCAGCTACGGG	Chondrocalcinosis 2
59151893	NM_000422.2(KRT17):c.275A>G (p.Asn92Ser)	GTCAYTGAGGTTCTGCATGGTGG, GCGGTCAYTGAGGTTCTGCATGG	Pachyonychia congenita type 2
121909499	NM_002427.3(MMP13):c.272T>C (p.Met91Thr)	GTCAYGAAAAAGCCAAGATGCGG, TCAYGAAAAAGCCAAGATGCGG G	
61748478	NM_000552.3(VWF):c.2384A>G (p.Tyr795Cys)	GTCAYAGTTCTGGCACGTTTTGG	von Willebrand disease type 2N

387906889	NM_006796.2(AFG3L2):c.1847A>G (p.Tyr616Cys)	GTAYAGAGGTATTGTTCTTTTGG	Spastic ataxia 5, autosomal recessive
118203907	NM_000130.4(F5):c.5189A>G (p.Tyr1730Cys)	GTAGYAGGCCCAAGCCCGACAGG	Factor V deficiency
118203945	NM_013319.2(UBIAD1):c.305A>G (p.Asn102Ser)	GTAAGTYGTACCAAATTACCGG	Schnyder crystalline corneal dystrophy
267607080	NM_005633.3(SOS1):c.1294T>C (p.Trp432Arg)	GGTYGGGAGGAAAAAGACATTGG	Noonan syndrome 4, Rasopathy
137852953	NM_012464.4(TLL1):c.1885A>G (p.Ile629Val)	GGTTAYGGTGCCGTTAAGTTTGG	Atrial septal defect 6
118203949	NM_013319.2(UBIAD1):c.695A>G (p.Asn232Ser)	GGTGTGTYTGAATGGAGAATGG	Schnyder crystalline corneal dystrophy
137852952	NM_012464.4(TLL1):c.713T>C (p.Val238Ala)	GGGATTGYTGTTCATGAATTGGG	Atrial septal defect 6
41460449	m.3394T>C	GGCYATATACAACACGCAAAGG	Leber optic atrophy
80357281	NM_007294.3(BRCA1):c.5291T>C (p.Leu1764Pro)	GGCYAGAAATCTGTTGCTATGG, GGCYAGAAATCTGTTGCTATGGG	Familial cancer of breast, Breast-ovarian cancer, familial 1
5030764	NM_000174.4(GP9):c.182A>G (p.Asn61Ser)	GGCTGYTGTGGCCAGCAGAAGG	Bernard-Soulier syndrome type C
72556282	NM_000531.5(OTC):c.526T>C (p.Tyr176His)	GGCTGATYACCTCACGCTCCAGG, GATYACCTCACGCTCCAGGTTGG	not provided
121913594	NM_000530.6(MPZ):c.242A>G (p.His81Arg)	GGCATAGYGAAGATCTATGAGG	Charcot-Marie-Tooth disease type 1B
587777736	NM_017617.3(NOTCH1):c.1285T>C (p.Cys429Arg)	GGCAAGYGCATCAACACGCTGGG, GGGCAAGYGCATCAACACGCTGG	Adams-Oliver syndrome 1, Adams- Oliver syndrome 5
63750912	NM_016835.4(MAPT):c.1839T>C (p.Asn613=)	GGATAAYATCAAACACGTCCTCCGG, GATAAYATCAAACACGTCCTCCGG G	Frontotemporal dementia
121918075	NM_000371.3(TTR):c.401A>G (p.Tyr134Cys)	GGAGYAGGGGCTCAGCAGGCGG, ATAGGAGYAGGGGCTCAGCAGGG	Amyloidogenic transthyretin amyloidosis
730882063	NM_004523.3(KIF11):c.2547+2T> C	GGAGGYAATAACTTTGTAAGTGG	Microcephaly with or without chorioretinopathy, lymphedema, or mental retardation
397516156	NM_000257.3(MYH7):c.2546T>C (p.Met849Thr)	GGAGAYGGCCTCCATGAAGGAGG	Primary familial hypertrophic cardiomyopathy,
118204430	NM_000035.3(ALDOB):c.442T>C (p.Trp148Arg)	GGAAGYGGCGTGCTGTGCTGAGG	Hereditary fructosuria
200198778	NM_013382.5(POMT2):c.1997A>G (p.Tyr666Cys)	GGAAGYAGTGGTGGAAGTAGAGG	Congenital muscular dystrophy, Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies, type A2, Muscular dystrophy, Congenital muscular dystrophy- dystroglycanopathy with mental retardation, type B2
754896795	NM_004006.2(DMD):c.6982A>T (p.Lys2328Ter)	GCTTTTYTCAAGCTGCCCAAGG	Duchenne muscular dystrophy, Becker muscular dystrophy, Dilated cardiomyopathy 3B
148924904	NM_000546.5(TP53):c.488A>G (p.Tyr163Cys)	GCTTGYAGATGGCCATGGCGCGG	Hereditary cancer-predisposing syndrome

786204770	NM_016035.4(COQ4):c.155T>C (p.Leu52Ser)	GCTGTGGCCGCCGGCTCCGCGG	COENZYME Q10 DEFICIENCY, PRIMARY, 7
121909520	NM_001100.3(ACTA1):c.350A>G (p.Asn117Ser)	CGGYTGGCCTTGGGATTGAGGG, GCGGYTGGCCTTGGGATTGAGGG, CGCGGYTGGCCTTGGGATTGAGG	Nemaline myopathy 3
587776879	NM_004656.3(BAP1):c.438- 2A>G	GCCYGGGGAAAAACAGATCAGG	Tumor predisposition syndrome
727504434	NM_000501.3(ELN):c.890- 2A>G	GCCYGAACACAGCCACAGAGG	Supravalvar aortic stenosis
119455953	NM_000391.3(TPP1):c.1093T>C (p.Cys365Arg)	GCCGGYGTGTGCTGTCTCTGG	Ceroid lipofuscinosis, neuronal, 2
121964983	NM_000481.3(AMT):c.125A>G (p.His42Arg)	GCCAGGYGGAAGTCATAGAGCGG	Non-ketotic hyperglycinemia
121908300	NM_001005741.2(GBA):c.751T>C (p.Tyr251His)	GCCAGAYACTTTGTGAAGTAAGG, CCAGAYACTTTGTGAAGTAAGG	Gaucher disease, type 1
786205083	NM_003494.3(DYSF):c.3443- 33A>G	GCCAGAGYAGTGGCTGGAGTGG	Limb-girdle muscular dystrophy, type 2B
121908133	NM_175073.2(APTX):c.602A>G (p.His201Arg)	GCCAAAYGGTAACGGCCCTTTGGG, AGCCAAAYGGTAACGGCCCTTTGG	Adult onset ataxia with oculomotor apraxia
587777195	NM_005017.3(PCYT1A):c.571T>C (p.Phe191Leu)	GCATGYTTGCTCCAACACAGAGG	Spondylometaphyseal dysplasia with cone-rod dystrophy
431905520	NM_014714.3(IFIT140):c.4078T>C (p.Cys1360Arg)	CAAGCAGYGTGAGCTGCTCTGG, GCAGYGTGAGCTGCTCTGGAGG	Renal dysplasia, retinal pigmentary dystrophy, cerebellar ataxia and skeletal dysplasia
121912889	NM_001844.4(COL2A1):c.4172A>G (p.Tyr1391Cys)	GCAGTGGYAGGTGATGTCTGGG	Spondyloperipheral dysplasia, Platyspondylic lethal skeletal dysplasia Torrance type
137854492	NM_001363.4(DKC1):c.1069A>G (p.Thr357Ala)	GCAGGYAGAGATGACCGCTGTGG	Dyskeratosis congenita X-linked
121434362	NM_152783.4(D2HGDH):c.1315A>G (p.Asn439Asp)	GCAGGYACCATCTCCTGGAGGG, TGCAGGYACCATCTCCTGGAGG	D-2-hydroxyglutaric aciduria 1
80338732	NM_002764.3(PFPS1):c.344T>C (p.Met115Thr)	GCAAATAYGCTATCTGTAGCAGG	Charcot-Marie-Tooth disease, X- linked recessive, type 5
387906675	NM_000313.3(PROSI):c.701A>G (p.Tyr234Cys)	GATTAYATCTGTAGCCTTCGGGG, AGATTAYATCTGTAGCCTTCGGG, GAGATTAYATCTGTAGCCTTCGG	Thrombophilia due to protein S deficiency, autosomal recessive

28935478	NM_000061.2(BTK):c.1082A>G (p.Tyr361Cys)	GATGGYAGTTAATGAGCTCAGGG, TGATGGYAGTTAATGAGCTCAGG	
201777056	NM_005050.3(ABCD4):c.956A>G (p.Tyr319Cys)	GATGAGGYAGATGCACACAAAAGG	METHYLMALONIC ACIDURIA AND HOMOCYSTINURIA, cblJ
121918528	NM_000098.2(CPT2):c.359A>G (p.Tyr120Cys)	GATAGGYACATATCAAACCAGGG, AGATAGGYACATATCAAACCAG G	Carnitine palmitoyltransferase II deficiency, infantile
267607014	NM_002942.4(ROBO2):c.2834T>C (p.Ile945Thr)	GAGAYTGGAAATTTGGCCGTGG	Vesicoureteral reflux 2
281865192	NM_025114.3(CEP290):c.2991+1655 A>G	GATAYTCACAATTACAACCTGGGG, AGATAYTCACAATTACAACCTGGG, GAGATAYTCACAATTACAACCTG	Leber congenital amaurosis 10
386833492	NM_000112.3(SLC26A2):c.- 26+2T>C	GAGAGGYGAGAAGAGGGGAACGGG	Diastrophic dysplasia
587779773	NM_001101.3(ACTB):c.356T>C (p.Met119Thr)	GAGAAGAYGACCCAGGTGAGTGG	Baraitser-Winter syndrome 1
121913512	NM_000222.2(KIT):c.1924A>G (p.Lys642Glu)	GACTTYGAGTTCAGACATGAGGG, GGACTTYGAGTTCAGACATGAGG	
28939072	NM_006329.3(FBLN5):c.506T>C (p.Ile169Thr)	GACAYTGATGAATGTCGCTATGG	Age-related macular degeneration 3
104894248	NM_000525.3(KCNJ11):c.776A>G (p.His259Arg)	GACAYGGTAGATGATCAGCGGGG, TGACAYGGTAGATGATCAGCGGG, ATGACAYGGTAGATGATCAGCGG	Islet cell hyperplasia
387907132	NM_016464.4(TMEM138):c.287A>G (p.His96Arg)	GACAYGAAGGGAGATGCTGAGGG, AGACAYGAAGGGAGATGCTGAGG	Joubert syndrome 16
121918170	NM_000275.2(OCA2):c.1465A>G (p.Asn489Asp)	GACATYTGAGGGTCCCGCATGG	Tyrosinase-positive oculocutaneous albinism
122467173	NM_014009.3(FOXP3):c.970T>C (p.Phe324Leu)	GACAGAGYTCCTCCACAACATGG	Insulin-dependent diabetes mellitus secretory diarrhea syndrome
137852268	NM_000133.3(F9):c.1328T>C (p.Ile443Thr)	GAAYATATACCAAGGTATCCCGG	Hereditary factor IX deficiency disease
149054177	NM_001999.3(FBN2):c.3740T>C (p.Met1247Thr)	GAATGTAYGATAATGAACGGAGG	not specified, Macular degeneration, early- onset

137854488	NM_212482.1(FN1):c.2918A>G (p.Tyr973Cys)	GAAGTAAAYAGGTGACCCAGGGG	Glomerulopathy with fibronectin deposits 2
786204027	NM_005957.4(MTHFR):c.1530+2T>C	GAAGGYGTGGTAGGGAGGCACGG, AAGGYGTGGTAGGGAGGCACGG, AGGYGTGGTAGGGAGGCACGGGG	Homocysteinemia due to MTHFR deficiency
104894223	NM_012193.3(FZD4):c.766A>G (p.Ile256Val)	GAAATAYGATGGGGCCTCAGGG, AGAAATAYGATGGGGCGCTCAGG	Retinopathy of prematurity
137854474	NM_000138.4(FBN1):c.3793T>C (p.Cys1265Arg)	CTTGYGTTATGATGGATTTCATGG	Marfan syndrome
587784418	NM_006306.3(SMC1A):c.3254A>G (p.Tyr1085Cys)	CTTAYAGATCTCATCAATGTTGG	Congenital muscular hypertrophy-cerebral syndrome
81002805	NM_000059.3(BRCA2):c.316+2T>C	CTTAGGYAAGTAATGCAATATGG	Familial cancer of breast, Breast-ovarian cancer, familial 2, Hereditary cancer-predisposing syndrome
121909653	NM_182925.4(FLT4):c.3104A>G (p.His1035Arg)	CTGYGGATGCACTGGGGTGCGGG, TCTGYGGATGCACTGGGGTGCGG	
786205107	NM_031226.2(CYP19A1):c.743+2T>C	CTGTGYAAGTAATACAACCTTTGG	Aromatase deficiency
587777037	NM_001283009.1(RTEL1):c.3730T>C (p.Cys1244Arg)	CTGTGTGYGCCAGGGCTGTGGGG	Dyskeratosis congenita, autosomal recessive, 5
794728380	NM_000238.3(KCNH2):c.1945+6T>C	CTGTGAGYGTGCCAGGGGCGGG, TGAGYGTGCCAGGGGCGGGCGG	Cardiac arrhythmia
267607987	NM_000251.2(MSH2):c.2005+2T>C	CTGGYAAAAAACCTGGTTTTTGG, TGGYAAAAAACCTGGTTTTTGG G	Hereditary Nonpolyposis Colorectal Neoplasms
397509397	NM_006876.2(B4GAT1):c.1168A>G (p.Asn390Asp)	TGATYTTACGCCCTCCTTTTGGGG, CTGATYTTACGCCCTCCTTTTGGG, GCTGATYTTACGCCCTCCTTTTGG	Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies, type A13
121918381	NM_000040.1(APOC3):c.280A>G (p.Thr94Ala)	CTGAAGYTGCTGACCTCAGGG, GCTGAAGYTGCTGACCTCAGG	
104894919	NM_001015877.1(PHF6):c.769A>G (p.Arg257Gly)	CTCYTGATGTTGTTGTGAGCTGG	Borjeson-Forssman-Lehmann syndrome
267606869	NM_005144.4(HR):c.-218A>G	CTCYAGGGCCGAGGTTGGAGG, GCTCYAGGGCCGAGGTTGGAGG, GGCGCTCYAGGGCCGAGGTTGG	Marie Unna hereditary hypotrichosis 1
139732572	NM_000146.3(FTL):c.1A>G (p.Met1Val)	CTCAYGGTTGGTTGGCAAGAAGG	L-ferritin deficiency
397515418	NM_018486.2(HDAC8):c.1001A>G (p.His334Arg)	CTCAYGATCTGGGATCTCAGAGG	Cornelia de Lange syndrome 5
372395294	NM_198056.2(SCN5A):c.1247A>G (p.Tyr416Cys)	CTCAYAGGCCATTGCGACCACGG	not provided
104895304	NM_000431.3(MVK):c.803T>C (p.Ile268Thr)	CTCAAYAGATGCCATCTCCCTGG	Hyperimmunoglobulin D with periodic fever, Mevalonic aciduria
587777188	NM_001165899.1(PDE4D):c.1850T>C (p.Ile617Thr)	CTATAYTGTTATCCCCTCTGGG, ACTATAYTGTTATCCCCTCTGG	Acrodysostosis 2, with or without hormone resistance
398123026	NM_003867.3(FGF17):c.560A>G (p.Asn187Ser)	CGTGGYTGGGGAAGGGCAGCTGG	Hypogonadotropic hypogonadism 20 with or without anosmia
121964924	NM_001385.2(DPYS):c.1078T>C (p.Trp360Arg)	CGTAATAYGGGAAAAAGGCGTGG, AATAYGGGAAAAAGGCGTGGTGG, ATAYGGGAAAAAGGCGTGGTGGG	Dihydropyrimidinase deficiency
587777301	NM_199189.2(MATR3):c.1864A>G (p.Thr622Ala)	CGGYTGAACCTCAGTCTTCTGG	Myopathy, distal, 2
200238879	NM_000527.4(LDLR):c.694+2T>C	ACTGCGGYATGGGCGGGGCCAGG, CTGCGGYATGGGCGGGGCCAGGG, CGGYATGGGCGGGGCCAGGGTGG	Familial hypercholesterolemia
142951029	NM_145046.4(CALR3):c.245A>G (p.Lys82Arg)	CGGTYTGAAGCGTGCAGAGATGG	Arrhythmogenic right ventricular cardiomyopathy, Familial hypertrophic cardiomyopathy 19, Hypertrophic cardiomyopathy
786200953	NM_006785.3(MALT1):c.1019-2A>C	CGCYTTGAAAAAAAAGAAAGGG, TCGCYTTGAAAAAAAAGAAAG	Combined immunodeficiency
120074192	NM_000218.2(KCNQ1):c.418A>G (p.Ser140Gly)	CGCYGAAGATGAGGCAGACCAGG	Atrial fibrillation, familial, 3, Atrial fibrillation
267606887	NM_005957.4(MTHFR):c.971A>G (p.Asn324Ser)	CGCGGYTGAGGGTGTAGAAGTGG	Homocystinuria due to MTHFR deficiency
118192117	NM_000540.2(RYR1):c.1205T>C (p.Met402Thr)	CGCAYGATCCACAGCACC AATGG	Congenital myopathy with fiber type disproportion, Central core disease
199473625	NM_198056.2(SCN5A):c.4978A>G (p.Ile1660Val)	CGAYTTGAAGAGGGCAGGCAGG, AGCCCGAYGTGAAGAGGGCAGG	Brugada syndrome
794726865	NM_000921.4(PDE3A):c.1333A>G (p.Thr445Ala)	CGAGGYGGTGGTGGTCCAAGTGG	Brachydactyly with hypertension
606231254	NM_005740.2(DNAL4):c.153+2T>C	CGAGGYATTGCCAGCAGTGCAGG	Mirror movements 3
786204826	NM_004771.3(MMP20):c.611A>G (p.His204Arg)	CGAAAYGTGATCTCCTCCACAGG	Amelogenesis imperfecta, hypomaturation type, IA2

796053139	NM_021007.2(SCN2A):c.4308+2T>C	CGAAATGYAAGTCTAGTTAGAGG, GAAATGYAAGTCTAGTTAGAGG	not provided
137854494	NM_005502.3(ABCA1):c.4429T>C (p.Cys1477Arg)	CCTGTGYGTCCCCAGGGGCAGG, CTGTGYGTCCCCAGGGGCAGGG, TGTGYGTCCCCAGGGGCAGGGG, GTGYGTCCCCAGGGGCAGGGG	Tangier disease
786205144	NM_001103.3(ACTN2):c.683T>C (p.Met228Thr)	CCTAAAAAGTTGGATGCTGAAGG	Dilated cardiomyopathy 1AA
199919568	NM_007254.3(PNKP):c.1029+2T>C	CCGGYGAGGCCCTGGGGCGGGG, TCCGGYGAGGCCCTGGGGCGGGG, ATCCGGYGAGGCCCTGGGGCGGG, GATCCGGYGAGGCCCTGGGGCGG	not provided
28939079	NM_018965.3(TREM2):c.401A>G (p.Asp134Gly)	TGAYCCAGGGGGTCTATGGGAGG, CGGTGAYCCAGGGGGTCTATGGG, CCGGTGTAYCCAGGGGGTCTATGG	Polycystic lipomembranous osteodysplasia with sclerosing leukoencephalopathy
193302855	NM_032520.4(GNPTG):c.610-2A>G	CCCYGAAGGTGGAGGATGCAGG, GCCCYGAAGGTGGAGGATGCAGG	Mucopolidiosis III Gamma
111033708	NM_000155.3(GALT):c.499T>C (p.Trp167Arg)	CCCTYGGGTGCAGGTTTGTGAGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
28933378	NM_000174.4(GP9):c.70T>C (p.Cys24Arg)	CCCAYGTACCTGCCGCGCCCTGG	Bernard Soulier syndrome, Bernard-Soulier syndrome type C
364897	NM_000157.3(GBA):c.680A>G (p.Asn227Ser)	CCAYTGGTCTTGAGCCAAGTGGG, TCCAYTGGTCTTGAGCCAAGTGG	Gaucher disease, Subacute neuronopathic Gaucher disease, Gaucher disease, type 1
796052551	NM_000833.4(GRIN2A):c.2449A>G (p.Met817Val)	CCAYGTTGTCAATGTCCAGCTGG	not provided
63751006	NM_002087.3(GRN):c.2T>C (p.Met1Thr)	CCAYGTGGACCCTGGTGAGCTGG	Frontotemporal dementia, ubiquitin-positive
786203997	NM_001031.4(RPS28):c.1A>G (p.Met1Val)	TGTCCAYGATGGCGCGCGGGCGG, CCAYGATGGCGCGCGGGCGGG	Diamond-Blackfan anemia with microtia and cleft palate
121908595	NM_002755.3(MAP2K1):c.389A>G (p.Tyr130Cys)	CCAYAGAAGCCACGATGTACGG	Cardiofaciocutaneous syndrome 3, Rasopathy
398122910	NM_000431.3(MVK):c.1039+2T>C	CCAGGYATCCCGGGGTAGGTGG, CAGGYATCCCGGGGTAGGTGGG	Porokeratosis, disseminated superficial actinic 1
119474039	NM_020365.4(EIF2B3):c.1037T>C (p.Ile346Thr)	CCAGAYTGTCAGCAAACACCTGG	Leukoencephalopathy with vanishing white matter
58777866	NM_000076.2(CDKN1C):c.*5+2T>C	CCAAGYGAGTACAGCGCACCTGG, CAAGYGAGTACAGCGCACCTGGG, AAGYGAGTACAGCGCACCTGGGG	Beckwith-Wiedemann syndrome
121918530	NM_005587.2(MEF2A):c.788A>G (p.Asn263Ser)	AGAYTACCACCACTGGTGGAGG, CCAAGAYTACCACCACTGGTGG	
483352818	NM_000211.4(ITGB2):c.1877+2T>C	CATGYGAGTGCAGGCGGAGCAGG	Leukocyte adhesion deficiency type 1
460184	NM_000186.3(CFH):c.3590T>C (p.Val1197Ala)	CAGYTGAATTTGTGTGTAACCGG	Atypical hemolytic-uremic syndrome 1
121908423	NM_004795.3(KL):c.578A>G (p.His193Arg)	CAGYGGTACAGGGTGACCACGGG, CCAGYGGTACAGGGTGACCACGG	
281860300	NM_005247.2(FGF3):c.146A>G (p.Tyr49Cys)	CAGYAGAGCTTGGCGGCCGGGG, GCAGYAGAGCTTGGCGGCCGGG, CGCAGYAGAGCTTGGCGGCCGG	Deafness with labyrinthine aplasia microtia and microdontia (LAMM)
28935488	NM_000169.2(GLA):c.806T>C (p.Val269Ala)	CAGTTAGYGATTGGCAACTTTGG	Fabry disease
587776514	NM_173560.3(RFX6):c.380+2T> C	CAGTGGYGAGACTCGCCCGCAGG, AGTGGYGAGACTCGCCCGCAGGG	Mitchell-Riley syndrome
104894117	NM_178138.4(LHX3):c.332A>G (p.Tyr111Cys)	CAGGTGGYACACGAAGTCTCTGGG	Pituitary hormone deficiency, combined 3
34878913	NM_000184.2(HBG2):c.125T>C (p.Phe42Ser)	CAGAGGTYCTTTGACAGCTTTGG	Cyanosis, transient neonatal
120074124	NM_000543.4(SMPD1):c.911T>C (p.Leu304Pro)	AGCACYTGTGAGGAAGTTCTCTGG, GCACYTGTGAGGAAGTTCTCTGGG, CACYTGTGAGGAAGTTCTCTGGGG	Sphingomyelin/cholesterol lipodosis, Niemann- Pick disease, type A, Niemann-Pick disease, type B
281860272	NM_005211.3(CSF1R):c.2320- 2A>G	CACYGAGGGAAAGCACTGCAGGG, GCACYGAGGGAAAGCACTGCAGG	Hereditary diffuse leukoencephalopathy with spheroids
128624216	NM_000033.3(ABCD1):c.443A>G (p.Asn148Ser)	CACTGYTGACGAAGGTAGCAGGG, GCACGYTGACGAAGGTAGCAGG	Adrenoleukodystrophy
398124257	NM_012463.3(ATP6V0A2):c.825+2 T>C	CACTGYGAGTAAGCTGGAAGTGG	Cutis laxa with osteodystrophy
267606679	NM_004183.3(BEST1):c.704T>C (p.Val235Ala)	CACTGGYGTATACACAGGTGAGG	Vitreoretinopathology dominant
397514518	NM_000344.3(SMN1):c.388T>C (p.Tyr130His)	CACTGGAYATGGAAATAGAGAGG	Kugelberg-Welander disease

143946794	NM_001946.3(DUSP6):c.566A>G (p.Asn189Ser)	CACTAYTGGGGTCTCGGTCAAGG	Hypogonadotropic hypogonadism 19 with or without anosmia
397516076	NM_000256.3(MYBPC3):c.821+2T>C	GCACGYGAGTGGCCATCTCAGG, CACGYGAGTGGCCATCTCAGGG	Familial hypertrophic cardiomyopathy 4, not specified
149977726	NM_001257988.1(TYMP):c.665A>G (p.Lys222Arg)	CACGAGTYTCTTACTGAGAATGG, GAGTYTCTTACTGAGAATGGAGG	
121917770	NM_003361.3(UMOD):c.383A>G (p.Asn128Ser)	CACAYTGACACATGTGGCCAGGG, CCACAYTGACACATGTGGCCAGG	Familial juvenile gout
121909008	NM_000492.3(CFTR):c.2738A>G (p.Tyr913Cys)	CACATAAYACGAACTGGTGCTGG	Cystic fibrosis
137852819	NM_003688.3(CASK):c.2740T>C (p.Trp914Arg)	CACAGYGGGTCCCTGTCTCCTGG, ACAGYGGGTCCCTGTCTCCTGGG	FG syndrome 4
74315320	NM_024009.2(GJB3):c.421A>G (p.Ile141Val)	CAAYGATGAGCTTGAAGATGAGG	Deafness, autosomal recessive
80356747	NM_001701.3(BAAT):c.967A>G (p.Ile323Val)	CAAYGAAGAGGAATGCCCTGG	Atypical hemolytic-uremic syndrome 1
180177324	NM_012203.1(GRHPR):c.934A>G (p.Asn312Asp)	CAAGTYGTTAGTGCCAACAAGG	Primary hyperoxaluria, type II
281860274	NM_005211.3(CSF1R):c.2381T>C (p.Ile794Thr)	CAAGAYTGGGGACTTCGGGCTGG	Hereditary diffuse leukoencephalopathy with spheroids
398122908	NM_005334.2(HCFC1):c.-970T>C	CAAGAYGGCGGCTCCCAGGGAGG	Mental retardation 3, X-linked
548076633	NM_002693.2(POLG):c.3470A>G (p.Asn157Ser)	CAAGAGGYTGGTGATCTGCAAGG	not provided
120074146	NM_000019.3(ACAT1):c.935T>C (p.Ile312Thr)	CAAGAAAYAGTAGGTAAGGCCAGG	Deficiency of acetyl-CoA acetyltransferase
397514489	NM_005340.6(HINT1):c.250T>C (p.Cys84Arg)	CAAGAAAYGTGCTGCTGATCTGG, AAGAAAYGTGCTGCTGATCTGGG	Gamstorp-Wohlfart syndrome
587783539	NM_178151.2(DCX):c.2T>C (p.Met1Thr)	CAAAATAYGGAACCTTGATTTTGG	Heterotopia
104894765	NM_005448.2(BMP15):c.704A>G (p.Tyr235Cys)	ATTGAAAYAGAGTAACAAGAAGG	Ovarian dysgenesis 2
137852429	NM_000132.3(F8):c.1892A>G (p.Asn631Ser)	ATGYTGAGGCTTGAACTCTGG	Hereditary factor VIII deficiency disease
72558441	NM_000531.5(OTC):c.779T>C (p.Leu260Ser)	ATGTATYAATTACAGACACTTGG	not provided
398123765	NM_003494.3(DYSF):c.1284+2T>C	ATGGYAAGGAGCAAGGGAGCAGG	Limb-girdle muscular dystrophy, type 2B
387906924	NM_020191.2(MRPS22):c.644T>C (p.Leu215Pro)	ATCYTAGGGTAAGGTGACTTAGG	Combined oxidative phosphorylation deficiency 5
397518039	NM_206933.2(USH2A):c.8559-2A>G	ATCYAAAGCAAAAGACAAGCAGG	Retinitis pigmentosa, Usher syndrome, type 2A
5742905	NM_000071.2(CBS):c.833T>C (p.Ile278Thr)	ATCAYTGGGGTGGATCCCGAAGG, TCAYTGGGGTGGATCCCGAAGGG	Homocystinuria due to CBS deficiency, Homocystinuria, pyridoxine-responsive
397507473	NM_004333.4(BRAF):c.1403T>C (p.Phe468Ser)	ATCATYTGGAACAGTCTACAAGG, TCATYTGGAACAGTCTACAAGG	Cardiofaciocutaneous syndrome, Rasopathy
786204056	NM_000264.3(PTCH1):c.3168+2T>C	ATCATTYGGAGTGTATTATAAAGG, TCATTGYGAGTGTATTATAAGGG, CATTGYGAGTGTATTATAAGGG	Gorlin syndrome
72558484	NM_000531.5(OTC):c.1005+2T>C	ATCATGGYAAGCAAGAAACAAGG	not provided
199473074	NM_000335.4(SCN5A):c.688A>G (p.Ile230Val)	ATAYAGTTTTTCAGGGCCCGGAGG, CTGATAYAGTTTTTCAGGGCCCGG	Brugada syndrome
111033273	NM_206933.2(USH2A):c.1606T>C (p.Cys536Arg)	ATATAGAYGCCTCTGCTCCAGG	Usher syndrome, type 2A
72556290	NM_000531.5(OTC):c.542A>G (p.Glu181Gly)	ATAGTGYCCTAAAAGGCACGGG	not provided
121918711	NM_004612.3(TGFBR1):c.1199A>G (p.Asp400Gly)	ATAGATGYCAGCACGTTTGAAGG	Loeys-Dietz syndrome 1
104886288	NM_000495.4(COL4A5):c.4699T>C (p.Cys1567Arg)	AGTAYGTGAAGCTCCAGCTGTGG	Alport syndrome, X-linked recessive
144637717	NM_016725.2(FOLR1):c.493+2T>C	CTTCAGGYGAGGGCTGGGGTGGG, AGGYGAGGGCTGGGGTGGGCAGG	not provided
72558492	NM_000531.5(OTC):c.1034A>G (p.Tyr345Cys)	AGGTGAGYAATCTGTACAGCAGGG	not provided
62638745	NM_000121.3(EPOR):c.1460A>G (p.Asn487Ser)	AGGGYTGGAGTAGGGGCCATCGG	Acute myeloid leukemia, M6 type, Familial erythrocytosis, 1

387907021	NM_031427.3(DNAL1):c.449A>G (p.Asn150Ser)	AGGGAYTGCCTACAAACACCAGG	Kartagener syndrome, Ciliary dyskinesia, primary, 16
397514488	NM_001161581.1(POCIA):c.398T>C (p.Leu133Pro)	AGCYGTGGGACAAGAGCAGCCGG	Short stature, onychodysplasia, facial dysmorphism, and hypotrichosis
154774633	NM_017882.2(CLN6):c.200T>C (p.Leu67Pro)	AGCYGGTATTCCTCTCAGTGG	Adult neuronal ceroid lipofuscinosis
111033700	NM_000155.3(GALT):c.482T>C (p.Leu161Pro)	AGCYGGGTGCCAGTACCCTTGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
128621198	NM_000061.2(BTK):c.1223T>C (p.Leu408Pro)	GAGCYGGGACTGGACAATTTGG, AGCYGGGACTGGACAATTTGGG	X-linked agammaglobulinemia
137852611	NM_000211.4(ITGB2):c.446T>C (p.Leu149Pro)	AGCYAGTGGCGACCTGCTCCGG	Leukocyte adhesion deficiency
121908838	NM_003722.4(TP63):c.697A>G (p.Lys233Glu)	AGCTTYTTTGTAGACAGGCATGG	Split-hand/foot malformation 4
397515869	NM_000169.2(GLA):c.1153A>G (p.Thr385Ala)	AGCTGTGYGATGAAGCAGGCAGG	not specified
118204064	NM_000237.2(LPL):c.548A>G (p.Asp183Gly)	GCTGGAYCGAGGCCCTTAAAAGGG, AGCTGGAYCGAGGCCCTTAAAAGG	Hyperlipoproteinemia, type I
128620186	NM_000061.2(BTK):c.2T>C (p.Met1Thr)	AGCTAYGGCCGAGTGATTCTGG	X-linked agammaglobulinemia
786204132	NM_014946.3(SPAST):c.1165A>G (p.Thr389Ala)	ATTGYCTTCCATTCCCAGGTGG, AGCATTGYCTTCCCATTCCCAGG CAGCAAGBACGTGGGCCTCTGGG, AGCAAGBACGTGGGCCTCTGGG, GCAAGBACGTGGGCCTCTGGGG	Spastic paraplegia 4, autosomal dominant
199473661	NM_000218.2(KCNQ1):c.550T>C (p.Tyr184His)	AGCAAGBACGTGGGCCTCTGGG, AGCAAGBACGTGGGCCTCTGGG, GCAAGBACGTGGGCCTCTGGGG	Congenital long QT syndrome, Cardiac arrhythmia
387907129	NM_024599.5(RHBDF2):c.557T>C (p.Ile186Thr)	AGAYTGTGGATCCGCTGGCCCGG	Howell-Evans syndrome
387906702	NM_006306.3(SMC1A):c.2351T>C (p.Ile784Thr)	AGAYTGGTGTGCGCAACATCCGG	Congenital muscular hypertrophy-cerebral syndrome
193929348	NM_000525.3(KCNJ11):c.544A>G (p.Ile182Val)	AGAYGAGGGTCTCAGCCCTGCCG	Permanent neonatal diabetes mellitus
121908934	NM_004086.2(COCH):c.1535T>C (p.Met512Thr)	AGATAYGGCTTCTAAACCGAAGG	Deafness, autosomal dominant 9
397514377	NM_000060.3(BTD):c.641A>G (p.Asn214Ser)	AGAGGYTGTGTTACGGTAGCGG	Biotinidase deficiency
72552295	NM_000531.5(OTC):c.2T>C (p.Met1Thr)	AGAAGAYGCTGTTAATCTGAGG	not provided
201893545	NM_016247.3(IMPG2):c.370T>C (p.Phe124Leu)	ACTYTTTGGGATCGACTTCCTGG	Macular dystrophy, vitelliform, 5
121434469	m.4290T>C	ACTYTGATAGAGTAAATAATAGG	
121918733	NM_006920.4(SCN1A):c.269T>C (p.Phe90Ser)	ACTTYTATAGTATTGAATAAAGG, CTTYTATAGTATTGAATAAAGG G	Severe myoclonic epilepsy in infancy
121434471	m.4291T>C	ACTYTGATAGAGTAAATAATAGG	Hypertension, hypercholesterolemia, and hypomagnesemia, mitochondrial Sideroblastic anemia with B-cell immunodeficiency, periodic fevers, and developmental delay
606231289	NM_001302946.1(TRNT1):c.497T>C (p.Leu166Ser)	ACTTYATTTGACTACTTTAATGG	
63750067	NM_000517.4(HBA2):c.*92A>G	CTTYATTCAAAGACCAGGAAGGG, ACTTYATTCAAAGACCAGGAAG G	Hemoglobin H disease, nondeletional
121918734	NM_006920.4(SCN1A):c.272T>C (p.Ile91Thr)	ACTTTTAYAGTATTGAATAAAGG, CTTTTAYAGTATTGAATAAAGG G	Severe myoclonic epilepsy in infancy
137854557	NM_000267.3(NF1):c.1466A>G (p.Tyr489Cys)	ACTTAYAGCTTCTGTCTCCAGG	Neurofibromatosis, type 1
397514626	NM_018344.5(SLC29A3):c.607T>C (p.Ser203Pro)	ACTGATAYCAGGTGAGAGCCAGG, CTGATAYCAGGTGAGAGCCAGGG	Histiocytosis-lymphadenopathy plus syndrome
118204440	NM_000512.4(GALNS):c.1460A>G (p.Asn487Ser)	ACGYTGAGCTGGGGCTGCCGCGG, CACGYTGAGCTGGGGCTGCCGCGG	Mucopolysaccharidosis, MPS-IV-A
587776843	NG_012088.1:g.2209A>G	ACCYTATGATCCGCCCGCCTTGG	
137853033	NM_001080463.1(DYNC2H1):c.4610A>G (p.Gln1537Arg)	ACCYGTGAAGGGAACAGAGATGG	Short-rib thoracic dysplasia 3 with or without polydactyly
28933698	NM_000435.2(NOTCH3):c.1363T>C (p.Cys455Arg)	TTCACCYGTATCTGTATGGCAGG, ACCYGTATCTGTATGGCAGGTGG	Cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy
587776766	NM_000463.2(UGT1A1):c.1085-2A>G	ACCYGAGATGCAAAATAGGGAGG, GTGACCYGAGATGCAAAATAGGG, GGTGACCYGAGATGCAAAATAGG	Crigler Najjar syndrome, type 1
587781628	NM_001128425.1(MUTYH):c.1187-2A>G	ACCYGAGAGGGAGGGCAGCCAGG	Hereditary cancer-predisposing syndrome, Carcinoma of colon
61755817	NM_000322.4(PRP2):c.736T>C (p.Trp246Arg)	ACCTGYGGGTGCGTGGCTGCAGG, CCTGYGGGTGCGTGGCTGCAGGG	Retinitis pigmentosa

121909184	NM_001089.2(ABCA3):c.1702A>G (p.Asn568Asp)	ACCGTYGTGGCCAGCAGGACGG	Surfactant metabolism dysfunction, pulmonary, 3
121434466	m.4269A>G	ACAYATTTCCTTAGGTTTGAGGGG, GACAYATTTCCTTAGGTTTGAGGG, AGACAYATTTCCTTAGGTTTGAGG	
794726768	NM_001165963.1(SCN1A):c.1048A>G (p.Met350Val)	ACAYATATCCCTCTGGACATTGG	Severe myoclonic epilepsy in infancy
28934876	NM_001382.3(DPAGT1):c.509A>G (p.Tyr170Cys)	ACAYAGTACAGGATTCCTGCGGG, GACAYAGTACAGGATTCCTGCGG	Congenital disorder of glycosylation type 1J
104894749	NM_000054.4(AVPR2):c.614A>G (p.Tyr205Cys)	ACAYAGGTGCGACGGCCCCAGGG, GACAYAGGTGCGACGGCCCCAGG	Nephrogenic diabetes insipidus, Nephrogenic diabetes insipidus, X-linked
128621205	NM_000061.2(BTK):c.1741T>C (p.Trp581Arg)	ACATTYGGGCTTTTGTAAGTGG	X-linked agammaglobulinemia
28940892	NM_000529.2(MC2R):c.761A>G (p.Tyr254Cys)	ACATGYAGCAGGCGCAGTAGGGG, GACATGYAGCAGGCGCAGTAGGG, AGACATGYAGCAGGCGCAGTAGG	ACTH resistance
794726844	NM_001165963.1(SCN1A):c.1046A>G (p.Tyr349Cys)	ACATAYATCCCTCTGGACATTGG	Severe myoclonic epilepsy in infancy
587783083	NM_003159.2(CDKL5):c.449A>G (p.Lys150Arg)	ACAGTYTTAGGACATCATTGTGG	not provided
397514651	NM_000108.4(DLD):c.140T>C (p.Ile47Thr)	ACAGTTAYAGGTTCTGGTCTGG, GTTAYAGGTTCTGGTCTGGAGG	Maple syrup urine disease, type 3
794727060	NM_001848.2(COL6A1):c.957+2T >C	ACAAGGYGAGCGTGGGCTGCTGG, CAAGGYGAGCGTGGGCTGCTGG	Ullrich congenital muscular dystrophy, Bethlem myopathy
72554346	NM_000531.5(OTC):c.284T>C (p.Leu95Ser)	ACAAGATYGTCTACAGAAACAGG	not provided
483353031	NM_002136.2(HNRNPA1):c.841T>C (p.Phe281Leu)	AATYTTGGAGGCAGAAGCTCTGG	Chronic progressive multiple sclerosis
104894271	NM_000315.2(PTH):c.52T>C (p.Cys18Arg)	AATYGTITTTCTTACAAAATCGG	Hypoparathyroidism familial isolated
267608260	NM_015599.2(PGM3):c.248T>C (p.Leu83Ser)	AATGTYGGCACCATCTGGGAGG	Immunodeficiency 23
267606900	NM_018109.3(MTPAP):c.1432A>G (p.Asn478Asp)	AATGGATYCTGAATGTACAGAGG	Ataxia, spastic, 4, autosomal recessive
796053169	NM_021007.2(SCN2A):c.387- 2A>G	AATAAAGYAGAATATCGTCAAGG	not provided
104894937	NM_000116.4(TAZ):c.352T>C (p.Cys118Arg)	AAGYGTGTGCCTGTGTGCCGAGG	3-Methylglutaconic aciduria type 2
104893911	NM_001018077.1(NR3C1):c.1712T>C (p.Val571Ala)	AAGYGATTGCAGCAGTGAATGG	Pseudohermaphroditism, female, with hypokalemia, due to glucocorticoid resistance
397514472	NM_004813.2(PEX16):c.992A>G (p.Tyr331Cys)	AAGYAGATTTCTGCCAGGTGGG, GAAGYAGATTTCTGCCAGGTGG, GTAGAAGYAGATTTCTGCCAGG	Peroxisome biogenesis disorder 8B
121918407	NM_001083112.2(GPD2):c.1904T>C (p.Phe635Ser)	AAGTYTGATGCAGACCAGAAAGG	Diabetes mellitus type 2
63751110	NM_000251.2(MSH2):c.595T>C (p.Cys199Arg)	AAGGAAYGTGTTTACC CGGAGG	Hereditary Nonpolyposis Colorectal Neoplasms
119450945	NM_000026.2(ADSL):c.674T>C (p.Met225Thr)	AAGAYGGTGACAGAAAAGGCAGG	Adenylosuccinate lyase deficiency
113993988	NM_002863.4(PYGL):c.2461T>C (p.Tyr821His)	AAGAAYATGCCAAAACATCTGG	Glycogen storage disease, type VI
119485091	NM_022041.3(GAN):c.1268T>C (p.Ile423Thr)	AAGAAAAYCTACGCCATGGGTGG, AAAAYCTACGCCATGGGTGGAGG	Giant axonal neuropathy
137852419	NM_000132.3(F8):c.1660A>G (p.Ser554Gly)	AACYAGAGTAATAGCGGGTCAGG	Hereditary factor VIII deficiency disease
121964967	NM_000071.2(CBS):c.1150A>G (p.Lys384Glu)	AACTYGGTCTGCGGGATGGGGG, GAACTYGGTCTGCGGGATGGGG, GGAACYGGTCTGCGGGATGGG, AGGAACTYGGTCTGCGGGATGG	Homocystinuria, pyridoxine-responsive
137852376	NM_000132.3(F8):c.1754T>C (p.Ile585Thr)	AACAGAYAATGTGACAGAAAGAGG	Hereditary factor VIII deficiency disease
121917930	NM_006920.4(SCN1A):c.3577T>C (p.Trp1193Arg)	AACAAYGGTGAACCTGAGAAGG	Generalized epilepsy with febrile seizures plus, type 1, Generalized epilepsy with febrile seizures plus, type 2
28939717	NM_003907.2(EIF2B5):c.271A>G (p.Thr91Ala)	AAATGYTTCCTGTACACCTGTGG	Leukoencephalopathy with vanishing white matter
80357276	NM_007294.3(BRCA1):c.122A>G (p.His41Arg)	AAATATGYGGTCAACTTTGTGG	Familial cancer of breast, Breast-ovarian cancer, familial 1

397515897	NM_000256.3(MYBPC3):c.1351+2T>C	AAAGGYGGGCTGGGACCTGAGG	Familial hypertrophic cardiomyopathy 4, Cardiomyopathy
397514491	NM_005340.6(HINT1):c.152A>G (p.His51Arg)	AAAAAYGTGTTGGTGCCTGAGGGG, GAAAAAYGTGTTGGTGCCTGAGGG, AGAAAAAYGTGTTGGTGCCTGAGGG	Gamstorp-Wohlfart syndrome
387907164	NM_020894.2(UVSSA):c.94T>C (p.Cys32Arg)	AAAATTYGC AAGTATGTCTTAGG, AAATTYGC AAGTATGTCTTAGG G	UV-sensitive syndrome 3
118161496	NM_025152.2(NUBPL):c.815-27T>C	TGGTTCYAAATGGATGTCTGCTGG, GGTTCYAAATGGATGTCTGCTGGG	Mitochondrial complex I deficiency
764313717	NM_005609.2(PYGM):c.425_528del	TGGCTGYCAGGGACCCAGCAAGG, CTGYCAGGGACCCAGCAAGGAGG	
28934568	NM_003242.5(TGFBR2):c.923T>C (p.Leu308Pro)	AGTTCYACGCGCTGAGGAGCGG	Loeys-Dietz syndrome 2
121913461	NM_007313.2(ABL1):c.814T>C (p.Tyr272His)	CCAGYACGGGGAGGTGTACGAGG, CAGYACGGGGAGGTGTACGAGGG	
377750405	NM_173551.4(ANKS6):c.1322A>G (p.Gln441Arg)	AGGGCYGTCGGACCTTCGAGTGG, GGGCYGTCGGACCTTCGAGTGGG, GGCYGTCGGACCTTCGAGTGGGG	Nephronophthisis 16
57639980	NM_001927.3(DES):c.1034T>C (p.Leu345Pro)	ATTCYCYGATGAGGCAGATGCGG, TTCCYCYGATGAGGCAGATGCGGG	Myofibrillar myopathy 1
147391618	NM_020320.3(RARS2):c.35A>G (p.Gln12Arg)	ATACCYGGCAAGCAATAGCGCGG	Pontocerebellar hypoplasia type 6
182650126	NM_002977.3(SCN9A):c.2215A>G (p.Ile739Val)	GTAAYTGCAAGATCTACAAAAGG	Small fiber neuropathy
80358278	NM_004700.3(KCNQ4):c.842T>C (p.Leu281Ser)	ACATYGACAACCATCGGCTATGG	DFNA 2 Nonsyndromic Hearing Loss
786204012	NM_005957.4(MTHFR):c.388T>C (p.Cys130Arg)	GACCYGCTGCCGTCAGCGCCTGG	Homocysteinemia due to MTHFR deficiency
786204037	NM_005957.4(MTHFR):c.1883T>C (p.Leu628Pro)	TCCCACYGGACAACCTGCTCTGG	Homocysteinemia due to MTHFR deficiency
202147607	NM_000140.3(FECH):c.1137+3A>G	GTAGAYACCTTAGAGAACAATGG	Erythropoietic protoporphyria
122456136	NM_005183.3(CACNA1F):c.2267T>C (p.Ile756Thr)	TGCCAYTGCTGTGGACAACCTGG	
786204851	NM_007374.2(SIX6):c.110T>C (p.Leu37Pro)	GTCGYGCCCGTGGCCCTGCGG	Cataract, microphthalmia and nystagmus
794728167	NM_000138.4(FBN1):c.1468+2T>C	ATTGGYACGTGATCCATCCTAGG	Thoracic aortic aneurysms and aortic dissections
121964909	NM_000027.3(AGA):c.214T>C (p.Ser72Pro)	GACGGCYCTGTAGGCTTTGGAGG	Aspartylglycosaminuria
121964978	NM_000170.2(GLDC):c.2T>C (p.Met1Thr)	CGGCCAYGCAGTCTGTGCCAGG, GGCCAYGCAGTCTGTGCCAGGG	Non-ketotic hyperglycinemia
121965008	NM_000398.6(CYB5R3):c.446T>C (p.Leu149Pro)	CTGCGGTCTACCAGGGCAAAGG	METHEMOGLOBINEMIA, TYPE I
121965064	NM_000128.3(F11):c.901T>C (p.Phe301Leu)	TGATYCTTTGGGAGAAGAACTGG	Hereditary factor XI deficiency disease
45517398	NM_000548.3(TSC2):c.5150T>C (p.Leu1717Pro)	GCCCYGCACGCAAAATGTGAGTGG, CCCYGCACGCAAAATGTGAGTGGG	Tuberous sclerosis syndrome
786205857	NM_015662.2(IFT172):c.770T>C (p.Leu257Pro)	TTGTGCGYAGGAAGTTATGACAGG	RETINITIS PIGMENTOSA 71
786205904	NM_001135669.1(XPR1):c.653T>C (p.Leu218Ser)	GCGTTCYACGTGTCCCCCTTTGG, GCGTTCYACGTGTCCCCCTTTGGG	BASAL GANGLIA CALCIFICATION,
104893704	NM_000388.3(CASR):c.2641T>C (p.Phe881Leu)	ACGCTYTC AAGGTGGCTGCCCGG, CGCTYTC AAGGTGGCTGCCCGGG	Hypercalciuric hypercalcemia
104893747	NM_198159.2(MITF):c.1195T>C (p.Ser399Pro)	ACTTYCCCTATTCCATCCACGG, CTTYCCCTATTCCATCCACGGG	Waardenburg syndrome type 2A
104893770	NM_000539.3(RHO):c.133T>C (p.Phe45Leu)	CATGYTTCGCTGATCGTGCTGG, ATGYTTCGCTGATCGTGCTGGG	Retinitis pigmentosa 4
28937596	NM_003907.2(EIF2B5):c.1882T>C (p.Trp628Arg)	AGGCCYGGAGCCCTGTTTTTAGG	Leukoencephalopathy with vanishing white matter
104893876	NM_001151.3(SLC25A4):c.293T>C (p.Leu98Pro)	GCAGCYCTTCTAGGGGGTGTGG	Autosomal dominant progressive external ophthalmoplegia with mitochondrial DNA deletions 2
104893883	NM_006005.3(WFS1):c.2486T>C (p.Leu829Pro)	ACCATCCYGGAGGGCCGCTGGG	WFS1-Related Disorders
104893962	NM_000165.4(GJA1):c.52T>C (p.Ser18Pro)	CTACYCAACTGCTGGAGGGAAGG	Oculodentodigital dysplasia

104893978	NM_000434.3(NEU1):c.718T>C (p.Trp240Arg)	GCCTCCYGGCGCTACGGAAGTGG, CCTCCYGGCGCTACGGAAGTGGG, CTCCYGGCGCTACGGAAGTGGGG	Sialidosis, type II
104894092	NM_002546.3(TNFRSF11B):c.349T>C (p.Phe117Leu)	TAGAGYTCTGCTTGAACATAGG	Hyperphosphatemia with bone disease
104894135	NM_000102.3(CYP17A1):c.316T>C (p.Ser106Pro)	CATCGCGYCCAACAACCGTAAGG, ATCGCGYCCAACAACCGTAAGGG	Complete combined 17-alpha-hydroxylase/17,20-lyase
104894151	NM_000102.3(CYP17A1):c.1358T>C (p.Phe453Ser)	AGCTCTCYCTCATCATGGCCTGG	Combined partial 17-alpha-hydroxylase/17,20-lyase deficiency
36015961	NM_000518.4(HBB):c.344T>C (p.Leu115Pro)	TGTGTGCGYGGCCATCACTTTGG	Beta thalassemia intermedia
104894472	NM_152443.2(RDH12):c.523T>C (p.Ser175Pro)	TCCYCGTGGCTCACCACATTTGG	Leber congenital amaurosis 13
104894587	NM_004870.3(MPDU1):c.356T>C (p.Leu119Pro)	TTCCYGGTCACTGACTACAGAGG	Congenital disorder of glycosylation type 1F
104894588	NM_004870.3(MPDU1):c.2T>C (p.Met1Thr)	AATAYGGCGCCGAGGCGGACGG	Congenital disorder of glycosylation type 1F
104894626	NM_000304.3(PMP22):c.82T>C (p.Trp28Arg)	TAGCAAYGGATCGTGGCAATGG	Charcot-Marie-Tooth disease, type IE
104894631	NM_018129.3(PNPO):c.784T>C (p.Ter262Gln)	ACCTYAACTCTGGACCTGCTGG	"Pyridoxal 5-phosphate-dependent epilepsy"
104894703	NM_032551.4(KISS1R):c.305T>C (p.Leu102Pro)	GCCCTGCGYGTACCCGCTGCCCGG, TGCYGTACCCGCTGCCCGGCTGG	
104894826	NM_000166.5(GJB1):c.407T>C (p.Val126Ala)	ATGYCATCAGCGTGGTGTCCCG	Dejerine-Sottas disease, X-linked hereditary motor and sensory neuropathy

104894859	NM_001122606.1(LAMP2):c.961T>C (p.Trp321Arg)	CAGCTACYGGGATGCCCCCTGG, AGCTACYGGGATGCCCCCTGGG	Danon disease
104894931	NM_006517.4(SLC16A2):c.1313T>C (p.Leu438Pro)	TGAGCYGGTGGGCCAATGCAGG	Allan-Herndon-Dudley syndrome
104894935	NM_000330.3(RS1):c.38T>C (p.Leu13Pro)	TTACTTCYCTTTGGCTATGAAGG	Juvenile retinoschisis
104895217	NM_001065.3(TNFRSF1A):c.175T>C (p.Cys59Arg)	TGCGYGTACCAAGTGCCACAAAGG	TNF receptor-associated periodic fever syndrome (TRAPS)
143889283	NM_003793.3(CTSF):c.692A>G (p.Tyr231Cys)	CTCCAYACTGAGCTGTGCCACGG	Ceroid lipofuscinosis, neuronal, 13
122459147	NM_001159702.2(FHL1):c.310T>C (p.Cys104Arg)	GGGGYGCTTCAAGGCCATTGTGG	Myopathy, reducing body, X-linked, childhood-onset
74552543	NM_020184.3(CNNM4):c.971T>C (p.Leu324Pro)	AAGCTCCYGGACTTTTTCTGGG	Cone-rod dystrophy amelogenesis imperfecta
199476117	m.10158T>C	AAAYCCACCCCTTACGAGTGCGG	Leigh disease, Leigh syndrome due to mitochondrial complex I deficiency, Mitochondrial complex I deficiency
794727808	NM_020451.2(SEPN1):c.872+2T>C	TTCCGGYGAGTGGGCCACACTGG	Congenital myopathy with fiber type disproportion, Eichsfeld type congenital muscular dystrophy
140547520	NM_005022.3(PFN1):c.350A>G (p.Glu117Gly)	CACCTYCTTTGCCCATCAGCAGG	Amyotrophic lateral sclerosis 18
397514359	NM_000060.3(BTD):c.445T>C (p.Phe149Leu)	TCACCGCYTCAATGACACAGAGG	Biotinidase deficiency
207460001	m.15197T>C	CTAYCCGCCATCCCATACATTTGG	Exercise intolerance
397514406	NM_000060.3(BTD):c.1214T>C (p.Leu405Pro)	TTCACCCYGGTCCCTGTCTGGGG	Biotinidase deficiency
397514516	NM_006177.3(NRL):c.287T>C (p.Met96Thr)	GAGGCCAYGGAGCTGCTGCAGGG	Retinitis pigmentosa 27
72554312	NM_000531.5(OTC):c.134T>C (p.Leu45Pro)	CTCACTCYAAAAAACTTTACCGG	Ornithine carbamoyltransferase deficiency
397514569	NM_178012.4(TUBB2B):c.350T>C (p.Leu117Pro)	GGTCCYGGATGTGGTGGGAAGG	Polymicrogyria, asymmetric
397514571	NM_000431.3(MVK):c.122T>C (p.Leu41Pro)	CGGCYCAACCCACAGCAATGG, GGCYCAACCCACAGCAATGGG	Porokeratosis, disseminated superficial actinic 1
794728390	NM_000238.3(KCNH2):c.2396T>C (p.Leu799Pro)	GCCATCCYGGGTATGGGTGGGG, CCATCCYGGGTATGGGTGGGG, CATCCYGGGTATGGGTGGGGGG	Cardiac arrhythmia
397514713	NM_001199107.1(TBC1D24):c.686T>C (p.Phe229Ser)	GGTCTYTGACGCTTCTCTGGTGG	Early infantile epileptic encephalopathy 16
397514719	NM_080605.3(B3GALT6):c.193A>G (p.Ser65Gly)	CGCYGGCCACCAGCACTGCCAGG	Spondyloepimetaphyseal dysplasia with joint laxity
730880608	NM_000256.3(MYBPC3):c.3796T>C (p.Cys1266Arg)	GAGYCCCGCCTGGAGGTGCGAGG	Cardiomyopathy
397515329	NM_001382.3(DPAGT1):c.503T>C (p.Leu168Pro)	AATCCYGTACTATGTCTACATGG, ATCCYGTACTATGTCTACATGGG, TCCYGTACTATGTCTACATGGGG	Congenital disorder of glycosylation type 1J

397515465	NM_018127.6(ELAC2):c.460T>C (p.Phe154Leu)	ATAYTTTCTGGTCCATTGAAAGG	Combined oxidative phosphorylation deficiency 17
397515557	NM_005211.3(CSF1R):c.2483T>C (p.Phe828Ser)	CATCTYTGACTGTGTCTACACGG	Hereditary diffuse leukoencephalopathy with spheroids
397515599	NM_194248.2(OTOF):c.3413T>C (p.Leu1138Pro)	AGGTGCGYTTCTGGGGCCTACGG, GGTGCGYTTCTGGGGCCTACGGG	Deafness, autosomal recessive 9
397515766	NM_000138.4(FBN1):c.2341T>C (p.Cys781Arg)	GGACAAYGTAGAAATACTCCTGG	Marfan syndrome
565779970	NM_001429.3(EP300):c.3573T>A (p.Tyr1191Ter)	CTTAYTACAGTTACCAGAACAGG	Rubinstein-Taybi syndrome 2
786200938	NM_080605.3(B3GALT6):c.1A>G (p.Met1Val)	AGCTTCAYGGCGCCCGCCGGG, TCAYGGCGCCCGCCGGGCGG	Spondyloepimetaphyseal dysplasia with joint laxity
28942087	NM_000229.1(LCAT):c.698T>C (p.Leu233Pro)	ATCTCTCYTGGGGCTCCCTGGGG, TCTCYTGGGGCTCCCTGGGGTGG	Norum disease
128621203	NM_000061.2(BTK):c.1625T>C (p.Leu542Pro)	TCGGCCYGTCCAGGTGAGTGTGG	X-linked agammaglobulinemia with growth hormone deficiency
397515412	NM_006383.3(CIB2):c.368T>C (p.Ile123Thr)	CTTCACTGCAAGGAGGACCTGG	Deafness, autosomal recessive 48
193929364	NM_000352.4(ABCC8):c.404T>C (p.Leu135Pro)	AAGCYGCTAATTGGTAGGTGAGG	Permanent neonatal diabetes mellitus
730880872	NM_000257.3(MYH7):c.1400T>C (p.Phe467Thr)	TCGAGAYCTTCGATGTGAGTTGG, CGAGAYCTTCGATGTGAGTTGGG	Cardiomyopathy
80356474	NM_002977.3(SCN9A):c.2543T>C (p.Ile848Thr)	AAGATCAYTGGTAACTCAGTAGG, AGATCAYTGGTAACTCAGTAGGG, GATCAYTGGTAACTCAGTAGGGG	Primary erythromelalgia
80356489	NM_001164277.1(SLC37A4):c.352T>C (p.Trp118Arg)	GGGCGGCCCCCATGTGGGAAGG	Glucose-6-phosphate transport defect
80356536	NM_152296.4(ATP1A3):c.2338T>C (p.Phe780Leu)	GCCCYTCTGCTGTTTCATCATGG	Dystonia 12
80356596	NM_194248.2(OTOF):c.3032T>C (p.Leu1011Pro)	GATGCGYGTGTTTCGACAACCTGG	Deafness, autosomal recessive 9, Auditory neuropathy, autosomal recessive, 1

80356689	NM_000083.2(CLCN1):c.857T>C (p.Val286Ala)	AGGAGYGTATTAGCATCGAGG	Myotonia congenita
118203884	m.4409T>C	AGGYCAGCTAAATAAGCTATCGG	Mitochondrial myopathy
58777625	NM_173596.2(SLC39A5):c.911T>C (p.Met304Thr)	AGAACAYGCTGGGGCTTTTGCGG	Myopia 24, autosomal dominant
587783087	NM_003159.2(CDKL5):c.602T>C (p.Leu201Pro)	ATTCYTGGGGAGCTTAGCGATGG	not provided
118203951	NM_013319.2(UBIAD1):c.511T>C (p.Ser171Pro)	TCTGGCYCTTTCTCTACACAGG, GGCYCTTTCTCTACACAGGAGG	Schnyder crystalline corneal dystrophy
118204017	NM_000018.3(ACADVL):c.1372T>C (p.Phe458Leu)	TCGCATCYTCCGGATCTTTGAGG, CGCATCYTCCGGATCTTTGAGGG, GCATCYTCCGGATCTTTGAGGGG	Very long chain acyl-CoA dehydrogenase deficiency
397518466	NM_000833.4(GRIN2A):c.2T>C (p.Met1Thr)	CTAYGGGCAGAGTGGGCTATTGG	Focal epilepsy with speech disorder with or without mental retardation
118204069	NM_000237.2(LPL):c.337T>C (p.Trp113Arg)	GGACYGGCTGTCACGGGCTCAGG	Hyperlipoproteinemia, type I
118204080	NM_000237.2(LPL):c.755T>C (p.Ile252Thr)	GTTGAYTGAGAGAGAGGACTTGG	Hyperlipoproteinemia, type I
118204111	NM_000190.3(HMBS):c.739T>C (p.Cys247Arg)	GCTTCGCGCATCGCTGAAAGGG	Acute intermittent porphyria
80357438	NM_007294.3(BRCA1):c.65T>C (p.Leu22Ser)	AAATCTYAGAGTGTCCCATCTGG	Familial cancer of breast, Breast-ovarian cancer, familial 1, Hereditary cancer-predisposing syndrome
139877390	NM_001040431.2(COA3):c.215A>G (p.Tyr72Cys)	CCAYCTGGGGAGGTAGGTTCCAGG	not provided, Mental retardation, autosomal dominant 31
793888527	NM_005859.4(PURA):c.563T>C (p.Ile188Thr)	GACCAYTGCCTGCCCGCGCAGG, ACCAATGCGCTGCCCGCGCAGGG, CCAYTGCCTGCCCGCGCAGGGG	Hereditary cancer-predisposing syndrome
561425038	NM_002878.3(RAD51D):c.1A>G (p.Met1Val)	CGCCCAAGTTCGCCCGCAGGCCGG	Hereditary cancer-predisposing syndrome
121907934	NM_024105.3(ALG12):c.473T>C (p.Leu158Pro)	TCCYGTGGCCCTCGCGGCTGG	Congenital disorder of glycosylation type 1G
80358207	NM_153212.2(GJB4):c.409T>C (p.Phe137Leu)	CCTCATCYTCAAGGCCGCCGTGG	Erythrokeratoderma variabilis
80358228	NM_002353.2(TACSTD2):c.557T>C (p.Leu186Pro)	TCGGCYGACCCCAAGTTCGTGG	Lattice corneal dystrophy Type III
121908076	NM_138691.2(TMC1):c.1543T>C (p.Cys515Arg)	AGGACCTYGTGGGAAACAATGG, ACCTYGTGGGAAACAATGGTGG, CCTYGTGGGAAACAATGGTGGG	Deafness, autosomal recessive 7

121908089	NM_017838.3(NHP2):c.415T>C (p.Tyr139His)	GGAGGCTYACGATGAGTGCCTGG, GGCTYACGATGAGTGCCTGGAGG	Dyskeratosis congenita autosomal recessive 1, Dyskeratosis congenita, autosomal recessive 2
121908154	NM_001243133.1(NLRP3):c.926T>C (p.Phe309Ser)	GGTGCCTYTGACGAGCACATAGG	Familial cold urticaria, Chronic infantile neurological, cutaneous and articular syndrome
121908158	NM_001033855.2(DCLRE1C):c.2T>C (p.Met1Thr)	GGCGCTAYGAGTTCTTTCGAGGG, GCGCTAYGAGTTCTTTCGAGGGG	Histiocytic medullary reticulosis
796052870	NM_018129.3(PNPO):c.2T>C (p.Met1Thr)	CCCCAYGACGTGCTGGCTGCGG, CCCCAYGACGTGCTGGCTGCGGG, CCCAYGACGTGCTGGCTGCGGGG	not provided
121908318	NM_020427.2(SLURP1):c.43T>C (p.Trp15Arg)	GCAGCCYGGAGCATGGGCTGTGG	Acroerythrodermatid
121908352	NM_022124.5(CDH23):c.5663T>C (p.Phe1888Ser)	CTCACCTYCAACATCACTGCGGG	Deafness, autosomal recessive 12
121908520	NM_000030.2(AGXT):c.613T>C (p.Ser205Pro)	CCTGTACYCGGGCTCCCAGAAGG	Primary hyperoxaluria, type I
121908618	NM_004273.4(CHST3):c.920T>C (p.Leu307Pro)	CGTGYGGCCTCGGCATGGTGG	Spondyloepiphyseal dysplasia with congenital joint dislocations
11694	NM_006432.3(NPC2):c.199T>C (p.Ser67Pro)	TATTCAGYCTAAAAGCAGCAAGG	Niemann-Pick disease type C2
121908739	NM_000022.2(ADA):c.320T>C (p.Leu107Pro)	CCTGCGYGGCCAACTCCAAGTGG	Severe combined immunodeficiency due to ADA deficiency
80359022	NM_000059.3(BRCA2):c.7958T>C (p.Leu2653Pro)	TGCTCTTCAACTAAAATACAGG	Familial cancer of breast, Breast-ovarian cancer, familial 2
121908902	NM_003880.3(WISP3):c.232T>C (p.Cys78Arg)	AAAATCYGTGCCAAGCAACCAGG, AAATCYGTGCCAAGCAACCAGGG, AATCYGTGCCAAGCAACCAGGGG	Progressive pseudorheumatoid dysplasia
121908947	NM_006892.3(DNMT3B):c.808T>C (p.Ser270Pro)	CAAGTTCYCCGAGGTGAGTCCGG, AAGTTCYCCGAGGTGAGTCCGGG, AGTTCYCCGAGGTGAGTCCGGGG	Centromeric instability of chromosomes 1,9 and 16 and immunodeficiency
121909028	NM_000492.3(CFTR):c.3857T>C (p.Phe1286Ser)	AGCCTYTGAGTGATACCACAGG	Cystic fibrosis
121909135	NM_000085.4(CLCNKB):c.1294T>C (p.Tyr432His)	CTTTGTCTATGGTGTGAGTCTGGGG	Barter syndrome type 3
121909143	NM_001300.5(KLF6):c.506T>C (p.Leu169Pro)	GGAGCYGCCCTCGCCAGGGAAGG	
121909182	NM_001089.2(ABCA3):c.302T>C (p.Leu101Pro)	GCACYTGTGATCAACATGCGAGG	Surfactant metabolism dysfunction, pulmonary, 3
121909200	NM_000503.5(EYA1):c.1459T>C (p.Ser487Pro)	CACTCYCGCTCATTCACTCCCGG	Melnick-Fraser syndrome
121909247	NM_004970.2(IGFALS):c.1618T>C (p.Cys540Arg)	GGACYGTGGCTGCCCTCTCAAGG	Acid-labile subunit deficiency
121909253	NM_005570.3(LMAN1):c.2T>C (p.Met1Thr)	AGAYGGCGGGATCCAGGCAAAGG	Combined deficiency of factor V and factor VIII, 1
121909385	NM_000339.2(SLC12A3):c.1868T>C (p.Leu623Pro)	CAACCYGGCCCTCAGTACTCGG	Familial hypokalemia-hypomagnesemia
121909497	NM_002427.3(MMP13):c.224T>C (p.Phe75Ser)	TTCTYCGGCTTAGAGGTGACTGG	Spondyloepimetaphyseal dysplasia, Missouri type
121909508	NM_000751.2(CHRND):c.188T>C (p.Leu63Pro)	AACCYCATCTCCCTGGTGAGAGG	MYASTHENIC SYNDROME, CONGENITAL, 3B, FAST- CHANNEL
121909519	NM_001100.3(ACTA1):c.287T>C (p.Leu96Pro)	CGAGCYTCGCGTGGCTCCCGAGG	Nemaline myopathy 3
121909572	NM_000488.3(SERPINC1):c.667T>C (p.Ser223Pro)	TGGGTGYCCAATAAGACCGAAGG	Antithrombin III deficiency
121909677	NM_000821.6(GGCX):c.896T>C (p.Phe299Ser)	TATGTYCTCTACGTCATGCTGG	Pseudoxanthoma elasticum-like disorder with multiple coagulation factor deficiency
121909727	NM_001018077.1(NR3C1):c.2209T>C (p.Phe737Leu)	CTATTGCYTCCAAACATTTTTGG	Glucocorticoid resistance, generalized
139573311	NM_000492.3(CFTR):c.1400T>C (p.Leu467Pro)	TTCACYTCTAATGGTGATTATGG, TCACYTCTAATGGTGATTATGGG	Cystic fibrosis
121912441	NM_000454.4(SOD1):c.341T>C (p.Ile114Thr)	CATCAYTGGCCGCACACTGGTGG	Amyotrophic lateral sclerosis type 1
121912446	NM_000454.4(SOD1):c.434T>C (p.Leu145Ser)	CGTTYGGCTTGTGGTGAATTGG, GTTYGGCTTGTGGTGAATTGGG	Amyotrophic lateral sclerosis type 1
121912463	NM_000213.3(ITGB4):c.1684T>C (p.Cys562Arg)	GGCCAGYGTGTGTGAGCCTGG	Epidermolysis bullosa with pyloric atresia
121912492	NM_002292.3(LAMB2):c.961T>C (p.Cys321Arg)	CCTCAACYGCGAGCAGTGTACAGG	Nephrotic syndrome, type 5, with or without ocular abnormalities
397516659	NM_001399.4(EDA):c.2T>C (p.Met1Thr)	GGCCAYGGGCTACCCGGAGGTGG	Hypohidrotic X-linked ectodermal dysplasia

111033589	NM_021044.2(DHH):c.485T>C (p.Leu162Pro)	GTTGCGYGGCGCGCCTCGCAGTGG	46,XY gonadal dysgenesis, complete, dhh-related
111033622	NM_000206.2(IL2RG):c.343T>C (p.Cys115Arg)	TGGCYGTCAGTTGCAAAAAAAGG	X-linked severe combined immunodeficiency
121912613	NM_001041.3(SI):c.1859T>C (p.Leu620Pro)	ATGCGYGGAGTTCAGTTTGTGG	Sucrase-isomaltase deficiency
121912619	NM_016180.4(SLC45A2):c.1082T>C (p.Leu361Pro)	GAGTTTCYCATCTACGAAAGAGG	Oculocutaneous albinism type 4
61750581	NM_000552.3(VWF):c.4837T>C (p.Ser1613Pro)	CTGCCYCTGATGAGATCAAGAGG	von Willebrand disease, type 2a
121912653	NM_000546.5(TP53):c.755T>C (p.Leu252Pro)	CATCCYCACCATCATCACACTGG	Li-Fraumeni syndrome 1
111033683	NM_000155.3(GALT):c.386T>C (p.Met129Thr)	AGGTCAYGTGCTCCACCCCTGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
111033752	NM_000155.3(GALT):c.677T>C (p.Leu226Pro)	CAGGAGCYACTCAGGAAGGTGGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
121912729	NM_000039.1(APOA1):c.593T>C (p.Leu198Ser)	GCGCTYGGCCGCGCGCTTGGG	Familial visceral amyloidosis, Ostertag type
769452	NM_000041.3(APOE):c.137T>C (p.Leu46Pro)	AACYGGCACTGGGTGCTTTTGG	
121912762	NM_016124.4(RHD):c.329T>C (p.Leu110Pro)	ACACYGTTCAAGTATTGGGATGG	
111033824	NM_000155.3(GALT):c.1138T>C (p.Ter380Arg)	CGCCYGACCACGCCGACCACAGG, GCCYGACCACGCCGACCACAGGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
111033832	NM_000155.3(GALT):c.980T>C (p.Leu327Pro)	TCCYGCCTCTGCCACTGTCCGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
730881974	NM_000455.4(STK11):c.545T>C (p.Leu182Pro)	GGGAACCYGCTGCTCACCACCGG, AACCYGCTGCTCACCACCGGTGG	Hereditary cancer-predisposing syndrome
1064644	NM_000157.3(GBA):c.703T>C (p.Ser235Pro)	GGGYCACTCAAGGGACAGCCCGG	Gaucher disease
796052090	NM_138413.3(HOGA1):c.533T>C (p.Leu178Pro)	GGACCYGCCTGTGGATGCAGTGG	Primary hyperoxaluria, type III
121913141	NM_000208.2(INSR):c.779T>C (p.Leu260Pro)	CTACCYGGACGGCAGGTGTGTGG	Leprechaunism syndrome
121913272	NM_006218.2(PIK3CA):c.1258T>C (p.Cys420Arg)	GGAACACYGTCCATTGGCATGGG, GAACACYGTCCATTGGCATGGGG	Congenital lipomatous overgrowth, vascular malformations, and epidermal nevi, Neoplasm of ovary, PIK3CA Related Overgrowth Spectrum
61751310	NM_000552.3(VWF):c.8317T>C (p.Cys2773Arg)	GCTCCYGTCTCTCCGACACGG	von Willebrand disease, type 2a
312262799	NM_024408.3(NOTCH2):c.1438T>C (p.Cys480Arg)	TTCACAYGTCTGTGCATGCCAGG	Alagille syndrome 2
121913570	NM_000426.3(LAMA2):c.7691T>C (p.Leu2564Pro)	ATCATTCTTTTGGGAAGTGGAGG, TCATTCTTTTGGGAAGTGGAGGG	Merosin deficient congenital muscular dystrophy
121913640	NM_000257.3(MYH7):c.1046T>C (p.Met349Thr)	AACTCCAYGTATAAGCTGACAGG	Familial hypertrophic cardiomyopathy 1, Cardiomyopathy
121913642	NM_000257.3(MYH7):c.1594T>C (p.Ser532Pro)	CATCATGYCCATCCTGGAAGAGG	Dilated cardiomyopathy 1S
119463996	NM_001079802.1(FKTN):c.527T>C (p.Phe176Ser)	GTAGTCTYTCATGAGAGGAGTGG	Limb-girdle muscular dystrophy -
587776456	NM_002049.3(GATA1):c.1240T>C (p.Ter414Arg)	GCTCAYGAGGGCACAGAGCATGG	GATA-1-related thrombocytopenia with dyserythropoiesis
63750654	NM_000184.2(HBG2):c.-228T>C	ATGCAAAYATCTGTCTGAAACGG	Fetal hemoglobin quantitative trait locus 1
587776519	NM_001999.3(FBN2):c.3725-15A>G	AGCAYTGCAACCACATTGTCCAGG	Congenital contractural arachnodactyly
78365220	NM_000402.4(G6PD):c.473T>C (p.Leu158Pro)	TGCCCYCCACTGGGGTCCACAGG	Anemia, nonspherocytic hemolytic, due to G6PD deficiency
63750741	NM_000179.2(MSH6):c.1346T>C (p.Leu449Pro)	CTGGGGCYGGTATTCATGAAAGG	Hereditary Nonpolyposis Colorectal Neoplasms
587776914	NM_017565.3(FAM20A):c.590-2A>G	GTAATCYGCAAAGGAGGAGAAGG, TAATCYGCAAAGGAGGAGAAGG	Enamel-renal syndrome
5030809	NM_000551.3(VHL):c.292T>C (p.Tyr98His)	CCCYACCAACGCTGCCGCCTGG	Von Hippel-Lindau syndrome, Hereditary cancer-predisposing syndrome
199476132	m.5728T>C	CAATCYACTTCTCCCGCCCGG, AATCYACTTCTCCCGCCCGG	Cytochrome-c oxidase deficiency, Mitochondrial complex I deficiency

62637012	NM_014336.4(AIPL1):c.715T>C (p.Cys239Arg)	CTGCCAGYGCCTGCTGAAGAAGG, CCAGYGCCTGCTGAAGAAGGAGG	Leber congenital amaurosis 4
199476199	NM_207352.3(CYP4V2):c.1021T>C (p.Ser341Pro)	AAACTGGYCTTATACCTGTTGG, AACTGGYCTTATACCTGTTGGG	Bietti crystalline corneoretinal dystrophy
58777183	NM_006702.4(PNPLA6):c.3053T>C (p.Phe1018Ser)	CCTYTAACCCGAGCATCCATCGG	Boucher Neuhauser syndrome
199476389	NM_000487.5(ARSA):c.899T>C (p.Leu300Ser)	GGTCTCTYCGGGTGTGGAAAGGG	Metachromatic leukodystrophy
199476398	NM_016599.4(MYOZ2):c.142T>C (p.Ser48Pro)	TTAYCCCATCTCAGTAACCGTGG	Familial hypertrophic cardiomyopathy 16
119456967	NM_001037633.1(SIL1):c.1370T>C (p.Leu457Pro)	TTGCGYGAAGGAGCTGAGATGAGG	Marinesco-Sjogren syndrome
730882253	NM_006888.4(CALM1):c.268T>C (p.Phe90Leu)	GGCAYTCCGAGTCTTTGACAAGG	Long QT syndrome 14
58777283	NM_012338.3(TSPAN12):c.413A>G (p.Tyr138Cys)	TAATCCAYAATTTGTCATCCTGG	Exudative vitreoretinopathy 5
58777306	NM_015884.3(MBTPS2):c.1391T>C (p.Phe464Ser)	GCTYTGCTTTGGATGGACAATGG	Palmoplantar keratoderma, mutilating, with periorificial keratotic plaques, X-linked
56378716	NM_000250.1(MPO):c.752T>C (p.Met251Thr)	TCACTCAYGTTTCATGCAATGGGG	Myeloperoxidase deficiency
58777390	NM_005026.3(PIK3CD):c.1246T>C (p.Cys416Arg)	GCAGGACYGCCCAATTGCCITGGG	Activated PI3K-delta syndrome
58777480	NM_003108.3(SOX11):c.178T>C (p.Ser60Pro)	TATGGYCCAAGATCGAACCGAGG	Mental retardation, autosomal dominant 27
58777663	NM_001288767.1(ARMC5):c.1379T>C (p.Leu460Pro)	GCCCCGACYGCGGGATGCTGGTGG	Acth-independent macronodular adrenal hyperplasia 2
61753033	NM_000350.2(ABCA4):c.5819T>C (p.Leu1940Pro)	AAGGCYACATGAACTAACCAAGG	Stargardt disease, Stargardt disease 1, Cone- rod dystrophy 3
200488568	NM_002972.3(SBF1):c.4768A>G (p.Thr1590Ala)	CAGGCGYCTCTTGCTCAGCCGG	Charcot-Marie-Tooth disease, type 4B3
132630274	NM_000377.2(WAS):c.809T>C (p.Leu270Pro)	CGGAGTCYGTTCCTCCAGGGCAGG	Severe congenital neutropenia X-linked
132630308	NM_001399.4(EDA):c.181T>C (p.Tyr61His)	CTGCYACCTAGAGTTGCGCTCGG	Hypohidrotic X-linked ectodermal dysplasia
60934003	NM_170707.3(LMNA):c.1589T>C (p.Leu530Pro)	ACGGCTCYCATCAACTCCACTGG, CGGCTCYCATCAACTCCACTGGG, GGCTCYCATCAACTCCACTGGGG	Benign scapulo-peroneal muscular dystrophy with cardiomyopathy
180177160	NM_000030.2(AGXT):c.1076T>C (p.Leu359Pro)	GGTGCYCGGATCGGCCTGCTGG, GTGCYCGGATCGGCCTGCTGGG	Primary hyperoxaluria, type I
180177222	NM_000030.2(AGXT):c.449T>C (p.Leu150Pro)	GTGCYGTCTTCTTAACCCACGG, TGTCYGTCTTCTTAACCCACGGG	Primary hyperoxaluria, type I
180177254	NM_000030.2(AGXT):c.661T>C (p.Ser221Pro)	GCTCATCYCCTCAGTGACAAGG	Primary hyperoxaluria, type I
180177264	NM_000030.2(AGXT):c.757T>C (p.Cys253Arg)	GGGGCYGTGACGACCAGCCAGG	Primary hyperoxaluria, type I
180177293	NM_000030.2(AGXT):c.893T>C (p.Leu298Pro)	GTATCYGCATGGGCGCTGCAGG	Primary hyperoxaluria, type I
376785840	NM_001282227.1(CECR1):c.1232A>G (p.Tyr411Cys)	GAAATCAYAGGACAAGCCTTTGG	Polyarteritis nodosa
587779393	NM_000257.3(MYH7):c.4937T>C (p.Leu1646Pro)	GAGCCYCCAGAGCTTGTGAAGG	Myopathy, distal, 1
587779410	NM_012434.4(SLC17A5):c.500T>C (p.Leu167Pro)	ATTGTACYCAGAGCACTAGAAGG	Sialic acid storage disease, severe infantile type
587779513	NM_000090.3(COL3A1):c.2337+2T>C (p.Gly762_Lys779del)	AGGYAACCTTAATACTACCTGG	Ehlers-Danlos syndrome, type 4
777539013	NM_020376.3(PNPLA2):c.757+2T >C	GAACGGYGCAGCCGCGGGCGGG, AACGGYGCAGCCGCGGGCGGG	Neutral lipid storage disease with myopathy
34557412	NM_012452.2(TNFRSF13B):c.310T>C (p.Cys104Arg)	ACTTCYGTGAGAACAAGCTCAGG	Immunoglobulin A deficiency 2, Common variable
796052970	NM_001165963.1(SCN1A):c.1094T>C (p.Phe365Ser)	CAAGCTYTGATACCTTCAGTTGG, AAGCTYTGATACCTTCAGTTGGG	not provided
724159989	NC_012920.1:m.7505T>C	CCTCCAYGACTTTTCAAAAAGG	Deafness, nonsyndromic sensorineural, mitochondrial
796053222	NM_014191.3(SCN8A):c.4889T>C (p.Leu1630Pro)	CGTCYGCATCAAAGGCGCCAAAGG, GTCYGCATCAAAGGCGCCAAAGGG	not provided
118192127	NM_000540.2(RYR1):c.10817T>C (p.Leu3606Pro)	TACTACCYGGACCAGGTGGGTGG, ACTACCYGGACCAGGTGGGTGGG, CTACCYGGACCAGGTGGGTGGGG	Central core disease
118192170	NM_000540.2(RYR1):c.14693T>C (p.Ile4898Thr)	AGGCAYTGGGGACGAGATCGAGG	Malignant hyperthermia susceptibility type 1, Central core disease

121917703	NM_005247.2(FGF3):c.466T>C (p.Ser156Pro)	GTACGTGYCTGTGAACGGCAAGG, TACGTGYCTGTGAACGGCAAGGG	Deafness with labyrinthine aplasia microtia and microdontia (LAMM)
690016549	NM_005211.3(CSF1R):c.2450T>C (p.Leu817Pro)	CCGCCYGCCTGTGAAGTGGATGG	Hereditary diffuse leukoencephalopathy with spheroids
690016552	NM_005211.3(CSF1R):c.2566T>C (p.Tyr856His)	GAATCCCYACCTGGCATCCTGG	Hereditary diffuse leukoencephalopathy with spheroids
121917738	NM_001098668.2(SFTPA2):c.593T>C (p.Phe198Ser)	GGAGACTYCCGCTACTCAGATGG, GAGACTYCCGCTACTCAGATGGG	Idiopathic fibrosing alveolitis, chronic form
690016559	NM_005211.3(CSF1R):c.1957T>C (p.Cys653Arg)	AGCCYGTACCCATGGAGGTAAGG, GCCYGTACCCATGGAGGTAAGGG	Hereditary diffuse leukoencephalopathy with spheroids
690016560	NM_005211.3(CSF1R):c.2717T>C (p.Ile906Thr)	GCAGAYCTGCTCCTTCTTCAGG	Hereditary diffuse leukoencephalopathy with spheroids
121917769	NM_003361.3(UMOD):c.376T>C (p.Cys126Arg)	GGCCACAYGTGTCATGTGGTGG, GCCACAYGTGTCATGTGGTGGG	Familial juvenile gout
121917773	NM_003361.3(UMOD):c.943T>C (p.Cys315Arg)	ATGGCACYGCCAGTGC AAACAGG	Glomerulocystic kidney disease with hyperuricemia and isosthenuria
121917818	NM_007255.2(B4GALT7):c.617T>C (p.Leu206Pro)	TGCTCTCCAAGCAGCACTACCGG	Ehlers-Danlos syndrome progeroid type
121917824	NM_021615.4(CHST6):c.827T>C (p.Leu276Pro)	GGACCYGGCGGGGAGCCGCTGG	Macular corneal dystrophy Type I
121917848	NM_000452.2(SLC10A2):c.728T>C (p.Leu243Pro)	TTTCYCTGGCTAGAATTGCTGG	Bile acid malabsorption, primary
121918006	NM_000478.4(ALPL):c.1306T>C (p.Tyr436His)	TGGACYATGGTGAGACCTCCAGG	Infantile hypophosphatasia
121918010	NM_000478.4(ALPL):c.979T>C (p.Phe327Leu)	CAAAGGCYCTCTTCTGTGGTGG, GGCYTCTTCTGTGGTGAAGG	Infantile hypophosphatasia
121918088	NM_000371.3(TTR):c.400T>C (p.Tyr134His)	CCCCYACTCCTATTCCACCACGG	
121918110	NM_001042465.1(PSAP):c.1055T>C (p.Leu352Pro)	GAAGCYGCCAAGTCCCTGTCCGG	Gaucher disease, atypical, due to saposin C deficiency
121918137	NM_003730.4(RNASET2):c.550T>C (p.Cys184Arg)	CCAGYGCCTTCCACCAAGCCAGG	Leukoencephalopathy, cystic, without megalencephaly
121918191	NM_001127628.1(FBP1):c.581T>C (p.Phe194Ser)	GGAGTYCATTGGTGACAAGG	Fructose-biphosphatase deficiency
121918306	NM_006946.2(SPTBN2):c.758T>C (p.Leu253Pro)	ACCAAGCYGCTGGATCCCGAAGG, AAGCYGCTGGATCCCGAAGGTGG, AGCYGCTGGATCCCGAAGGTGGG	Spinocerebellar ataxia 5
121918505	NM_000141.4(FGFR2):c.799T>C (p.Ser267Pro)	AATGCCYCCACAGTGGTCCGAGG	Pfeiffer syndrome, Neoplasm of stomach
121918643	NM_003126.2(SPTA1):c.620T>C (p.Leu207Pro)	GTGGAGCYGGTAGCTAAAGAAGG, TGGAGCYGGTAGCTAAAGAAGGG	Hereditary pyropeikilocytosis, Elliptocytosis 2
121918646	NM_001024858.2(SPTB):c.604T>C (p.Trp202Arg)	CTCCAGCYGGAAGGATGGCTTGG	Spherocytosis type 2
121918648	NM_001024858.2(SPTB):c.6055T>C (p.Ser2019Pro)	ATGCCYCTGTGGCTGAGGCGTGG	
727504166	NM_000543.4(SMPD1):c.475T>C (p.Cys159Arg)	TGAGGCCYGTGGCCTGCTCCTGG, GAGGCCYGTGGCCTGCTCCTGGG	Niemann-Pick disease, type A, Niemann-Pick disease, type B
193922915	NM_000434.3(NEU1):c.1088T>C (p.Leu363Pro)	CAGCYATGGCCAGGCCCCAGTGG	Sialidosis, type II
727504419	NM_000501.3(ELN):c.889+2T> C	CAGGYAACATCTGTCCAGCAGG, AGGYAACATCTGTCCAGCAGGG	Supravalvar aortic stenosis
376395543	NM_000256.3(MYBPC3):c.26- 2A>G	GAGACYGAAGGGCCAGGTGGAGG	Primary familial hypertrophic cardiomyopathy, Familial hypertrophic cardiomyopathy 4, Cardiomyopathy
1169305	NM_000545.6(HNF1A):c.1720G>A (p.Gly574Ser)	GATGCGYGGCAGGGTCTGGCTGG, ATGCGYGGCAGGGTCTGGCTGGG, TGCGYGGCAGGGTCTGGCTGGGG	Maturity-onset diabetes of the young, type 3
730880130	NM_000527.4(LDLR):c.1468T>C (p.Trp490Arg)	CTACYGGACCGACTCTGTCTGG, TACYGGACCGACTCTGTCTGGG	Familial hypercholesterolemia
281860286	NM_018713.2(SLC30A10):c.500T>C (p.Phe167Ser)	GGCGCTTYCGGGGGCCTCAGGG	Hyper manganeseemia with dystonia, polycythemia and cirrhosis
730880306	NM_145693.2(LPIN1):c.1441+2T> C	AAGGYACCGGGGCTCGCGCGG, AGGYACCGGGGCTCGCGCGGG	Myoglobinuria, acute recurrent, autosomal recessive
74315452	NM_000454.4(SOD1):c.338T>C (p.Ile113Thr)	TTGCAYCATTGGCCGCACACTGG	Amotrophic lateral sclerosis type 1
730880455	NM_000169.2(GLA):c.41T>C (p.Leu14Pro)	CGCGCYTGGCTTCGCTTCTGG	not provided
267606656	NM_054027.4(ANKH):c.1015T>C (p.Cys339Arg)	AGCTCYGTTTCGTGATGTTTGG	Cranio metaphyseal dysplasia, autosomal dominant

267606687	NM_033409.3(SLC52A3):c.1238T>C (p.Val413Ala)	AGTTACGYCAAGGTGATGCTGGG	Brown-Vialetto-Van laere syndrome
267606721	NM_001928.2(CFD):c.640T>C (p.Cys214Arg)	GGTGYGCGGGGGCGTGTCTGAGG, GTGYGCGGGGGCGTGTCTGAGGG	Complement factor d deficiency
267606747	NM_001849.3(COL6A2):c.2329T>C (p.Cys777Arg)	CGCCYCGACAAGCCACAGCAGG	Ullrich congenital muscular dystrophy
431905515	NM_001044.4(SLC6A3):c.671T>C (p.Leu224Pro)	CTGCACCYCCACCAGAGCCATGG	Infantile Parkinsonism-dystonia
267606857	NM_000180.3(GUCY2D):c.2846T>C (p.Ile949Thr)	AGAGAYCGCCAACATGTCACTGG	Cone-rod dystrophy 6
267606880	NM_022489.3(INF2):c.125T>C (p.Leu42Pro)	GCTGICYCAGATGCCCTCTGTGG	Focal segmental glomerulosclerosis 5
515726191	NM_015713.4(RRM2B):c.581A>G (p.Glu194Gly)	AACTCCTYCTACAGCAGCAAAGG	RRM2B-related mitochondrial disease
267606917	NM_004646.3(NPHS1):c.793T>C (p.Cys265Arg)	GCTGCCGYGCGTGGCCGAGGGG, CTGCCGYGCGTGGCCGAGGGG	Finnish congenital nephrotic syndrome
267607104	NM_001199107.1(TBC1D24):c.751T>C (p.Phe251Leu)	CAAGTTCYTCCACAAGGTGAGGG, TTCYTCCACAAGGTGAGGGCCGG	Myoclonic epilepsy, familial infantile
267607182	NM_144631.5(ZNF513):c.1015T>C (p.Cys339Arg)	TGGGCGCYGCATGCGAGGAGAGG, CGCYGCATGCGAGGAGAGGCTGG	Retinitis pigmentosa 58
267607211	NM_000229.1(LCAT):c.508T>C (p.Trp170Arg)	TATGACYGGCGGCTGGAGCCCGG	Norum disease
267607215	NM_016269.4(LEF1):c.181T>C (n.Ser61Pro)	GAACGAGYCTGAAATCATCCCGG	Sebaceous tumors, somatic
587783580	NM_178151.2(DCX):c.683T>C (n.Leu278Pro)	AAAAAACYCTACTCTGGATGG	Heterotopia
587783644	NM_004004.5(GJB2):c.107T>C (p.Leu36Pro)	GATCCYCGTTGTGGCTGCAAAGG	Hearing impairment
587783653	NM_005682.6(ADGRG1):c.1460T>C (p.Leu487Pro)	CCCTGCYCACCTGCCTTTCCTGG	Polymicrogyria, bilateral frontoparietal
587783863	NM_000252.2(MTM1):c.958T>C (p.Ser320Pro)	GGAACTTTAAAAAAGTGAAGG	Severe X-linked myotubular myopathy
267607751	NM_000249.3(MLH1):c.453+2T>C	ATCACGGYAAAGAATGGTACATGG, TCACGGYAAAGAATGGTACATGGG	Hereditary Nonpolyposis Colorectal Neoplasms
119103227	NM_000411.6(HLCS):c.710T>C (p.Leu237Pro)	CTATCYTTCACGGGAGGGAAGG	Holocarboxylase synthetase deficiency
119103237	NM_005787.5(ALG3):c.211T>C (n.Trp71Arg)	GATTGACYGGAAGGCCTACATGG	Congenital disorder of glycosylation type 1D
398122806	NM_003172.3(SURF1):c.679T>C (p.Trp227Arg)	CCACYGGCATTATCGAGACCTGG	Congenital myasthenic syndrome, acetazolamide-responsive
80338747	NM_004525.2(LRP2):c.7564T>C (p.Tyr2522His)	GTACCTGYACTGGGCTGACTGGG	Donnai Barrow syndrome
398122838	NM_001271723.1(FBXO38):c.616T>C (p.Cys206Arg)	TTCTCYGTATCCAATGCTAAGG	Distal hereditary motor neuropathy 2D
398122989	NM_014495.3(ANGPTL3):c.883T>C (p.Phe295Leu)	ACAAAACYTCAATGAAACGTGGG	Hypobetalipoproteinemia, familial, 2
80338945	NM_004004.5(GJB2):c.269T>C (p.Leu90Pro)	GCTCCYAGTGGCCATGCACGTGG	Deafness, autosomal recessive 1A, Hearing impairment
80338956	NM_000334.4(SCN4A):c.2078T>C (p.Ile693Thr)	AAGATCAITGGCAATTCAGTGGG, AGATCAITGGCAATTCAGTGGGG, GATCAITGGCAATTCAGTGGGGG	Hyperkalemic Periodic Paralysis Type 1, Paramyotonia congenita of von Eulenburg
267608131	NM_000179.2(MSH6):c.4001+2T>C	CGGYAACTAACTATAATGG	Hereditary Nonpolyposis Colorectal Neoplasms
587784573	NM_004963.3(GUCY2C):c.2782T>C (p.Cys928Arg)	TCCYGTGCTGCTGGAGTTGTGG, CCCYGTGCTGCTGGAGTTGTGGG	Meconium ileus
267608511	NM_003159.2(CDKL5):c.659T>C (p.Leu220Pro)	CCAACTTTTACTATTGAGAAGG	Early infantile epileptic encephalopathy 2
373842615	NM_000118.3(ENG):c.1273- 2A>G	CGCCYCGGGGATAAAGCCAGG, CGCCYCGGGGATAAAGCCAGGG	Haemorrhagic telangiectasia 1
185492581	NM_000335.4(SCN5A):c.376A>G (p.Lys126Glu)	GAATCTYCACAGCCGCTCTCCGG	Brugada syndrome
200533370	NM_133499.2(SYN1):c.1699A>G (p.Thr567Ala)	GATGYCTGACGGTAGCCTGTGG, ATGYCTGACGGTAGCCTGTGGG	Epilepsy, X-linked, with variable learning disabilities and behavior disorders, not specified
118203981	NM_148960.2(CLDN19):c.269T>C (p.Leu90Pro)	GCTCCYGGGCTTCGTGGCCATGG	Hypomagnesemia 5, renal, with ocular involvement
137853892	NM_001235.3(SERPINH1):c.233T>C (p.Leu78Pro)	GTCGYAGGGCTCGTGTCTGCTGG, TCGCYAGGGCTCGTGTCTGCTGGG	Osteogenesis imperfecta type 10
118204024	NM_000263.3(NAGLU):c.142T>C	GGCCGACYTCTCCGTGTCTGGTGG	Mucopolysaccharidosis, MPS-III-B
690016563	NM_005211.3(CSF1R):c.1745T>C (p.Leu582Pro)	CAACCYGCAGTTTGGTGAGATGG	Hereditary diffuse leukoencephalopathy with spheroids
58380626	NM_000526.4(KRT14):c.1243T>C (p.Tyr415His)	CGCCACCYACCGCCGCTGTGG, CACCYACCGCCGCTGTGGAGG, ACCYACCGCCGCTGTGGAGGG	Epidermolysis bullosa herpetiformis, Dowling-Meara

113994151	NM_207346.2(TSEN54):c.277T>C (p.Ser93Pro)	TTGAAGYCTCCCGGGTGAAGCGG, AAGYCTCCCGGGTGAAGCGCGG	Pontocerebellar hypoplasia type 4
113994206	NM_004937.2(CTNS):c.473T>C (o.Leu158Pro)	TGGTCYAGCTTCGACTTCGTGG	Cystinosis
62516109	NM_000277.1(PAH):c.638T>C (p.Leu213Pro)	CCACTTCYTGAAAAGTACTGTGG	Phenylketonuria
370011798	NM_001302946.1(TRNT1):c.668T>C (p.Ile223Thr)	GCAAYTGCAAAAATGCAAAAAGG	Sideroblastic anemia with B-cell immunodeficiency, periodic fevers, and developmental delay
62517167	NM_000277.1(PAH):c.293T>C (p.Leu98Ser)	AAGATCTYGAGGCATGACATTGG	Mild non-PKU hyperphenylalanemia
12021720	NM_001918.3(DBT):c.1150G>A (p.Gly384Ser)	GACYCACAGAGCCCAATTTCTGG	Intermediate maple syrup urine disease type 2
104886289	NM_000495.4(COL4A5):c.4756T>C (p.Cys1586Arg)	TCCCCATYGTCTCAGGGATGGG	Alport syndrome, X-linked recessive
370471013	NC_012920.1:m.5559A>G	CAACYTACTGAGGGCTTTGAAGG	Leigh disease
121434215	NM_000487.5(ARSA):c.410T>C (p.Leu137Pro)	GCCTTCCYGGCCCCCATCAGGG	Metachromatic leukodystrophy, adult type
386134128	NM_000096.3(CP):c.1123T>C (o.Tyr375His)	ACACTACYACATTGCCGCTGAGG	Deficiency of ferroxidase
121434275	NM_001127328.2(ACADM):c.1136T>C (p.Ile379Thr)	GTGCAGAYACTTGGAGGCAATGG	Medium-chain acyl-coenzyme A dehydrogenase deficiency
121434276	NM_001127328.2(ACADM):c.742T>C (p.Cys248Arg)	CAGCGAYGTTAGATACTAGAGG	Medium-chain acyl-coenzyme A dehydrogenase deficiency
121434284	NM_002225.3(IVD):c.134T>C (p.Leu45Pro)	ATGGGCYAAGCGAGGAGCAGAGG	ISOVALERIC ACIDEMIA, TYPE I
121434334	NM_005908.3(MANBA):c.1513T>C (p.Ser505Pro)	ATTACGYCCAGTCTACAAATGG, TTACGYCCAGTCTACAAATGGG, TACGYCCAGTCTACAAATGGGG CGCCCGGYACGGCATCGCTGGG, GCCCGGYACGGCATCGCTGGGG	Beta-D-mannosidosis
121434366	NM_000159.3(GCDH):c.883T>C (p.Tyr295His)	CGCCCGGYACGGCATCGCTGGG, GCCCGGYACGGCATCGCTGGGG	Glutaric aciduria, type 1
60715293	NM_000424.3(KRT5):c.541T>C (p.Ser181Pro)	GTTTGCCYCCCTTCATCGACAAGG	Epidermolysis bullosa herpetiformis, Dowling-Meara
121434409	NM_001003722.1(GLE1):c.2051T>C (p.Ile684Thr)	AAGGACAYTCCTGTCCCAAGGG	Lethal arthrogyposis with anterior horn cell disease
121434434	NM_001287.5(CLCN7):c.2297T>C (p.Leu766Pro)	GGGCCYCGGCACCTGGTGGTGG	Osteopetrosis autosomal recessive 4
121434455	NM_000466.2(PEX1):c.1991T>C (p.Leu664Pro)	GATGACCYTGACCTCATTGCTGG	Zellweger syndrome
199422317	NM_001099274.1(TINF2):c.862T>C (p.Phe288Leu)	CTGYTTCCCTTTAGGAATCTCGG	Aplastic anemia
104895221	NM_001065.3(TNFRSF1A):c.349T>C (p.Cys117Arg)	CTCTTCTYGCACAGTGGACCGGG	TNF receptor-associated periodic fever syndrome (TRAPS)
137854459	NM_000138.4(FBN1):c.4987T>C (p.Cys1663Arg)	GGGACAYGTTACAACACCGTTGG	Marfan syndrome
387907075	NM_024027.4(COLEC11):c.505T>C (p.Ser169Pro)	CAGCTGYCCTGCCAGGGCCCGGG, AGCTGYCCTGCCAGGGCCCGGGG, GCTGYCCTGCCAGGGCCCGGGG, CTGYCCTGCCAGGGCCCGGGGG	Carnevale syndrome
1048095	NM_000352.4(ABCC8):c.674T>C (p.Leu225Pro)	TGCGYTCAAAAGGCACCTACTGG	Permanent neonatal diabetes mellitus
796065347	NM_019074.3(DLL4):c.1168T>C (p.Cys390Arg)	GAAYGTCCCCCAACTTCACCGG	Adams-Oliver syndrome, ADAMS-OLIVER SYNDROME 6
137852347	NM_000402.4(G6PD):c.1054T>C (p.Tyr352His)	AGGGYACCTGGACGACCCACGG	Anemia, nonspherocytic hemolytic, due to G6PD deficiency
74315327	NM_213653.3(HFE2):c.302T>C (p.Leu101Pro)	GGACCYGCCTTCCATTCGGCGG	Hemochromatosis type 2A
137852579	NM_000044.3(AR):c.2033T>C (p.Leu678Pro)	GTCCYGGAAGCCATTGAGCCAGG	
137852636	NM_001166107.1(HMGCS2):c.520T>C (p.Phe174Leu)	CCCTCYTCAATGCTGCCAACTGG	mitochondrial 3-hydroxy-3-methylglutaryl-CoA synthase deficiency
137852661	NM_033163.3(FGF8):c.118T>C (p.Phe40Leu)	TTCCCTGYTCCGGGTGGCCGGG	Kallmann syndrome 6
121912967	NM_005215.3(DCC):c.503T>C (p.Met168Thr)	AGCCCAAGCCAAACATCCACTGG	
137852806	NM_001039523.2(CHRNA1):c.901T>C (p.Phe301Leu)	TGTGYTCCTTCTGGTTCATCGTGG	Myasthenic syndrome, congenital, fast-channel
137852850	NM_182760.3(SUMF1):c.463T>C (p.Ser155Pro)	GGCGACYCCTTTGTCTTTGAAGG	Multiple sulfatase deficiency

137852886	NM_000158.3(GBE1):c.671T>C (p.Leu224Pro)	AATGTACYACCAAGAATCAAAGG	Glycogen storage disease, type IV, GLYCOGEN STORAGE DISEASE IV, NONPROGRESSIVE HEPATIC
137852911	NM_000419.3(ITGA2B):c.641T>C (p.Leu214Pro)	CTGGTGCYGGGGCTCTGGCGG	Glanzmann thrombasthenia
137852948	NM_138694.3(PKHD1):c.10658T>C (p.Ile3553Thr)	GAGCCCAYTGAAATACGCTCAGG	Polycystic kidney disease, infantile type
137852964	NM_024960.4(PANK2):c.178T>C (p.Ser60Pro)	ATTGACYCAGTCGGATTCAATGG	
137853020	NM_006899.3(IDH3B):c.395T>C (p.Leu132Pro)	TGCGGCYAGGTAGGTGGTCTGG, GCGGCYAGGTAGGTGGTCTGGG	Retinitis pigmentosa 46
137853249	NM_033500.2(HK1):c.1550T>C (p.Leu517Ser)	GACTTCTYGGCCCTGGATCTTGG, TTCTYGGCCCTGGATCTTGGAGG	Hemolytic anemia due to hexokinase deficiency
137853270	NM_000444.5(PHEX):c.1664T>C (p.Leu555Pro)	AGCYCCAGAAGCCTTTCTTTTGG	Familial X-linked hypophosphatemic vitamin D refractory rickets
137853325	NM_003639.4(IKBKG):c.1249T>C (p.Cys417Arg)	TGGAGYGCATTGAGTAGGGCCCG	Hypohidrotic ectodermal dysplasia with immune deficiency, Hyper-IgM immunodeficiency, X- linked, with hypohidrotic ectodermal dysplasia
28932769	NM_002055.4(GFAP):c.1055T>C (p.Leu352Pro)	GGACCYGCCTCAATGTCAAGCTGG	Alexander disease
397507439	NM_002769.4(PRSS1):c.116T>C (p.Val39Ala)	TACCAGGYGTCCTGAATTCTGG	Hereditary pancreatitis
387906446	NM_000132.3(F8):c.1729T>C (p.Ser577Pro)	AAAGAAYCTGTAGATCAAAGAGG	Hereditary factor VIII deficiency disease
387906482	NM_000133.3(F9):c.1031T>C (p.Ile344Thr)	ACGAACAYCTTCTCAAATTTGG	Hereditary factor IX deficiency disease
387906508	NM_000131.4(F7):c.983T>C (p.Phe328Ser)	GACGYCTCTGAGAGGACGCTGG	Factor VII deficiency
387906532	NM_001040113.1(MYH11):c.3791T>C (p.Leu1264Pro)	GAAGCYGGAGGCGCAGGTGCAGG	Aortic aneurysm, familial thoracic 4
387906658	NM_002465.3(MYBP1):c.2566T>C (p.Tyr856His)	CAAACCYATATCCGAGAGTTGG	Distal arthrogyrosis type 1B
387906701	NM_003491.3(NAA10):c.109T>C (p.Ser37Pro)	TGGCCTTYCCTGGCCCCAGGTGG, GGCCTTYCCTGGCCCCAGGTGGG	N-terminal acetyltransferase deficiency
387906717	NM_000377.2(WAS):c.881T>C (p.Ile294Thr)	GACTTCAYTGAGGACCAAGGTGG, ACTTCAYTGAGGACCAAGGTGGG	Severe congenital neutropenia X-linked
387906809	NM_000287.3(PEX6):c.1601T>C (p.Leu534Pro)	CTTCYGGGCCGGGACCGTGATGG, TTCYGGGCCGGGACCGTGATGGG	Peroxisome biogenesis disorder 4B
387906965	NM_024513.3(FYCO1):c.4127T>C (p.Leu1376Pro)	CAGCCYATCCCATCACTGTGG	Cataract, autosomal recessive congenital 2
387906967	NM_006147.3(IRF6):c.65T>C (p.Leu222Pro)	GCCYCTACCTGGGCTCATCTGG	Van der Woude syndrome, Poptiteal pterygium syndrome
387906982	NM_025132.3(WDR19):c.20T>C (p.Leu7Pro)	TCTCACYGCTAGAAAAGACTTGG	Asphyxiating thoracic dystrophy 5
387907072	NM_032446.2(MEGF10):c.2320T>C (p.Cys774Arg)	GGGCAGYGTACTTGCCGCACTGG	Myopathy, areflexia, respiratory distress, and dysphagia, early-onset, Myopathy, areflexia, respiratory distress, and dysphagia, early- onset, mild variant
137854499	NM_005502.3(ABCA1):c.6026T>C (p.Phe2009Ser)	GAGTYCTTTGCCTTTTGAGAGG	Familial hypoalphalipoproteinemia
387907117	NM_000196.3(HSD11B2):c.1012T>C (p.Tyr338His)	CCGCCGCYATTACCCCGCCAGG, CGCCGCYATTACCCCGCCAGGG	Apparent mineralocorticoid excess
387907170	NM_004453.3(ETFDH):c.1130T>C (p.Leu377Pro)	CCAAAACYACCTTTCTTGGTGG	
387907205	NM_033360.3(KRAS):c.211T>C (p.Tyr71His)	GGACCAGYACATGAGGACTGGGG, CCAGYACATGAGGACTGGGGAGG, CAGYACATGAGGACTGGGGAGGG	Cardiofaciocutaneous syndrome 2
387907240	NM_024110.4(CARD14):c.467T>C (p.Leu156Pro)	CAGCAGCYGCAGGAGCACCTGGG	Pityriasis rubra pilaris
387907282	NM_152296.4(ATP1A3):c.2431T>C (p.Ser811Pro)	TGCCATCYCACTGGCGTACGAGG	Alternating hemiplegia of childhood 2
387907361	NM_005120.2(MED12):c.3493T>C (p.Ser1165Pro)	AGGACYCTGAGCCAGGGGCCCGG	Ohdo syndrome, X-linked
28933970	NM_006194.3(PAX9):c.62T>C (p.Leu21Pro)	GGCCGCYGCCAACGCCATCCGG	Toothagenesis, selective, 3
137854472	NM_000138.4(FBN1):c.3128A>G (p.Lys1043Arg)	TGCACYTGCCGTGGGTGCAGAGG	
727504261	NM_000257.3(MYH7):c.2708A>G (p.Glu903Gly)	AGCGCYCCTCAGCATCTGCCAGG	Cardiomyopathy, not specified
81002853	NM_000059.3(BRCA2):c.476- 2A>G	ACCACYGGGGTAAAAAAGGGG, TACCACYGGGGTAAAAAAGGGG	Familial cancer of breast, Breast-ovarian cancer, familial 2, Hereditary cancer- predisposing syndrome

119473032	NM_021020.3(LZTS1):c.355A>G (p.Lys119Glu)	CCCTYCTCGGAGCCCTGTAGAGG	
193922801	NM_000540.2(RYR1):c.7043A>G (p.Glu2348Gly)	TTCYCCTCCACGCTCTCGCCTGG	not provided
36210419	NM_000218.2(KCNQ1):c.652A>G (p.Lys218Glu)	GCCCCYGGAGCCCACGCAGAGG	Torsades de pointes, Cardiac arrhythmia
121964989	NM_000108.4(DLD):c.1483A>G (p.Arg495Gly)	TTCTCYAAAAGCTTCTGATAAGG	Maple syrup urine disease, type 3
28936669	NM_000095.2(COMP):c.1418A>G (p.Asp473Gly)	ATTGYCGTCGTCGTCGTCGAGG	
28936696	NM_018488.2(TBX4):c.1592A>G (p.Gln531Arg)	GTACYGTAAGGAAGATTCTCGGG, GGTACYGTAAGGAAGATTCTCGG	Ischiopatellar dysplasia
121965077	NM_000137.2(FAH):c.1141A>G (p.Arg381Gly)	TCCYGGTCTGACCATTCCCCAGG	Tyrosinemia type I
794728203	NM_000138.4(FBN1):c.3344A>G (p.Asp1115Gly)	ACTCAYCAATATCTGCAAAATGG	Thoracic aortic aneurysms and aortic dissections
786205436	NM_003002.3(SDHD):c.275A>G (p.Asp92Gly)	GAATAGYCCATCGCAGAGCAAGG	Fatal infantile mitochondrial cardiomyopathy
72551317	NM_000784.3(CYP27A1):c.776A>G (p.Lys259Arg)	AGTCCACYTGGGGAGGAAGGTGG	Cholesterol storage disease

786205687	NM_016218.2(POLK):c.1385A>G (p.Asn462Ser)	ATTACAYTCTTCAACTTAATGG	Malignant tumor of prostate
794728280	NM_000138.4(FBN1):c.7916A>G (p.Tyr2639Cys)	TGTTTCACTGGAAGCCGGCGGG, CTGTTCACTGGAAGCCGGCGGG	Thoracic aortic aneurysms and aortic dissections
28937317	NM_000335.4(SCN5A):c.3971A>G (p.Asn1324Ser)	GCAYTGACCACCACCTCAAGTGG	Long QT syndrome 3, Congenital long QT syndrome
786205854	NM_144499.2(GNAT1):c.386A>G (p.Asp129Gly)	CGGAGYCCTTCCACAGCCGCTGG	NIGHT BLINDNESS, CONGENITAL
104893776	NM_000539.3(RHO):c.533A>G (p.Tyr178Cys)	GGATGYACCTGAGGACAGGCAGG	Retinitis pigmentosa 4
28937590	NM_001257342.1(BCSIL):c.232A>G (p.Ser78Gly)	GACACYGAGGTGCTGAGTACGGG, CGACACYGAGGTGCTGAGTACGG	GRACILE syndrome
104893866	NM_000320.2(QDPR):c.449A>G (p.Tyr150Cys)	TGCCGYACCCGATCATACTGGG, ATGCCGYACCCGATCATACTGG	Dihydropteridine reductase deficiency
587776590	NM_015629.3(PRPF31):c.527+3A>G	GACAYACCCCTGGGTGGTGGAGG, GCGGACAYACCCCTGGGTGGTGG	Retinitis pigmentosa 11
104894015	NM_000162.3(GCK):c.641A>G (p.Tyr214Cys)	GTAGYAGCAGGAGATCATCGTGG	Hyperinsulinemic hypoglycemia familial 3
202247823	NM_000532.4(PCCB):c.1606A>G (p.Asn536Asp)	ATATYTGATGTTTTCTCCAAGG	Propionic acidemia
104894199	NM_000073.2(CD3G):c.1A>G (p.Met1Val)	CCAYGTCAGTCTCTGTCTCCGG	Immunodeficiency 17
104894208	NM_001814.4(CTSC):c.857A>G (p.Gln286Arg)	CTCCYAGGGCTTAGGATTGGGG, CCTCCYAGGGCTTAGGATTGGG, ACCTCCYAGGGCTTAGGATTGG	Papillon-Lefevre syndrome, Haim-Munk syndrome
104894211	NM_001814.4(CTSC):c.1040A>G (p.Tyr347Cys)	TCCTACAYAGTGGTACTCAGAGG	Papillon-Lefevre syndrome, Periodontitis
104894290	NM_000448.2(RAG1):c.2735A>G (p.Tyr912Cys)	CTGYACTGGCAGAGGGATTCTGG	Histiocytic medullary reticulosis
104894354	NM_000217.2(KCNA1):c.676A>G (p.Thr226Ala)	GCGYTTCCACGATGAAGAAGGGG, AGCGYTTCCACGATGAAGAAGGG, CAGCGYTTCCACGATGAAGAAGG	Episodic ataxia type 1
104894425	NM_014239.3(EIF2B2):c.638A>G (p.Glu213Gly)	AGTGTGTCYAACTGCTTTGG	Leukoencephalopathy with vanishing white matter, Ovarioleukodystrophy
104894450	NM_000270.3(PNP):c.383A>G (p.Asp128Gly)	ATAYCTCCAACCTCAAACCTGGG, GATAYCTCCAACCTCAAACCTGG	Purine-nucleoside phosphorylase deficiency
147394623	NM_024887.3(DHDDS):c.124A>G (p.Lys42Glu)	GGCACTYCTTGGCATAGCGACGG	Retinitis pigmentosa 59
60723330	NM_005557.3(KRT16):c.374A>G (p.Asn125Ser)	GCGGTCAYTGAGGTTCTGCATGG	Pachyonychia congenita, type 1, Palmoplantar keratoderma, nonepidermolytic, focal
104894634	NM_030665.3(RAI1):c.4685A>G (p.Gln1562Arg)	CTGCTGCGYTCGTCGTCGCTTGG	Smith-Magenis syndrome
104894730	NM_000363.4(TNNB3):c.532A>G (p.Lys178Glu)	CCTYCTTACCTTGGAGGTTGG, CCTCCTYCTTACCTTGGAGG	Familial restrictive cardiomyopathy 1
104894816	NM_002049.3(GATA1):c.653A>G (p.Asp218Gly)	GTCCTGYCCCTCCGCCACAGTGG	GATA-1-related thrombocytopenia with dyserythropoiesis
794726773	NM_001165963.1(SCN1A):c.1662+3A>G	GTGCCAYACCTGGTGTGGGGAGG	Severe myoclonic epilepsy in infancy

104894861	NM_000202.6(IDS):c.404A>G (p.Lys135Arg)	AAAGACTYTTCCACCCGACATGG	Mucopolysaccharidosis, MPS-II
104894874	NM_000266.3(NDP):c.125A>G (p.His42Arg)	TGGYGCCCTCATGCAGCGTCGAGG	
191205969	NM_002420.5(TRPM1):c.296T>C (p.Leu99Pro)	AAGCYCTTAATATCTGTGCATGG	Congenital stationary night blindness, type 1C
794727073	NM_019109.4(ALG1):c.1188-2A>G	TAAACYGCAGAGAGAACCAAGGG, GTAAACYGCAGAGAGAACCAAG G	Congenital disorder of glycosylation type 1K
281875236	NM_001004334.3(GPR179):c.659A>G (p.Tyr220Cys)	CCCACAYATCCATCTGCCTGCGG	Congenital stationary night blindness, type 1E
28939094	NM_015915.4(ATL1):c.1222A>G (p.Met408Val)	CACCCAYCTTCTCACCCCTCGG	Spastic paraplegia 3
281875324	NM_005359.5(SMAD4):c.989A>G (p.Glu330Gly)	ATCCATTYCAAAGTAAGCAATGG	Juvenile polyposis syndrome, Hereditary cancer-predisposing syndrome
77173848	NM_000037.3(ANK1):c.-1087T>C	GGGCCYGGCCCGCACGTCACAGG	Spherocytosis, type 1, autosomal recessive
150181226	NM_001159772.1(CANT1):c.671T>C (p.Leu224Pro)	CGTCYGTACGTGGGGCCTGGG, GCGTCYGTACGTGGGGCCTGG	Desbuquois syndrome
397514253	NM_000041.3(APOE):c.237-2A>G	CGCCCYGCGCCGAGAGGGCCGG, GCGCCCYGCGCCGAGAGGGCCGG	Familial type 3 hyperlipoproteinemia
397514348	NM_000060.3(BTD):c.278A>G (p.Tyr93Cys)	GTTCAAYAGATGTCAAGGTTCTGG	Biotinidase deficiency
397514415	NM_000060.3(BTD):c.1313A>G (p.Tyr438Cys)	GGCAYACAGCTCTTTGGATAAAG	Biotinidase deficiency
397514501	NM_007171.3(POMT1):c.430A>G (p.Asn144Asp)	GAGCATYCTCTGTTTCAAAGAGG	Limb-girdle muscular dystrophy-
370382601	NM_174917.4(ACSF3):c.1A>G (p.Met1Val)	GGCAGCAITGCACTGACAGGCGG	not provided
72554332	NM_000531.5(OTC):c.238A>G (p.Lys80Glu)	AAGGACTYCCCTTGCAATAAAGG	Ornithine carbamoyltransferase deficiency
397514599	NM_033109.4(PNPT1):c.1424A>G	GACTYCAGATGTAACTCTTATGG	Deafness, autosomal recessive 70

	(p.Glu475Gly)		
397514650	NM_000108.4(DLD):c.1444A>G (p.Arg482Gly)	GACTCYAGCTATATCTTACAGG	Maple syrup urine disease, type 3
397514675	NM_003156.3(STIM1):c.251A>G (p.Asp84Gly)	TTCCACAYCCACATCACCATTGG	Myopathy with tubular aggregates
794728378	NM_000238.3(KCNH2):c.1913A>G (p.Lys638Arg)	ATCYTCTCTGAGTTGGTGTGGG, GATCYTCTCTGAGTTGGTGTGG	Cardiac arrhythmia
397514711	NM_002163.2(IRF8):c.238A>G (p.Thr80Ala)	AACCTCGYCTTCCAAGTGGCTGG	Autosomal dominant CD11C+/CD135+ dendritic cell deficiency
397514729	NM_000388.3(CASR):c.85A>G (p.Lys29Glu)	CCCCCTYCTTTGGGCTCGCTGG	Hypocalcemia, autosomal dominant 1, with barrter syndrome
397514743	NM_022114.3(PRDM16):c.2447A>G (p.Asn816Ser)	GCCGCCGYTTTGGCTGGCACGGG	Left ventricular noncompaction 8
397514757	NM_005689.2(ABCB6):c.508A>G (p.Ser170Gly)	TGGGCGYGTCCAAGACACCAGGG, GTGGGCGYGTCCAAGACACCAGG	Dyschromatosis universalis hereditaria 3
28940313	NM_152443.2(RDH12):c.677A>G (p.Tyr226Cys)	CACTGCGYAGGTGGTGACCCCGG	Leber congenital amaurosis 13
794728538	NM_000218.2(KCNQ1):c.1787A>G (p.Glu596Gly)	GTCYCTACTCGGTTACAGGCGGG, TGCTYCTACTCGGTTACAGGCGG	Cardiac arrhythmia
794728569	NM_000218.2(KCNQ1):c.605A>G (p.Asp202Gly)	AGGYCTGTGGAGTGCAGGAGAGG	Cardiac arrhythmia
794728573	NM_000218.2(KCNQ1):c.1515-2A>G	GCCYGCAGTGGAGAGAGGAGAGG	Cardiac arrhythmia
370874727	NM_003494.3(DYSF):c.3349-2A>G	CCGCCYGGAGACACGAAGCTGG	Limb-girdle muscular dystrophy, type 2B
794728859	NM_198056.2(SCN5A):c.2788-2A>G	ACCYGTCGAGATAATGGGTCAGG	not provided
794728887	NM_198056.2(SCN5A):c.4462A>G (p.Thr1488Ala)	CCTCTGYCATGAAGATGTCCTGG	not provided
28940878	NM_000372.4(TYR):c.125A>G (p.Asp42Gly)	CTCCTGYCCCCGCTCCACGGTGG	Tyrosinase-negative oculocutaneous albinism
397515420	NM_172107.2(KCNQ2):c.1636A>G (p.Met546Val)	GCAYGACACTGCAGGGGGGTGGG, CGCAYGACACTGCAGGGGGGTGG, AACCGCAYGACACTGCAGGGGGG	Early infantile epileptic encephalopathy 7

397515428	NM_001410.2(MEGF8):c.7099A>G (p.Ser2367Gly)	GACYCCCGTGAAATGATCCCGG	Carpenter syndrome 2
143601447	NM_201631.3(TGM5):c.122T>C (p.Leu41Pro)	TCAACCYCACCTGTACTTCAGG	Peeling skin syndrome, acral type
397515519	NM_000207.2(INS):c.*59A>G	GGGCTTATTCCATCTCTCTCGG	Permanent neonatal diabetes mellitus
397515523	NM_000370.3(TTPA):c.191A>G (p.Asp64Gly)	CAGGYCCAGATCGAAATCCCGGG, CCAGGYCCAGATCGAAATCCCGG	Ataxia with vitamin E deficiency
397515891	NM_000256.3(MYBPC3):c.1224- 2A>G	TACTTGCGYGTAGAACAGAAGGGG	Familial hypertrophic cardiomyopathy 4, Cardiomyopathy
397516082	NM_000256.3(MYBPC3):c.927- 2A>G	GTCCCYGTGTCCCGCAGTCTAGG	Familial hypertrophic cardiomyopathy 4, Cardiomyopathy
397516138	NM_000257.3(MYH7):c.2206A>G (p.Ile736Val)	TATCAAYGAACTGTCCCTCAGGG, CTATCAAYGAACTGTCCCTCAGG	Familial hypertrophic cardiomyopathy 1, Cardiomyopathy, not specified
1154510	NM_002150.2(HPD):c.97G>A (p.Ala33Thr)	ATGACGYGGCCTGAATCACAGGG, AATGACGYGGCCTGAATCACAGG	4-Alpha-hydroxyphenylpyruvate hydroxylase deficiency
397516330	NM_000260.3(MYO7A):c.6439- 2A>G	ATATCCYGGGGAGCAGAAAAGGG, GATATCCYGGGGAGCAGAAAAGG	Usher syndrome, type 1
72556271	NM_000531.5(OTC):c.482A>G (p.Asn161Ser)	CAGCCCAYTGATAATTGGGATGG	not provided
606231260	NM_023073.3(C5orf42):c.3290- 2A>G	ATCYATCAAATACAAAATTGG	Orofaciodigital syndrome 6
587777521	NM_004817.3(TJP2):c.1992- 2A>G	CAGCTCYGAGAAGAAACCACGGG, TCAGCTCYGAGAAGAAACCACGG	Progressive familial intrahepatic cholestasis 4
730880846	NM_000257.3(MYH7):c.617A>G (p.Lys206Arg)	CTTCYTGCTGCGGTCCCAATGG	Cardiomyopathy
397517978	NM_206933.2(USH2A):c.12067- 2A>G	TTCCCYGTAAGAAAATTAACAGG	Usher syndrome, type 2A, Retinitis pigmentosa 39
606231409	NM_000216.2(ANOS1):c.1A>G (p.Met1Val)	GCACCAYGCTGCGGGTCGAGGG, GGCACCAYGCTGCGGGTCGAGG	Kallmann syndrome 1
80356546	NM_003334.3(UBA1):c.1639A>G (p.Ser547Gly)	TGGCYGTACCCGGATATGTGG	Arthrogryposis multiplex congenita, distal, X- linked
80356584	NM_194248.2(OTOF):c.766- 2A>G	GACCYGCAGGCAGGAGAAGGGGG, TGACCYGCAGGCAGGAGAAGGGG, CTGACCYGCAGGCAGGAGAAGGG, GCTGACCYGCAGGCAGGAGAAGG	Deafness, autosomal recessive 9
730880930	NM_000257.3(MYH7):c.1615A>G (p.Met539Val)	GGAACAYGCACTCTCTCCAGG	Cardiomyopathy
118203947	NM_013319.2(UBIAD1):c.355A>G (p.Arg119Gly)	TCCYGTCACTCTTTTTGTGG	Schneider crystalline corneal dystrophy
60171927	NM_000526.4(KRT14):c.368A>G (p.Asn123Ser)	GCGGTCAYTGAGTTCTGCATGG	Epidermolysis bullosa herpetiformis, Dowling- Meara
199422248	NM_001363.4(DKCI):c.941A>G (p.Lys314Arg)	AATCYTGGCCCCATAGCAGATGG	Dyskeratosis congenita X-linked
72558467	NM_000531.5(OTC):c.929A>G (p.Glu310Gly)	TCCACTYCTTCTGGCTTTCTGGG, ATCCACTYCTTCTGGCTTTCTGG	not provided
72558478	NM_000531.5(OTC):c.988A>G (p.Arg330Gly)	ACTTTCYGTITTTCTGCCTCTGGG, CACTTTCYGTITTTCTGCCTCTGG	not provided
118204455	NM_000505.3(F12):c.158A>G (p.Tyr53Cys)	GGTGGYACTGGAAGGGGAAGTGG	
80357477	NM_007294.3(BRCA1):c.5453A>G (p.Asp1818Gly)	TTGYCCTCTGTCCAGGCATCTGG	Familial cancer of breast, Breast-ovarian cancer, familial 1
121907908	NM_024426.4(WT1):c.1021A>G (p.Ser341Gly)	CGCYCTGTACCCTGTGCTGTGG	Mesothelioma
121907926	NM_000280.4(PAX6):c.1171A>G (p.Thr391Ala)	GTGGYGCCCGAGGTGCCCATTTGG	Optic nerve aplasia, bilateral
121908023	NM_024740.2(ALG9):c.860A>G (p.Tyr287Cys)	TTAYACAAAACAATGTTGAGTGG	Congenital disorder of glycosylation type 1L
121908148	NM_001243133.1(NLRP3):c.1880A>G (p.Glu627Gly)	ACAATYCCAGCTGGCTGGGCTGG	Familial cold urticaria
121908166	NM_006492.2(ALX3):c.608A>G (p.Asn203Ser)	CGGYTCTGGAACACAGCTGGGG, GCGGYTCTGGAACACAGACCTGGG, TGCGGYTCTGGAACACAGACCTGG	Frontonasal dysplasia 1

121908184	NM_020451.2(SEPN1):c.1A>G (p.Met1Val)	CCCAYGGCTGCGGCTGGCGGCGG, CCGCCCAAYGGCTGCGGCTGGCGG	Eichsfeld type congenital muscular dystrophy
121908258	NM_130468.3(CHST14):c.878A>G (p.Tyr293Cys)	AAGTCAYAGTGCACGGCACAAGG	Ehlers-Danlos syndrome, musculocontractural type
121908383	NM_001128425.1(MUTYH):c.1241A>G (p.Gln414Arg)	AAGCYGCTCTGAGGGCTCCAGG	Neoplasm of stomach
121908580	NM_004328.4(BCSL1):c.148A>G (p.Thr50Ala)	GTGYGATCATGTAATGGCGCCGG	Mitochondrial complex III deficiency
121908584	NM_016417.2(GLRX5):c.294A>G (p.Gln98=)	CCTGACCYGTGCGGAGCTCCGGG	Anemia, sideroblastic, pyridoxine-refractory, autosomal recessive
121908635	NM_022817.2(PER2):c.1984A>G (p.Ser662Gly)	GCCACACYCTCTGCCTTGCCCGG	Advanced sleep phase syndrome, familial
121908655	NM_003839.3(TNFRSF11A):c.508A>G (p.Arg170Gly)	GGGTCYGCATTTGTCCGTGGAGG	Osteopetrosis autosomal recessive 7
29001653	NM_000539.3(RHO):c.886A>G (p.Lys296Glu)	CGCTCTYGGCAAAGAACGCTGGG, GCGCTCTYGGCAAAGAACGCTGG	Retinitis pigmentosa 4
56307355	NM_006502.2(POLH):c.1603A>G (p.Lys535Glu)	AGACTTTYCTGCTTAAAGAAGGG	Xeroderma pigmentosum, variant type
121908919	NM_002977.3(SCN9A):c.1964A>G (p.Lys655Arg)	CCTTTTCYGTGTATTGATTGG	Generalized epilepsy with febrile seizures plus, type 7, not specified
121908939	NM_006892.3(DNMT3B):c.2450A>G (p.Asp817Gly)	GACACGYCTGTGTAGTGACAGG	Centromeric instability of chromosomes 1,9 and 16 and immunodeficiency
121909088	NM_001005360.2(DNM2):c.1684A>G (p.Lys562Glu)	ACTYCTTCTTTCTCTGAGGG, TACTYCTTCTTTCTCTGAGG	Charcot-Marie-Tooth disease, dominant intermediate b, with neutropenia
120074112	NM_000483.4(APOC2):c.1A>G (p.Met1Val)	GCCAYAGTGTCCAGAGACCTGG	Apolipoprotein C2 deficiency
121909239	NM_000314.6(PTEN):c.755A>G (p.Asp252Gly)	ATAYCACCACACACAGGTAACGG	Macrocephaly/autism syndrome
121909251	NM_198217.2(ING1):c.515A>G (p.Asn172Ser)	TGGYTGACAGACAGTACGTGGG, CTGGYTGACAGACAGTACGTGG	Squamous cell carcinoma of the head and neck
121909396	NM_001174089.1(SLC4A11):c.2518A>G (p.Met840Val)	GATCAYCTTATGTAGGGCAGGG, AGATCAYCTTATGTAGGGCAGG	Corneal dystrophy and perceptive deafness
121909533	NM_000034.3(ALDOA):c.386A>G (p.Asp129Gly)	CCAYCCAACCCTAAGAGAAGAGG	HNSHA due to aldolase A deficiency
128627255	NM_004006.2(DMD):c.835A>G (p.Thr279Ala)	TGACCGYGATCTGCAGAGAAGGG, CTGACCGYGATCTGCAGAGAAGG	Dilated cardiomyopathy 3B
116929575	NM_001085.4(SERPINA3):c.1240A>G (p.Met414Val)	GCTCAYGAAGAAGATGTTCTGGG, TGCTCAYGAAGAAGATGTTCTGG	
61748392	NM_004992.3(MECP2):c.410A>G (p.Glu137Gly)	CAACYCCACTTTAGAGCGAAAGG	Mental retardation, X-linked, syndromic 13
61748906	NM_001005741.2(GBA):c.667T>C (p.Trp223Arg)	CCCACTYGGCTCAAGACCAATGG	Gaucher disease, type 1
199473024	NM_000238.3(KCNH2):c.3118A>G (p.Ser1040Gly)	CTGCYCTCCACGTCGCCCCGGG, CCTGCYCTCCACGTCGCCCCGGG, GCCTGCYCTCCACGTCGCCCCGG	Sudden infant death syndrome
794728365	NM_000238.3(KCNH2):c.1129-2A>G	GGACCYGCACCCGGGAAGGCCGG	Cardiac arrhythmia
72556293	NM_000531.1(OTC):c.548A>G (p.Tyr183Cys)	AGAGCTAYAGTGTCTCTAAAAGG	not provided
111033244	NM_000441.1(SLC26A4):c.1151A>G (p.Glu384Gly)	TGAATYCCTAAGGAAGAGACTGG	Pendred syndrome, Enlarged vestibular aqueduct syndrome
111033415	NM_000260.3(MYO7A):c.1344-2A>G	AGCYGCAGGGGCACAGGGATGGG, AAGCYGCAGGGGCACAGGGATGG	Usher syndrome, type 1
121912439	NM_000454.4(SOD1):c.302A>G (p.Glu101Gly)	AGAATCTYCAATAGACACATCGG	Amotrophic lateral sclerosis type 1
111033567	NM_002769.4(PRSS1):c.68A>G (p.Lys23Arg)	ATCYTGTATCATCATCAAAGGG, GATCYTGTATCATCATCAAAGG	Hereditary pancreatitis
121912565	NM_000901.4(NR3C2):c.2327A>G (p.Gln776Arg)	TCATCYGTTTGCTGCTAAGCGG	Pseudohypaldosteronism type 1 autosomal dominant
121912574	NM_000901.4(NR3C2):c.2915A>G (p.Glu972Gly)	CCGACYCCACCTTGGGCAGCTGG	Pseudohypaldosteronism type 1 autosomal dominant
121912589	NM_001173464.1(KIF21A):c.2839A>G (p.Met947Val)	ATTCAATCTGCCTCCATGTTGG	Fibrosis of extraocular muscles, congenital, 1
111033661	NM_000155.3(GALT):c.253-2A>G	ATTCACCYACCACAAGGATAGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
111033669	NM_000155.3(GALT):c.290A>G (p.Asn97Ser)	GAAGTCGYTGTCAAACAGGAAGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase

111033682	NM_000155.3(GALT):c.379A>G (p.Lys127Glu)	TGACCTYACTGGGTGGTGACGGG, ATGACCTYACTGGGTGGTGACGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
111033786	NM_000155.3(GALT):c.950A>G (p.Gln317Arg)	CAGCYGCCAATGGTTCAGTTGG	Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase
121912765	NM_001202.3(BMP4):c.278A>G (p.Glu93Gly)	CCTCCYCCCCAGACTGAAGCCGG	Microphthalmia syndromic 6
121912856	NM_000094.3(COL7A1):c.425A>G (p.Lys142Arg)	CACCYTGGGGACACCAGGTCGGG, TCACCYTGGGGACACCAGGTCGG	Epidermolysis bullosa dystrophica inversa, autosomal recessive
199474715	NM_152263.3(TPM3):c.505A>G (p.Lys169Glu)	CCAACTYACGAGCCACCTACAGG	Congenital myopathy with fiber type disproportion
199474718	NM_152263.3(TPM3):c.733A>G (p.Arg245Gly)	ATCYCTCAGCAAACCTCAGCACGG	Congenital myopathy with fiber type disproportion
121912895	NM_001844.4(COL2A1):c.2974A>G (p.Arg992Gly)	CCTCYCTCACCACGTTGCCAGG	Spondyloepimetaphyseal dysplasia Strudwick type
121913074	NM_000129.3(F13A1):c.851A>G (p.Tyr284Cys)	ATAGGCAYAGATATTGTCCAGG	Factor xiii, a subunit, deficiency of
121913145	NM_000208.2(INSR):c.707A>G (p.His236Arg)	GCTGYGCAACAGAGGCCCTTCGG	Leprechaunism syndrome
312262745	NM_025137.3(SPG11):c.2608A>G (p.Ile870Val)	ACTTAYCCTGGGGAGAAGGATGG	Spastic paraplegia 11, autosomal recessive
121913682	NM_000222.2(KIT):c.2459A>G (p.Asp820Gly)	AGAAACATTCTGTATGTCTCTGG	Mast cell disease, systemic
587776757	NM_000151.3(G6PC):c.230+4A> G	GTTCYTACCACCTAAAGACGAGG	Glycogen storage disease type 1A
61752063	NM_000330.3(RS1):c.286T>C (p.Trp96Arg)	TTCTTCGYGGACTGCAACAAGG	Juvenile retinoschisis
367543065	NM_024549.5(TCTN1):c.221- 2A>G	AGCAACYGCAGAAAAAGAGGGG, CAGCAACYGCAGAAAAAGAGGG G	Joubert syndrome 13
5030773	NM_000894.2(LHB):c.221A>G (p.Gln74Arg)	CCACCYAGGCGAGGGCGGCAGG	Isolated lutropin deficiency
199476092	NM_000264.3(PTCH1):c.2479A>G (p.Ser827Gly)	CGTTACYGAAACTCCTGTGTAGG	Gorlin syndrome, Holoprosencephaly 7, not specified
398123158	NM_000117.2(EMD):c.450- 2A>G	CGTCCCCYAGGCAAAAAGAGGGG	not provided
199476103	RMRP:n.71A>G	ACTTYCCCTAGGCGGAAAGGGG, GACTTYCCCTAGGCGGAAAGGG, GGACTTYCCCTAGGCGGAAAGG CTCYCTGCCACGTAATACAGGGG, ACTCYCTGCCACGTAATACAGGG, AACTCYCTGCCACGTAATACAGG GGGTCGYAGCGAACTGAGAAGGG, TGGGTCGYAGCGAACTGAGAAGG	Metaphyseal chondrodysplasia, McKusick type, Metaphyseal dysplasia without hypotrichosis
5030856	NM_000277.1(PAH):c.1169A>G (p.Gln390Gly)	CTCYCTGCCACGTAATACAGGGG, ACTCYCTGCCACGTAATACAGGG, AACTCYCTGCCACGTAATACAGG	Phenylketonuria, Hyperphenylalaninemia, non- pku
5030860	NM_000277.1(PAH):c.1241A>G (p.Tyr414Cys)	GGGTCGYAGCGAACTGAGAAGGG, TGGGTCGYAGCGAACTGAGAAGG	Phenylketonuria, Hyperphenylalaninemia, non- pku
587777055	NM_020988.2(GNAO1):c.521A>G (p.Asp174Gly)	GGATGYCCTGCTCGGTGGGCTGG	Early infantile epileptic encephalopathy 17
587777223	NM_024301.4(FKRP):c.1A>G (p.Met1Val)	CCGCAYGGGGCCGAAGTCTGGGG, GCCGCAYGGGGCCGAAGTCTGGG, AGCCGCAYGGGGCCGAAGTCTGG	Congenital muscular dystrophy- dystroglycanopathy with brain and eye anomalies type A5
587777479	NM_003108.3(SOX11):c.347A>G (p.Tyr116Cys)	GTACTTGYAGTCGGGTAGTCGG	Mental retardation, autosomal dominant 27
587777496	NM_020435.3(GJC2):c.-170A>G	TTGYTCCCCCTCGGCCTCAGGG, ATTGYTCCCCCTCGGCCTCAGG	Leukodystrophy, hypomyelinating, 2
587777507	NM_022552.4(DNMT3A):c.1943T>C (p.Leu648Pro)	CTCCYGGTGTGAAGGACTTGGG, GCTCCYGGTGTGAAGGACTTGG	Tatton-Brown-rahman syndrome
587777557	NM_018400.3(SCN3B):c.482T>C (p.Met161Thr)	AATCAYGATGTACATCCTTCTGG	Atrial fibrillation, familial, 16
587777569	NM_001030001.2(RPS29):c.149T>C (p.Ile50Thr)	GATAYCGGTTTCATTAAGGTAGG	Diamond-Blackfan anemia 13
587777657	NM_153334.6(SCARF2):c.190T>C (p.Cys64Arg)	CCACGYGCTGCGCTGGCTGAAGG	Marden Walker like syndrome
587777689	NM_005726.5(TSFM):c.57+4A> G	ACTTCYACCCGGGTAGCTCCCGG	Combined oxidative phosphorylation deficiency 3
796052005	NM_000255.3(MUT):c.329A>G (p.Tyr110Cys)	GCAYACTGGCGGATGGTCCAGGG, AGCAYACTGGCGGATGGTCCAGG	not provided
587777809	NM_144596.3(TTC8):c.115- 2A>G	GITCCYGGAAAGCATTAAAGAAGG	Retinitis pigmentosa 51
587777878	NM_000166.5(GJB1):c.580A>G (p.Met194Val)	TAGCAYGAAGACGGTGAAGACGG	X-linked hereditary motor and sensory neuropathy

74315420	NM_001029871.3(RSPO4):c.194A>G (p.Gln65Arg)	CGTACYGGCGGATGCCTTCCCGG	Anonychia
180177219	NM_000030.2(AGXT):c.424-2A>G (p.Gly_142Gln145del)	AGGCCCYGAGGAAGCAGGGACGG	Primary hyperoxaluria, type I
367610201	NM_002693.2(POLG):c.1808T>C (p.Met603Thr)	CTCAYGGCACTTACCTGGGATGG	not provided
180177319	NM_012203.1(GRHPR):c.84-2A>G	TCACAGCYGCGGGAAAGGGAGG	Primary hyperoxaluria, type II
796052068	NM_000030.2(AGXT):c.777-2A>G	GGTACCYGGAAGACACGAGGGGG, TGGTACCYGGAAGACACGAGGGGG	Primary hyperoxaluria, type I
61754010	NM_000552.3(VWF):c.1583A>G (p.Asn528Ser)	TGCCAYTGTAATTCACACACAGG	von Willebrand disease, type 2a
587778866	NM_000321.2(RB1):c.1927A>G (p.Lys643Glu)	ATTYCAATGGCTTCTGGGTCTGG	Retinoblastoma
74435397	NM_006331.7(EMG1):c.257A>G (p.Asp86Gly)	ATAYCTGGCCGCGCTTCCCCAGG	Bowen-Conradi syndrome
796052527	NM_000156.5(GAMT):c.1A>G (p.Met1Val)	CGCTCAYGCTGCAGGCTGGACGG	not provided
796052637	NM_172107.2(KCNQ2):c.848A>G (p.Lys283Arg)	GTACYTGCCCCGTAGCCAATGG	not provided
724159963	NM_032228.5(FAR1):c.1094A>G (p.Asp365Gly)	GATAYCATAACAGGAATGCTGGGG, AGATAYCATAACAGGAATGCTGGG, TAGATAYCATAACAGGAATGCTGG	Peroxisomal fatty acyl-coa reductase 1 disorder
587779722	NM_000090.3(COL3A1):c.1762-2A>G (p.Gly588_Gln605del)	CACCCYAAAGAAGAAGTGCTCGG	Ehlers-Danlos syndrome, type 4
118192102	m.8296A>G	TTTACAGYGGGCTCTAGAGGGGG	Diabetes-deafness syndrome maternally transmitted
727502787	NM_001077494.3(NFKB2):c.2594A>G (p.Asp865Gly)	CTGYCTTCTCACCTCTGTCTGG	Common variable immunodeficiency 10
727503036	NM_000117.2(EMD):c.266-2A>G	AGCCYTGGAAGGGGGCAGCGG	Emery-Dreifuss muscular dystrophy 1, X-linked
690016544	NM_005861.3(STUB1):c.194A>G (p.Asn65Ser)	GGCCCGYTGGTGTAATACACGG	Spinocerebellar ataxia, autosomal recessive 16
690016554	NM_005211.3(CSF1R):c.2655-2A>G	GTATCYGGGAGATAGGACAGAGG	Hereditary diffuse leukoencephalopathy with spheroids
118192185	NM_172107.2(KCNQ2):c.1A>G (p.Met1Val)	GCACCAYGGTGCCTGGCGGGAGG	Benign familial neonatal seizures 1
121917869	NM_012064.3(MIP):c.401A>G (p.Glu134Gly)	AGATCYCCACTGTGGTTGCCTGG	Cataract 15, multiple types
121918014	NM_000478.4(ALPL):c.1250A>G (p.Asn417Ser)	AGGCCAYTGCCATACAGGATGG	Infantile hypophosphatasia
121918036	NM_000174.4(GP9):c.110A>G (p.Asp37Gly)	GCAGYCCACCCACAGCCCATGG	Bernard-Soulier syndrome type C
121918089	NM_000371.3(TTR):c.379A>G (p.Ile127Val)	CGGCAAYGGTGTAGCGCGGGGG, GCGGCAAYGGTGTAGCGCGGGGG	Amyloidogenic transthyretin amyloidosis
121918121	NM_000823.3(GHRHR):c.985A>G (p.Lys329Glu)	CGACTYGGAGAGACGCCTGCAGG	Isolated growth hormone deficiency type 1B
121918333	NM_015335.4(MED13L):c.6068A>G (p.Asp2023Gly)	ATATCAYCTAGAGGGAAGGGGG, CATATCAYCTAGAGGGAAGGGGG	Transposition of great arteries
121918605	NM_001035.2(RYR2):c.12602A>G (p.Gln4201Arg)	CGCCAGCYGCATTTCAAAGATGG	Catecholaminergic polymorphic ventricular tachycardia
587781262	NM_002764.3(PRPS1):c.343A>G (p.Met115Val)	TAGCAYATTTGCAACAAGCTTGG	Charcot-Marie-Tooth disease, X-linked recessive, type 5, Deafness, high-frequency sensorineural, X-linked
121918608	NM_001161766.1(AHCY):c.344A>G (p.Tyr115Cys)	GCGGGYACTTGGTGTGGATGAGG	Hypermethioninemia with s-adenosylhomocysteine hydrolase deficiency
121918613	NM_000702.3(ATPIA2):c.1033A>G (p.Thr345Ala)	CTGYCAGGGTCAGGCACACCTGG	Familial hemiplegic migraine type 2
587781339	NM_000535.5(PMS2):c.904-2A>G	GCAGACCYGCACAAAATACAAGG	Hereditary cancer-predisposing syndrome
121918691	NM_001128177.1(THRB):c.1324A>G (p.Met442Val)	CTTCAYGTGCAGGAAGCGGCTGG	Thyroid hormone resistance, generalized, autosomal dominant
121918692	NM_001128177.1(THRB):c.1327A>G (p.Lys443Glu)	CCACCTCATGTGCAGGAAGCGG	Thyroid hormone resistance, generalized, autosomal dominant
727504333	NM_000256.3(MYBPC3):c.2906-2A>G	CCGTTTCYGTGGGTATAGAGTGGG, GCCGTTTCYGTGGGTATAGAGTGG	Familial hypertrophic cardiomyopathy 4
730880805	NM_006204.3(PDE6C):c.1483-2A>G	CTTTCYGTGAAATAAGGATGGG, TCTTTCYGTGAAATAAGGATGG	Achromatopsia 5

281860296	NM_000551.3(VHL):c.586A>T (p.Lys196Ter)	GGTCTTYCTGCACATTGGGTGG	Von Hippel-Lindau syndrome
730880444	NM_000169.2(GLA):c.370- 2A>G	GTGAACCYGAAATGAGAGGGAGG	not provided
756328339	NM_000256.3(MYBPC3):c.1227- 2A>G	GTACCYGGGTGGGGCCGCAGGG, TGTACCYGGGTGGGGGCCGCAGG	Familial hypertrophic cardiomyopathy 4, Cardiomyopathy
267606643	NM_013411.4(AK2):c.494A>G (p.Asp165Gly)	TCAYCTTTCATGGGCTCTTTTGG	Reticular dysgenesis
267606705	NM_005188.3(CBL):c.1144A>G (p.Lys382Glu)	TATTTYACATAGTTGGAATGTGG	Noonan syndrome-like disorder with or without juvenile myelomonocytic leukemia
62642934	NM_000277.1(PAH):c.916A>G (p.Ile306Val)	GGCCAAYTTCCTGTAATTGGGGG, AGGCCAAYTTCCTGTAATTGGGGG	Phenylketonuria, Hyperphenylalaninemia, non- pku
267606782	NM_000117.2(EMD):c.1A>G (p.Met1Val)	TCCAYGGCGGGTGC GGCTCAGG	Emery-Dreifuss muscular dystrophy, X-linked
267606820	NM_014053.3(FLVCR1):c.361A>G (p.Asn121Asp)	AGGCGTYGACCAGCGAGTACAGG	Posterior column ataxia with retinitis pigmentosa

[00318] In some embodiments, any of the base editors provided herein may be used to treat a disease or disorder. For example, any base editors provided herein may be used to correct one or more mutations associated with any of the diseases or disorders provided herein. Exemplary diseases or disorders that may be treated include, without limitation, 3-Methylglutaconic aciduria type 2, 46,XY gonadal dysgenesis, 4-Alpha-hydroxyphenylpyruvate hydroxylase deficiency, 6-pyruvoyl-tetrahydropterin synthase deficiency, achromatopsia, Acid-labile subunit deficiency, Acrodysostosis, acroerythrokeratoderma, ACTH resistance, ACTH-independent macronodular adrenal hyperplasia, Activated PI3K-delta syndrome, Acute intermittent porphyria, Acute myeloid leukemia, Adams-Oliver syndrome 1/5/6, Adenylosuccinate lyase deficiency, Adrenoleukodystrophy, Adult neuronal ceroid lipofuscinosis, Adult onset ataxia with oculomotor apraxia, Advanced sleep phase syndrome, Age-related macular degeneration, Alagille syndrome, Alexander disease, Allan-Herndon-Dudley syndrome, Alport syndrome, X-linked recessive, Alternating hemiplegia of childhood, Alveolar capillary dysplasia with misalignment of pulmonary veins, Amelogenesis imperfecta, Amyloidogenic transthyretin amyloidosis, Amyotrophic lateral sclerosis, Anemia (nonspherocytic hemolytic, due to G6PD deficiency), Anemia (sideroblastic, pyridoxine-refractory, autosomal recessive), Anonychia, Antithrombin III deficiency, Aortic aneurysm, Aplastic anemia, Apolipoprotein C2 deficiency, Apparent mineralocorticoid excess, Aromatase deficiency, Arrhythmogenic right ventricular cardiomyopathy, Familial hypertrophic cardiomyopathy, Hypertrophic cardiomyopathy, Arthrogryposis multiplex congenital, Aspartylglycosaminuria, Asphyxiating thoracic dystrophy, Ataxia with vitamin E deficiency, Ataxia (spastic), Atrial fibrillation, Atrial septal defect, atypical hemolytic-uremic syndrome, autosomal dominant CD11C+/CD1C+ dendritic cell deficiency, Autosomal dominant progressive external ophthalmoplegia with mitochondrial DNA deletions, Baraitser-Winter syndrome, Bartter syndrome, Basa ganglia calcification, Beckwith-Wiedemann syndrome, Benign familial neonatal seizures, Benign scapuloperoneal muscular dystrophy, Bernard Soulier syndrome, Beta thalassemia intermedia, Beta-D-mannosidosis, Bietti crystalline corneoretinal dystrophy, Bile acid malabsorption, Biotinidase deficiency, Borjeson-Forssman-Lehmann syndrome, Boucher Neuhauser syndrome, Bowen-Conradi syndrome, Brachydactyly, Brown-Vialetto-Van laere syndrome, Brugada syndrome, Cardiac arrhythmia, Cardiofaciocutaneous syndrome, Cardiomyopathy, Carnevale syndrome, Carnitine palmitoyltransferase II deficiency, Carpenter syndrome, Cataract, Catecholaminergic polymorphic

ventricular tachycardia, Central core disease, Centromeric instability of chromosomes 1,9 and 16 and immunodeficiency, Cerebral autosomal dominant arteriopathy, Cerebro-oculo-facio-skeletal syndrome, Ceroid lipofuscinosis, Charcot-Marie-Tooth disease, Cholestanol storage disease, Chondrocalcinosis, Chondrodysplasia, Chronic progressive multiple sclerosis, Coenzyme Q10 deficiency, Cohen syndrome, Combined deficiency of factor V and factor VIII, Combined immunodeficiency, Combined oxidative phosphorylation deficiency, Combined partial 17-alpha-hydroxylase/17,20-lyase deficiency, Complement factor d deficiency, Complete combined 17-alpha-hydroxylase/17,20-lyase deficiency, Cone-rod dystrophy, Congenital contractural arachnodactyly, Congenital disorder of glycosylation, Congenital lipomatous overgrowth, Neoplasm of ovary, PIK3CA Related Overgrowth Spectrum, Congenital long QT syndrome, Congenital muscular dystrophy, Congenital muscular hypertrophy-cerebral syndrome, Congenital myasthenic syndrome, Congenital myopathy with fiber type disproportion, Eichsfeld type congenital muscular dystrophy, Congenital stationary night blindness, Corneal dystrophy, Cornelia de Lange syndrome, Craniometaphyseal dysplasia, Crigler Najjar syndrome, Crouzon syndrome, Cutis laxa with osteodystrophy, Cyanosis, Cystic fibrosis, Cystinosis, Cytochrome-c oxidase deficiency, Mitochondrial complex I deficiency, D-2-hydroxyglutaric aciduria, Danon disease, Deafness with labyrinthine aplasia microtia and microdontia (LAMM), Deafness, Deficiency of acetyl-CoA acetyltransferase, Deficiency of ferroxidase, Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase, Dejerine-Sottas disease, Desbuquois syndrome, DFNA, Diabetes mellitus type 2, Diabetes-deafness syndrome, Diamond-Blackfan anemia, Diastrophic dysplasia, Dihydropteridine reductase deficiency, Dihydropyrimidinase deficiency, Dilated cardiomyopathy, Disseminated atypical mycobacterial infection, Distal arthrogryposis, Distal hereditary motor neuronopathy, Donnai Barrow syndrome, Duchenne muscular dystrophy, Becker muscular dystrophy, Dyschromatosis universalis hereditaria, Dyskeratosis congenital, Dystonia, Early infantile epileptic encephalopathy, Ehlers-Danlos syndrome, Eichsfeld type congenital muscular dystrophy, Emery-Dreifuss muscular dystrophy, Enamel-renal syndrome, Epidermolysis bullosa dystrophica inversa, Epidermolysis bullosa herpetiformis, Epilepsy, Episodic ataxia, Erythrokeratoderma variabilis, Erythropoietic protoporphyria, Exercise intolerance, Exudative vitreoretinopathy, Fabry disease, Factor V deficiency, Factor VII deficiency, Factor xiii deficiency, Familial adenomatous polyposis, breast cancer, ovarian cancer, cold urticarial, chronic infantile neurological, cutaneous and articular syndrome, hemiplegic migraine, hypercholesterolemia, hypertrophic cardiomyopathy,

hypoalphalipoproteinemia, hypokalemia-hypomagnesemia, juvenile gout, hyperlipoproteinemia, visceral amyloidosis, hypophosphatemic vitamin D refractory rickets, FG syndrome, Fibrosis of extraocular muscles, Finnish congenital nephrotic syndrome, focal epilepsy, Focal segmental glomerulosclerosis, Frontonasal dysplasia, Frontotemporal dementia, Fructose-biphosphatase deficiency, Gamstorp-Wohlfart syndrome, Ganglioside sialidase deficiency, GATA-1-related thrombocytopenia, Gaucher disease, Giant axonal neuropathy, Glanzmann thrombasthenia, Glomerulocystic kidney disease, Glomerulopathy, Glucocorticoid resistance, Glucose-6-phosphate transport defect, Glutaric aciduria, Glycogen storage disease, Gorlin syndrome, Holoprosencephaly, GRACILE syndrome, Haemorrhagic telangiectasia, Hemochromatosis, Hemoglobin H disease, Hemolytic anemia, Hemophagocytic lymphohistiocytosis, Carcinoma of colon, Myhre syndrome, leukoencephalopathy, Hereditary factor IX deficiency disease, Hereditary factor VIII deficiency disease, Hereditary factor XI deficiency disease, Hereditary fructosuria, Hereditary Nonpolyposis Colorectal Neoplasm, Hereditary pancreatitis, Hereditary pyropoikilocytosis, Elliptocytosis, Heterotaxy, Heterotopia, Histiocytic medullary reticulosis, Histiocytosis-lymphadenopathy plus syndrome, HNSHA due to aldolase A deficiency, Holocarboxylase synthetase deficiency, Homocysteinemia, Howel-Evans syndrome, Hydatidiform mole, Hypercalciuric hypercalcemia, Hyperimmunoglobulin D, Mevalonic aciduria, Hyperinsulinemic hypoglycemia, Hyperkalemic Periodic Paralysis, Paramyotonia congenita of von Eulenburg, Hyperlipoproteinemia, Hypermanganesemia, Hypermethioninemia, Hyperphosphatasemia, Hypertension, hypomagnesemia, Hypobetalipoproteinemia, Hypocalcemia, Hypogonadotropic hypogonadism, Hypogonadotropic hypogonadism, Hypohidrotic ectodermal dysplasia, Hyper-IgM immunodeficiency, Hypohidrotic X-linked ectodermal dysplasia, Hypomagnesemia, Hypoparathyroidism, Idiopathic fibrosing alveolitis, Immunodeficiency, Immunoglobulin A deficiency, Infantile hypophosphatasia, Infantile Parkinsonism-dystonia, Insulin-dependent diabetes mellitus, Intermediate maple syrup urine disease, Ischiopatellar dysplasia, Islet cell hyperplasia, Isolated growth hormone deficiency, Isolated lutropin deficiency, Isovaleric acidemia, Joubert syndrome, Juvenile polyposis syndrome, Juvenile retinoschisis, Kallmann syndrome, Kartagener syndrome, Kugelberg-Welander disease, Lattice corneal dystrophy, Leber congenital amaurosis, Leber optic atrophy, Left ventricular noncompaction, Leigh disease, Mitochondrial complex I deficiency, Leprechaunism syndrome, Arthrogryposis, Anterior horn cell disease, Leukocyte adhesion deficiency, Leukodystrophy, Leukoencephalopathy, Ovarioleukodystrophy, L-ferritin deficiency, Li-Fraumeni syndrome,

Limb-girdle muscular dystrophy- dystroglycanopathy, Loeys-Dietz syndrome, Long QT syndrome, Macrocephaly/autism syndrome, Macular corneal dystrophy, Macular dystrophy, Malignant hyperthermia susceptibility, Malignant tumor of prostate, Maple syrup urine disease, Marden Walker like syndrome, Marfan syndrome, Marie Unna hereditary hypotrichosis, Mast cell disease, Meconium ileus, Medium-chain acyl-coenzyme A dehydrogenase deficiency, Melnick-Fraser syndrome, Mental retardation, Merosin deficient congenital muscular dystrophy, Mesothelioma, Metachromatic leukodystrophy, Metaphyseal chondrodysplasia, Methemoglobinemia, methylmalonic aciduria, homocystinuria, Microcephaly, chorioretinopathy, lymphedema, Microphthalmia, Mild non-PKU hyperphenylalanemia, Mitchell-Riley syndrome, mitochondrial 3-hydroxy-3-methylglutaryl-CoA synthase deficiency, Mitochondrial complex I deficiency, Mitochondrial complex III deficiency, Mitochondrial myopathy, Mucopolidosis III, Mucopolysaccharidosis, Multiple sulfatase deficiency, Myasthenic syndrome, Mycobacterium tuberculosis, Myeloperoxidase deficiency, Myhre syndrome, Myoclonic epilepsy, Myofibrillar myopathy, Myoglobinuria, Myopathy, Myopia, Myotonia congenital, Navajo neurohepatopathy, Nemaline myopathy, Neoplasm of stomach, Nephrogenic diabetes insipidus, Nephronophthisis, Nephrotic syndrome, Neurofibromatosis, Neutral lipid storage disease, Niemann-Pick disease, Non-ketotic hyperglycinemia, Noonan syndrome, Noonan syndrome-like disorder, Norum disease, Macular degeneration, N-terminal acetyltransferase deficiency, Oculocutaneous albinism, Oculodentodigital dysplasia, Ohdo syndrome, Optic nerve aplasia, Ornithine carbamoyltransferase deficiency, Orofaciodigital syndrome, Osteogenesis imperfecta, Osteopetrosis, Ovarian dysgenesis, Pachyonychia, Palmoplantar keratoderma, nonepidermolytic, Papillon-Lefevre syndrome, Haim-Munk syndrome, Periodontitis, Peeling skin syndrome, Pendred syndrome, Peroxisomal fatty acyl-coa reductase 1 disorder, Peroxisome biogenesis disorder, Pfeiffer syndrome, Phenylketonuria, Phenylketonuria, Hyperphenylalaninemia, non-PKU, Pituitary hormone deficiency, Pityriasis rubra pilaris, Polyarteritis nodosa, Polycystic kidney disease, Polycystic lipomembranous osteodysplasia, Polymicrogyria, Pontocerebellar hypoplasia, Porokeratosis, Posterior column ataxia, Primary erythromelalgia, hyperoxaluria, Progressive familial intrahepatic cholestasis, Progressive pseudorheumatoid dysplasia, Propionic acidemia, Pseudohermaphroditism, Pseudohypoaldosteronism, Pseudoxanthoma elasticum-like disorder, Purine-nucleoside phosphorylase deficiency, Pyridoxal 5-phosphate-dependent epilepsy, Renal dysplasia, retinal pigmentary dystrophy, cerebellar ataxia, skeletal dysplasia, Reticular dysgenesis, Retinitis pigmentosa, Usher syndrome, Retinoblastoma, Retinopathy, RRM2B-related

mitochondrial disease, Rubinstein-Taybi syndrome, Schnyder crystalline corneal dystrophy, Sebaceous tumor, Severe congenital neutropenia, Severe myoclonic epilepsy in infancy, Severe X-linked myotubular myopathy, onychodysplasia, facial dysmorphism, hypotrichosis, Short-rib thoracic dysplasia, Sialic acid storage disease, Sialidosis, Sideroblastic anemia, Small fiber neuropathy, Smith-Magenis syndrome, Sorsby fundus dystrophy, Spastic ataxia, Spastic paraplegia, Spermatogenic failure, Spherocytosis, Sphingomyelin/cholesterol lipidosis, Spinocerebellar ataxia, Split-hand/foot malformation, Spondyloepimetaphyseal dysplasia, Platyspondylic lethal skeletal dysplasia, Squamous cell carcinoma of the head and neck, Stargardt disease, Sucrase-isomaltase deficiency, Sudden infant death syndrome, Supravalvar aortic stenosis, Surfactant metabolism dysfunction, Tangier disease, Tatton-Brown-rahman syndrome, Thoracic aortic aneurysms and aortic dissections, Thrombophilia, Thyroid hormone resistance, TNF receptor-associated periodic fever syndrome (TRAPS), Tooth agenesis, Torsades de pointes, Transposition of great arteries, Treacher Collins syndrome, Tuberous sclerosis syndrome, Tyrosinase-negative oculocutaneous albinism, Tyrosinase-positive oculocutaneous albinism, Tyrosinemia, UDPglucose-4-epimerase deficiency, Ullrich congenital muscular dystrophy, Bethlem myopathy Usher syndrome, UV-sensitive syndrome, Van der Woude syndrome, popliteal pterygium syndrome, Very long chain acyl-CoA dehydrogenase deficiency, Vesicoureteral reflux, Vitreoretinopathology, Von Hippel-Lindau syndrome, von Willebrand disease, Waardenburg syndrome, Warsaw breakage syndrome, WFS1-Related Disorders, Wilson disease, Xeroderma pigmentosum, X-linked agammaglobulinemia, X-linked hereditary motor and sensory neuropathy, X-linked severe combined immunodeficiency, and Zellweger syndrome.

[00319] The development of nucleobase editing advances both the scope and effectiveness of genome editing. The nucleobase editors described here offer researchers a choice of editing with virtually no indel formation (NBE2), or more efficient editing with a low frequency (here, typically $\leq 1\%$) of indel formation (NBE3). That the product of base editing is, by definition, no longer a substrate likely contributes to editing efficiency by preventing subsequent product transformation, which can hamper traditional Cas9 applications. By removing the reliance on double-stranded DNA cleavage and stochastic DNA repair processes that vary greatly by cell state and cell type, nucleobase editing has the potential to expand the type of genome modifications that can be cleanly installed, the efficiency of these modifications, and the type of cells that are amenable to editing. It is likely that recent engineered Cas9 variants^{69,70, 82} or delivery methods⁷¹

with improved DNA specificity, as well as Cas9 variants with altered PAM specificities,⁷² can be integrated into this strategy to provide additional nucleobase editors with improved DNA specificity or that can target an even wider range of disease-associated mutations. These findings also suggest that engineering additional fusions of dCas9 with enzymes that catalyze additional nucleobase transformations will increase the fraction of the possible DNA base changes that can be made through nucleobase editing. These results also suggest architectures for the fusion of other DNA-modifying enzymes, including methylases and demethylases, that may enable additional types of programmable genome and epigenome base editing.

Materials and Methods

[00320] Cloning. DNA sequences of all constructs and primers used in this paper are listed in the Supplementary Sequences. Plasmids containing genes encoding NBE1, NBE2, and NBE3 will be available from Addgene. PCR was performed using VeraSeq ULtra DNA polymerase (Enzymatics), or Q5 Hot Start High-Fidelity DNA Polymerase (New England Biolabs). NBE plasmids were constructed using USER cloning (New England Biolabs). Deaminase genes were synthesized as gBlocks Gene Fragments (Integrated DNA Technologies), and Cas9 genes were obtained from previously reported plasmids.¹⁸ Deaminase and fusion genes were cloned into pCMV (mammalian codon-optimized) or pET28b (*E. coli* codon-optimized) backbones. sgRNA expression plasmids were constructed using site-directed mutagenesis. Briefly, the primers listed in the Supplementary Sequences were 5' phosphorylated using T4 Polynucleotide Kinase (New England Biolabs) according to the manufacturer's instructions. Next, PCR was performed using Q5 Hot Start High-Fidelity Polymerase (New England Biolabs) with the phosphorylated primers and the plasmid pFYF1320 (EGFP sgRNA expression plasmid) as a template according to the manufacturer's instructions. PCR products were incubated with DpnI (20 U, New England Biolabs) at 37 °C for 1 h, purified on a QIAprep spin column (Qiagen), and ligated using QuickLigase (New England Biolabs) according to the manufacturer's instructions. DNA vector amplification was carried out using Mach1 competent cells (ThermoFisher Scientific).

[00321] *In vitro* deaminase assay on ssDNA. Sequences of all ssDNA substrates are listed in the Supplementary Sequences. All Cy3-labelled substrates were obtained from Integrated DNA Technologies (IDT). Deaminases were expressed *in vitro* using the TNT T7 Quick Coupled Transcription/Translation Kit (Promega) according to the manufacturer's instructions using 1 µg of plasmid. Following protein expression, 5 µL of lysate was combined with 35 µL of ssDNA (1.8

μM) and USER enzyme (1 unit) in CutSmart buffer (New England Biolabs) (50 mM potassium acetate, 29 mM Trisacetate, 10 mM magnesium acetate, 100 $\mu\text{g}/\text{mL}$ BSA, pH 7.9) and incubated at 37 °C for 2 h. Cleaved U-containing substrates were resolved from full-length unmodified substrates on a 10% TBE-urea gel (Bio-Rad).

[00322] Expression and purification of His₆-rAPOBEC1-linker-dCas9 fusions. *E. Coli* BL21 STAR (DE3)-competent cells (ThermoFisher Scientific) were transformed with plasmids encoding pET28b-His₆-rAPOBEC1-linker-dCas9 with GGS, (GGS)₃, (SEQ ID NO: 596) XTEN, or (GGS)₇ (SEQ ID NO: 597) linkers. The resulting expression strains were grown overnight in Luria-Bertani (LB) broth containing 100 $\mu\text{g}/\text{mL}$ of kanamycin at 37 °C. The cells were diluted 1:100 into the same growth medium and grown at 37 °C to OD₆₀₀ = ~0.6. The culture was cooled to 4 °C over a period of 2 h, and isopropyl - β -D-1- thiogalactopyranoside (IPTG) was added at 0.5 mM to induce protein expression. After ~16 h, the cells were collected by centrifugation at 4,000 g and resuspended in lysis buffer (50 mM tris(hydroxymethyl)-aminomethane (Tris)-HCl, pH 7.0, 1 M NaCl, 20% glycerol, 10 mM tris(2-carboxyethyl)phosphine (TCEP, Soltec Ventures)). The cells were lysed by sonication (20 s pulse-on, 20 s pulse-off for 8 min total at 6 W output) and the lysate supernatant was isolated following centrifugation at 25,000 g for 15 min. The lysate was incubated with His-Pur nickel-nitriloacetic acid (nickel-NTA) resin (ThermoFisher Scientific) at 4 °C for 1 h to capture the His-tagged fusion protein. The resin was transferred to a column and washed with 40 mL of lysis buffer. The His-tagged fusion protein was eluted in lysis buffer supplemented with 285 mM imidazole, and concentrated by ultrafiltration (Amicon-Millipore, 100-kDa molecular weight cut-off) to 1 mL total volume. The protein was diluted to 20 mL in low-salt purification buffer containing 50 mM tris(hydroxymethyl)-aminomethane (Tris)-HCl, pH 7.0, 0.1 M NaCl, 20% glycerol, 10 mM TCEP and loaded onto SP Sepharose Fast Flow resin (GE Life Sciences). The resin was washed with 40 mL of this low-salt buffer, and the protein eluted with 5 mL of activity buffer containing 50 mM tris(hydroxymethyl)-aminomethane (Tris)-HCl, pH 7.0, 0.5 M NaCl, 20% glycerol, 10 mM TCEP. The eluted proteins were quantified on a SDSPAGE gel.

[00323] *In vitro* transcription of sgRNAs. Linear DNA fragments containing the T7 promoter followed by the 20-bp sgRNA target sequence were transcribed *in vitro* using the primers listed in the Supplementary Sequences with the TranscriptAid T7 High Yield Transcription Kit (ThermoFisher Scientific) according to the manufacturer's instructions. sgRNA products were

purified using the MEGAclear Kit (ThermoFisher Scientific) according to the manufacturer's instructions and quantified by UV absorbance.

[00324] Preparation of Cy3-conjugated dsDNA substrates. Sequences of 80-nucleotide unlabeled strands are listed in the Supplementary Sequences and were ordered as PAGE-purified oligonucleotides from IDT. The 25-nt Cy3-labeled primer listed in the Supplementary Sequences is complementary to the 3' end of each 80-nt substrate. This primer was ordered as an HPLC-purified oligonucleotide from IDT. To generate the Cy3-labeled dsDNA substrates, the 80-nt strands (5 μ L of a 100 μ M solution) were combined with the Cy3-labeled primer (5 μ L of a 100 μ M solution) in NEBuffer 2 (38.25 μ L of a 50 mM NaCl, 10 mM Tris-HCl, 10 mM MgCl₂, 1 mM DTT, pH 7.9 solution, New England Biolabs) with dNTPs (0.75 μ L of a 100 mM solution) and heated to 95 °C for 5 min, followed by a gradual cooling to 45 °C at a rate of 0.1 °C/s. After this annealing period, Klenow exo⁻ (5 U, New England Biolabs) was added and the reaction was incubated at 37 °C for 1 h. The solution was diluted with Buffer PB (250 μ L, Qiagen) and isopropanol (50 μ L) and purified on a QIAprep spin column (Qiagen), eluting with 50 μ L of Tris buffer.

[00325] Deaminase assay on dsDNA. The purified fusion protein (20 μ L of 1.9 μ M in activity buffer) was combined with 1 equivalent of appropriate sgRNA and incubated at ambient temperature for 5 min. The Cy3-labeled dsDNA substrate was added to final concentration of 125 nM and the resulting solution was incubated at 37 °C for 2 h. The dsDNA was separated from the fusion by the addition of Buffer PB (100 μ L, Qiagen) and isopropanol (25 μ L) and purified on a EconoSpin micro spin column (Epoch Life Science), eluting with 20 μ L of CutSmart buffer (New England Biolabs). USER enzyme (1 U, New England Biolabs) was added to the purified, edited dsDNA and incubated at 37 °C for 1 h. The Cy3-labeled strand was fully denatured from its complement by combining 5 μ L of the reaction solution with 15 μ L of a DMSO-based loading buffer (5 mM Tris, 0.5 mM EDTA, 12.5% glycerol, 0.02% bromophenol blue, 0.02% xylene cyan, 80% DMSO). The full-length C-containing substrate was separated from any cleaved, U-containing edited substrates on a 10% TBE-urea gel (Bio-Rad) and imaged on a GE Amersham Typhoon imager.

[00326] Preparation of *in vitro*-edited dsDNA for high-throughput sequencing (HTS). The oligonucleotides listed in the Supplementary Sequences were obtained from IDT. Complementary sequences were combined (5 μ L of a 100 μ M solution) in Tris buffer and annealed by heating to 95 °C for 5 min, followed by a gradual cooling to 45 °C at a rate of 0.1 °C/s to generate 60-bp

dsDNA substrates. Purified fusion protein (20 μ L of 1.9 μ M in activity buffer) was combined with 1 equivalent of appropriate sgRNA and incubated at ambient temperature for 5 min. The 60-mer dsDNA substrate was added to final concentration of 125 nM and the resulting solution was incubated at 37 °C for 2 h. The dsDNA was separated from the fusion by the addition of Buffer PB (100 μ L, Qiagen) and isopropanol (25 μ L) and purified on a EconoSpin micro spin column (Epoch Life Science), eluting with 20 μ L of Tris buffer. The resulting edited DNA (1 μ L was used as a template) was amplified by PCR using the HTS primer pairs specified in the Supplementary Sequences and VeraSeq Ultra (Enzymatics) according to the manufacturer's instructions with 13 cycles of amplification. PCR reaction products were purified using RapidTips (Diffinity Genomics), and the purified DNA was amplified by PCR with primers containing sequencing adapters, purified, and sequenced on a MiSeq high-throughput DNA sequencer (Illumina) as previously described.⁷³

[00327] Cell culture. HEK293T (ATCC CRL-3216), U2OS (ATCC-HTB-96) and ST486 cells (ATCC) were maintained in Dulbecco's Modified Eagle's Medium plus GlutaMax (ThermoFisher) supplemented with 10% (v/v) fetal bovine serum (FBS) and penicillin/streptomycin (1x, Amresco), at 37 °C with 5% CO₂. HCC1954 cells (ATCC CRL-2338) were maintained in RPMI-1640 medium (ThermoFisher Scientific) supplemented as described above. Immortalized rat astrocytes containing the ApoE4 isoform of the *APOE* gene (Taconic Biosciences) were cultured in Dulbecco's Modified Eagle's Medium plus GlutaMax (ThermoFisher Scientific) supplemented with 10% (v/v) fetal bovine serum (FBS) and 200 μ g/mL Geneticin (ThermoFisher Scientific).

[00328] Transfections. HEK293T cells were seeded on 48-well collagen-coated BioCoat plates (Corning) and transfected at approximately 85% confluency. Briefly, 750 ng of NBE and 250 ng of sgRNA expression plasmids were transfected using 1.5 μ l of Lipofectamine 2000 (ThermoFisher Scientific) per well according to the manufacturer's protocol. Astrocytes, U2OS, HCC1954, HEK293T and ST486 cells were transfected using appropriate AMAXA NUCLEOFECTOR™ II programs according to manufacturer's instructions. 40 ng of infrared RFP (Addgene plasmid 45457)⁷⁴ was added to the nucleofection solution to assess nucleofection efficiencies in these cell lines. For astrocytes, U2OS, and ST486 cells, nucleofection efficiencies were 25%, 74%, and 92%, respectively. For HCC1954 cells, nucleofection efficiency was <10%. Therefore, following trypsinization, the HCC1954 cells were filtered through a 40 micron strainer (Fisher Scientific), and the nucleofected HCC1954 cells were collected on a Beckman Coulter

MoFlo XDP Cell Sorter using the iRFP signal (abs 643 nm, em 670 nm). The other cells were used without enrichment of nucleofected cells.

[00329] High-throughput DNA sequencing of genomic DNA samples. Transfected cells were harvested after 3 d and the genomic DNA was isolated using the Agencourt DNAdvance Genomic DNA Isolation Kit (Beckman Coulter) according to the manufacturer's instructions. On-target and off-target genomic regions of interest were amplified by PCR with flanking HTS primer pairs listed in the Supplementary Sequences. PCR amplification was carried out with Phusion high-fidelity DNA polymerase (ThermoFisher) according to the manufacturer's instructions using 5 ng of genomic DNA as a template. Cycle numbers were determined separately for each primer pair as to ensure the reaction was stopped in the linear range of amplification (30, 28, 28, 28, 32, and 32 cycles for EMX1, FANCF, HEK293 site 2, HEK293 site 3, HEK293 site 4, and RNF2 primers, respectively). PCR products were purified using RapidTips (Diffinity Genomics). Purified DNA was amplified by PCR with primers containing sequencing adaptors. The products were gel-purified and quantified using the QUANT-ITTM PicoGreen dsDNA Assay Kit (ThermoFisher) and KAPA Library Quantification Kit-Illumina (KAPA Biosystems). Samples were sequenced on an Illumina MiSeq as previously described.⁷³

[00330] Data analysis. Sequencing reads were automatically demultiplexed using MiSeq Reporter (Illumina), and individual FASTQ files were analyzed with a custom Matlab script provided in the Supplementary Notes. Each read was pairwise aligned to the appropriate reference sequence using the Smith-Waterman algorithm. Base calls with a Q-score below 31 were replaced with N's and were thus excluded in calculating nucleotide frequencies. This treatment yields an expected MiSeq base-calling error rate of approximately 1 in 1,000. Aligned sequences in which the read and reference sequence contained no gaps were stored in an alignment table from which base frequencies could be tabulated for each locus.

[00331] Indel frequencies were quantified with a custom Matlab script shown in the Supplementary Notes using previously described criteria⁷¹. Sequencing reads were scanned for exact matches to two 10-bp sequences that flank both sides of a window in which indels might occur. If no exact matches were located, the read was excluded from analysis. If the length of this indel window exactly matched the reference sequence the read was classified as not containing an indel. If the indel window was two or more bases longer or shorter than the reference sequence, then the sequencing read was classified as an insertion or deletion, respectively.

[00332] All publications, patents, patent applications, publication, and database entries (*e.g.*, sequence database entries) mentioned herein, *e.g.*, in the Background, Summary, Detailed Description, Examples, and/or References sections, are hereby incorporated by reference in their entirety as if each individual publication, patent, patent application, publication, and database entry was specifically and individually incorporated herein by reference. In case of conflict, the present application, including any definitions herein, will control.

Supplementary Sequences

[00333] Primers used for generating sgRNA transfection plasmids. rev_sgRNA_plasmid was used in all cases. The pFYF1320 plasmid was used as template as noted in Materials and Methods section. SEQ ID NOs: 329-338 appear from top to bottom below, respectively.

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rev_sgRNA_plasmid    GGTGTTTCGTCCTTTCCACAAG
fwd_p53_Y163C       GCTTGCAGATGGCCATGGCGGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC
fwd_p53_N239D       TGTCACACATGTAGTTGTAGGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC
fwd_APOE4_C158R     GAAGCGCCTGGCAGTGTACCGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC
fwd_EMX1            GAGTCCGAGCAGAAGAAGAAGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC
fwd_FANCF           GGAATCCCTTCTGCAGCACCGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC
fwd_HEK293_2        GAACACAAAGCATAGACTGCGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC
fwd_HEK293_3        GGCCCAGACTGAGCACGTGAGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC
fwd_HEK293_4        GGCACCTGCGGCTGGAGGTGGGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC
fwd_RNF2            GTCATCTTAGTCATTACCTGGTTTTAGAGCTAGAAATAGCAAGTTAAAATAAGGC

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[00334] Sequences of all ssDNA substrates used in in vitro deaminase assays. SEQ ID NOs: 339-341 appear from top to bottom below, respectively.

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rAPOBEC1 substrate   Cy3-ATTATTATTATTCCGCGGATTTATTTATTTATTTATTTATTT
hAID/pmCDA1 substrate Cy3-ATTATTATTATTAGCTATTTATTTATTTATTTATTTATTT
hAPOBEC3G substrate  Cy3-ATTATTATTATTCCCGGATTTATTTATTTATTTATTTATTT

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[00335] Primers used for generating PCR products to serve as substrates for T7 transcription of sgRNAs for gel-based deaminase assay. rev_gRNA_T7 was used in all cases.

The pFYF1320 plasmid was used as template as noted in Materials and Methods section. SEQ ID NOs: 342-365 appear from top to bottom below, respectively.

rev_sgRNA_T7	AAAAAAAGCACCGACTCGGTG
fwd_sgRNA_T7_dsDNA_2	TAATACGACTCACTATAGGCCGCGGATTTATTTATTTAAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_3	TAATACGACTCACTATAGGTCCGCGGATTTATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_4	TAATACGACTCACTATAGGTTCCGCGGATTTATTTATTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_5	TAATACGACTCACTATAGGATTCGCGGATTTATTTATTGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_6	TAATACGACTCACTATAGGTATTCGCGGATTTATTTATGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_7	TAATACGACTCACTATAGGTTATTCCGCGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_8	TAATACGACTCACTATAGGATTATTCCGCGGATTTATTTGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_9	TAATACGACTCACTATAGGTATTATTCCGCGGATTTATTGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_10	TAATACGACTCACTATAGGATTATTATCCGCGGATTTATGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_11	TAATACGACTCACTATAGGTATTATATCCGCGGATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_12	TAATACGACTCACTATAGGTTATTATATTCCGCGGATTTGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_13	TAATACGACTCACTATAGGATTATTATATTCCGCGGATTTGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_14	TAATACGACTCACTATAGGTATTATTATATTCCGCGGATGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_15	TAATACGACTCACTATAGGATTATTATTATTACCOCGGAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_18	TAATACGACTCACTATAGGATTATTATTATTATTACCOCGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_ncC	TAATACGACTCACTATAGGATATTAATTTATTTATTTAAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_APOE4_C112R	TAATACGACTCACTATAGGGGAGGACGTGCGCGGCCCGCTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_APOE4_C158R	TAATACGACTCACTATAGGGAAGCGCCTGGCAGTGTACCGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_CTNNB1_T41A	TAATACGACTCACTATAGGCTGTGGCAGTGGCACCCAGAAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_HRAS_Q61R	TAATACGACTCACTATAGGCOCTCCCGGCCGGCGGTATCCGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_53_Y163C	TAATACGACTCACTATAGGGCTTGCAGATGGCCATGGCGGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_53_Y236C	TAATACGACTCACTATAGGACACATGCAGTTGTAGTGGAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_dsDNA_53_N239D	TAATACGACTCACTATAGGTGTCACACATGTAGTTGTAGGTTTTAGAGCTAGAAATAGCA

[00336] Sequences of 80-nucleotide unlabeled strands and Cy3-labeled universal primer used in gel-based dsDNA deaminase assays. SEQ ID NOs: 366-390 appear from top to bottom below, respectively.

Cy3-primer	Cy3-GTAGGTAGTTAGGATGAATGGAAGGTTGGTA
dsDNA_2	GTCCATGGATCCAGAGGTCATCCATTAATAATAATAATCCGCGGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_3	GTCCATGGATCCAGAGGTCATCCATAAATAATAATAATCCGCGGAAGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_4	GTCCATGGATCCAGAGGTCATCCATAAATAATAATCCGCGGAAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_5	GTCCATGGATCCAGAGGTCATCCAAAATAATAATAATCCGCGGAATGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_6	GTCCATGGATCCAGAGGTCATCCAAAATAATAATCCGCGGAATAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_7	GTCCATGGATCCAGAGGTCATCCATAAATAATAATCCGCGGAATAAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_8	GTCCATGGATCCAGAGGTCATCCAAAATAATAATCCGCGGAATAATGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_9	GTCCATGGATCCAGAGGTCATCCAAAATAATAATCCGCGGAATAATAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_10	GTCCATGGATCCAGAGGTCATCCAAAATAATAATCCGCGGATAATAATGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_11	GTCCATGGATCCAGAGGTCATCCATAAATCCGCGGAATAATAAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_12	GTCCATGGATCCAGAGGTCATCCAAAATCCGCGGAATAATAAAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_13	GTCCATGGATCCAGAGGTCATCCAAAATCCGCGGAATAATAAATGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_14	GTCCATGGATCCAGAGGTCATCCAAATCCGCGGAATAATAAATAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_15	GTCCATGGATCCAGAGGTCATCCATCCGCGGAATAATAAATAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_18	GTCCATGGATCCAGAGGTCATCCAGCGGTAATAATAAATAGGCTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_noC	GTCCATGGATCCAGAGGTCATCCATAAATAATAATAATAATTACTATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_8U	5Dy3-GTAGGTAGTTAGGATGAATGGAAGGTTGGTA
dsDNA_APOE_C112R	GCAACTCGCCCGGCTACTGCACCAGGCGCCGCGCACGTCCTCCATGTCTACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_APOE_C158R	CGGCGCCCTGGCGGGCCCGGCTGGTACACTGCCAGGCGCTTCTGCAGTACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_CTNNB1_T41A	GTCTTACCTGGACTCTGGAATCCATTCCTGTGTCACCTGCCACAGTCTCTACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_HRAS_Q61R	GGAGACGTGCTGTTGGACATCCTGATACCGCGGCGGEGAGGAGTACTACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_p53_Y163C	ACCCCGCCCGGCAACCGGCTCCGCGCCATGGCCATCTGCAAGCAGTCATACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_p53_Y238C	AGGTTGGCTCTGACTGTACCAACATCCACTACAACCTGCATGTGTAACAGTACCAACCTTCCATTCATCCTAACTACCTAC
dsDNA_p53_N239D	TGCTCTGACTGTACCAACATCCACTACAACCTACATGTGTGACAGTTCCTACCAACCTTCCATTCATCCTAACTACCTAC

[00337] Primers used for generating PCR products to serve as substrates for T7 transcription of sgRNAs for high-throughput sequencing. rev_gRNA_T7 (above) was used in all cases. The pFYF1320 plasmid was used as template as noted in Materials and Methods section. SEQ ID NOs: 391-442 appear from top to bottom below, respectively.

fwd_sgRNA_T7_HTS_base TAATACGACTCACTATAGGTTATTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_1A TAATACGACTCACTATAGGATATTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_1C TAATACGACTCACTATAGGCTATTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_1G TAATACGACTCACTATAGGGTATTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_2A TAATACGACTCACTATAGGTAATTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_2C TAATACGACTCACTATAGGTCATTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_2G TAATACGACTCACTATAGGTGATTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_3T TAATACGACTCACTATAGGTTTTTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_3C TAATACGACTCACTATAGGTTCTTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_3G TAATACGACTCACTATAGGTTGTTTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_4A TAATACGACTCACTATAGGTTAATTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_4C TAATACGACTCACTATAGGTTACTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_4G TAATACGACTCACTATAGGTTAGTTTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_5A TAATACGACTCACTATAGGTTATATCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_5C TAATACGACTCACTATAGGTTATCTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_5G TAATACGACTCACTATAGGTTATGTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_6A TAATACGACTCACTATAGGTTATTACGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_6C TAATACGACTCACTATAGGTTATTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_6G TAATACGACTCACTATAGGTTATTGCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_8A TAATACGACTCACTATAGGTTATTTTCATGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_8T TAATACGACTCACTATAGGTTATTTCTTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_8C TAATACGACTCACTATAGGTTATTTTCCTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_9A TAATACGACTCACTATAGGTTATTTTCGAGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_9C TAATACGACTCACTATAGGTTATTTTCGCGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_9G TAATACGACTCACTATAGGTTATTTTCGGGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_10A TAATACGACTCACTATAGGTTATTTTCGTAGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_10T TAATACGACTCACTATAGGTTATTTTCGTTGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_10C TAATACGACTCACTATAGGTTATTTTCGTGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_11A TAATACGACTCACTATAGGTTATTTTCGTGAATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_11T TAATACGACTCACTATAGGTTATTTTCGTGTATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_11C TAATACGACTCACTATAGGTTATTTTCGTGCATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_12T TAATACGACTCACTATAGGTTATTTTCGTGGTTTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_12C TAATACGACTCACTATAGGTTATTTTCGTGGCTTTATTTAGTTTTAGAGCTAGAAATAGCA

fwd_sgRNA_T7_HTS_12G	TAATACGACTCACTATAGGTTATTTTCGTGGGTTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_13A	TAATACGACTCACTATAGGTTATTTTCGTGGAATTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_13C	TAATACGACTCACTATAGGTTATTTTCGTGGACTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_13G	TAATACGACTCACTATAGGTTATTTTCGTGGAGTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_multic	TAATACGACTCACTATAGGTTCCCCCCCCGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_TCGCACCC_odd	TAATACGACTCACTATAGGGCGACCCCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_CCTCGCAC_odd	TAATACGACTCACTATAGGCTCGCACGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_ACCCTCGC_odd	TAATACGACTCACTATAGGCCOCTCGCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_GCACCCTC_odd	TAATACGACTCACTATAGGCCACCCCTCGTGGATTTATTTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_TCGCACCC_even	TAATACGACTCACTATAGGTCGCACCCCGTGGATTTATTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_CCTCGCAC_even	TAATACGACTCACTATAGGCCTCGCACGTGGATTTATTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_ACCCTCGC_even	TAATACGACTCACTATAGGACCCTCGCGTGGATTTATTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_GCACCCTC_even	TAATACGACTCACTATAGGGCACCCCTCGTGGATTTATTAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_EMX1	TAATACGACTCACTATAGGGAGTCCGAGCAGAAGAAGAAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_FANCF	TAATACGACTCACTATAGGGGAATCCCTTCTGCAGCAOCCGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_HEK293_site2	TAATACGACTCACTATAGGGAACACAAAGCATAGACTGCGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_HEK293_site3	TAATACGACTCACTATAGGGGCCAGACTGAGCAOCTGAGTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_HEK293_site4	TAATACGACTCACTATAGGGGCACTGCGGCTGGAGGTGGSTTTTAGAGCTAGAAATAGCA
fwd_sgRNA_T7_HTS_RNF2	TAATACGACTCACTATAGGGTCATCTTAGTCATTACCTGGTTTTAGAGCTAGAAATAGCA

[00338] Sequences of *in vitro*-edited dsDNA for high-throughput sequencing (HTS). Shown are the sequences of edited strands. Reverse complements of all sequences shown were also obtained. dsDNA substrates were obtained by annealing complementary strands as described in Materials and Methods. Oligonucleotides representing the EMX1, FANCF, HEK293 site 2, HEK293 site 3, HEK293 site 4, and RNF2 loci were originally designed for use in the gel-based deaminase assay and therefore have the same 25-nt sequence on their 5'- ends (matching that of the Cy3-primer). SEQ ID NOs: 443-494 appear from top to bottom below, respectively.

Base sequence	ACGTAAACGGCCACAAGTTCTTATTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
1A	ACGTAAACGGCCACAAGTTCATATTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
1C	ACGTAAACGGCCACAAGTTCTTATTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
1G	ACGTAAACGGCCACAAGTTCGTATTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
2A	ACGTAAACGGCCACAAGTTCATATTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
2C	ACGTAAACGGCCACAAGTTCTCATTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
2G	ACGTAAACGGCCACAAGTTCGTATTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
3T	ACGTAAACGGCCACAAGTTCCTTTTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
3C	ACGTAAACGGCCACAAGTTCCTTCTTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
3G	ACGTAAACGGCCACAAGTTCCTGTTTTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
4A	ACGTAAACGGCCACAAGTTCCTTAATTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
4C	ACGTAAACGGCCACAAGTTCCTTACTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
4G	ACGTAAACGGCCACAAGTTCCTTAGTTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
5A	ACGTAAACGGCCACAAGTTCCTATATTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
5C	ACGTAAACGGCCACAAGTTCCTTATCTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
5G	ACGTAAACGGCCACAAGTTCCTTATGTCTGTGGATTTATTTATGGCATCTTCTTCAAGGACG
6A	ACGTAAACGGCCACAAGTTCCTTATTACGTGGATTTATTTATGGCATCTTCTTCAAGGACG
6C	ACGTAAACGGCCACAAGTTCCTTATTCCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
6G	ACGTAAACGGCCACAAGTTCCTTATTGCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
8A	ACGTAAACGGCCACAAGTTCCTTATTTTCATGGATTTATTTATGGCATCTTCTTCAAGGACG
8T	ACGTAAACGGCCACAAGTTCCTTATTTCTTGGATTTATTTATGGCATCTTCTTCAAGGACG
8C	ACGTAAACGGCCACAAGTTCCTTATTTTCCTGGATTTATTTATGGCATCTTCTTCAAGGACG
9A	ACGTAAACGGCCACAAGTTCCTTATTTTCGAGGATTTATTTATGGCATCTTCTTCAAGGACG
9C	ACGTAAACGGCCACAAGTTCCTTATTTTCGCGGATTTATTTATGGCATCTTCTTCAAGGACG
9G	ACGTAAACGGCCACAAGTTCCTTATTTTCGGGGATTTATTTATGGCATCTTCTTCAAGGACG
10A	ACGTAAACGGCCACAAGTTCCTTATTTTCGTAGATTTATTTATGGCATCTTCTTCAAGGACG
10T	ACGTAAACGGCCACAAGTTCCTTATTTTCGTTGATTTATTTATGGCATCTTCTTCAAGGACG
10C	ACGTAAACGGCCACAAGTTCCTTATTTTCGTGATTTATTTATGGCATCTTCTTCAAGGACG
11A	ACGTAAACGGCCACAAGTTCCTTATTTTCGTGAATTTATTTATGGCATCTTCTTCAAGGACG
11T	ACGTAAACGGCCACAAGTTCCTTATTTTCGTGTATTTATTTATGGCATCTTCTTCAAGGACG
11C	ACGTAAACGGCCACAAGTTCCTTATTTTCGTGCATTTATTTATGGCATCTTCTTCAAGGACG

12T ACGTAAACGGCCACAAGTTCTTATTTTCGTGGTTTTATTTATGGCATCTTCTTCAAGGACG
12C ACGTAAACGGCCACAAGTTCTTATTTTCGTGGCTTTATTTATGGCATCTTCTTCAAGGACG
12G ACGTAAACGGCCACAAGTTCTTATTTTCGTGGGTTTATTTATGGCATCTTCTTCAAGGACG
13A ACGTAAACGGCCACAAGTTCTTATTTTCGTGGAATTATTTATGGCATCTTCTTCAAGGACG
13C ACGTAAACGGCCACAAGTTCTTATTTTCGTGGACTTATTTATGGCATCTTCTTCAAGGACG
13G ACGTAAACGGCCACAAGTTCTTATTTTCGTGGAGTTATTTATGGCATCTTCTTCAAGGACG
multic ACGTAAACGGCCACAAGTTCTTCCCCCCCCGATTTATTTATGGCATCTTCTTCAAGGACG
TCGCACCC_odd ACGTAAACGGCCACAAGTTTCGCACCCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
CCTCGCAC_odd ACGTAAACGGCCACAAGTTCTCGCACGTGGATTTATTTATGGCATCTTCTTCAAGGACG
ACCCTCGC_odd ACGTAAACGGCCACAAGTTACCCTCGCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
GCACCCTC_odd ACGTAAACGGCCACAAGTTGCACCCCTCGTGGATTTATTTATGGCATCTTCTTCAAGGACG
TCGCACCC_even ACGTAAACGGCCACAAGTATTCGCACCCGTGGATTTATTATGGCATCTTCTTCAAGGACG
CCTCGCAC_even ACGTAAACGGCCACAAGTATCCTCGCACGTGGATTTATTATGGCATCTTCTTCAAGGACG
ACCCTCGC_even ACGTAAACGGCCACAAGTATACCCTCGCGTGGATTTATTATGGCATCTTCTTCAAGGACG
GCACCCTC_even ACGTAAACGGCCACAAGTATGCACCCCTCGTGGATTTATTATGGCATCTTCTTCAAGGACG
EMX1_invitro GTAGGTAGTTAGGATGAATGGAAGGTTGGTAGGCCTGAGTCCGAGCAGAAGAAGAAGGGCTCCCATCACATCAACCGGTG
FANCF_invitro GTAGGTAGTTAGGATGAATGGAAGGTTGGTACTCATGGAATCCCTTCTGCAGCACCTGGATCGCTTTCCGAGCTTCTGG
HEK293_site2_invitro GTAGGTAGTTAGGATGAATGGAAGGTTGGTAAACTGGAACACAAAGCATAGACTGCGGGGCGGGCCAGCCTGAATAGCTG
HEK293_site3_invitro GTAGGTAGTTAGGATGAATGGAAGGTTGGTACTTGGGGCCAGACTGAGCACGTGATGGCAGAGGAAAGGAAGCCCTGCT
HEK293_site4_invitro GTAGGTAGTTAGGATGAATGGAAGGTTGGTACCCTGGCACTCCGGCTGGAGGTGGGGTTAAAGCGGAGACTCTGGTGC
RNF2_invitro GTAGGTAGTTAGGATGAATGGAAGGTTGGTATGGCAGTCATCTTAGTCATTACCTGAGGTGTTCCGTTGTAACCTCATATAA

[00339] Primers for HTS of *in vitro* edited dsDNA. SEQ ID NOs: 495-503 appear from top to bottom below, respectively.

fwd_invitro_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNACGTAAACGGCCACAA
rev_invitro_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCGTCTTGAAGAAGATGC
fwd_invitro_HEK_targets ACACTCTTTCCCTACACGACGCTCTTCCGATCTNMMNGTAGGTAGTTAGGATGAATGGAA
rev_EMX1_invitro TGGAGTTCAGACGTGTGCTCTTCCGATCTCACCAGTTGATGTGATGG
rev_FANCF_invitro TGGAGTTCAGACGTGTGCTCTTCCGATCTCCAGAAGCTCGGAAAAGC
rev_HEK293_site2_invitro TGGAGTTCAGACGTGTGCTCTTCCGATCTCAGCTATTCAGGCTGGC
rev_HEK293_site3_invitro TGGAGTTCAGACGTGTGCTCTTCCGATCTAGCAGGGCTTCCCTTC
rev_HEK293_site4_invitro TGGAGTTCAGACGTGTGCTCTTCCGATCTGCACCAGAGTCTCCG
rev_RNF2_invitro TGGAGTTCAGACGTGTGCTCTTCCGATCTTTATATGAGTTACAACGAACACC

[00340] Primers for HTS of on-target and off-target sites from all mammalian cell culture experiments. SEQ ID NOs: 504-579 and 1869-1900 appear from top to bottom below, respectively.

fwd_EMX1_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCAGCTCAGCCTGAGTGTGA
rev_EMX1_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTCTCGTGGGTTTGTGGTTGC
fwd_FANCF_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCATTGCAGAGAGGCGTATCA
rev_FANCF_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTGGGGTCCCAGGTGCTGAC
fwd_HEK293_site2_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNCCAGCCCCATCTGTCAAAC
rev_HEK293_site2_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTTGAATGGATTCCCTGGAAACAATGA
fwd_HEK293_site3_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNATGTGGGCTGCCTAGAAAAGG
rev_HEK293_site3_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTCCCAGCCAAACTTGTCAACC
fwd_HEK293_site4_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNGAACCCAGGTAGCCAGAGAC
rev_HEK293_site4_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTTCCCTTCAACCCGAACGGAG
fwd_RNF2_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCTCTTCTTTATTTCCAGCAATGT
rev_RNF2_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTGTTTTTCATGTTCTAAAAATGTATCCCA
fwd_p53_Y163C_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNTACAGTACTCCCCTGCCCTC
rev_p53_Y163C_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTGCTGCTCACCATCGCTATCT
fwd_p53_N239D_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCCTCATCTTGGGCCTGTGTT
rev_p53_N239D_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTAAATCGGTAAGAGGTGGGCC
fwd_APOE4_C158R_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNGCGGACATGGAGGACGTG
rev_APOE4_C158R_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTCTGTTCCACCAGGGGGCCC
fwd_EMX1_off1_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNTGCCCAATCATTGATGCTTTT
rev_EMX1_off1_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTAGAAACATTTACCATAGACTATCACCT
fwd_EMX1_off2_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNAGTAGCCTCTTTCTCAATGTGC
rev_EMX1_off2_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTGCTTTTCAAGGATGCAGTCT
fwd_EMX1_off3_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNGAGCTAGACTCCGAGGGGA
rev_EMX1_off3_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTTCCCTCGTCTCTGCTCTCACTT
fwd_EMX1_off4_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNAGAGGCTGAAGAGGAAGACCA
rev_EMX1_off4_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTGGCCCAGCTGTGCATTCTAT

fwd_EMX1_off5_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCCAAGAGGGGCCAAGTCTG
 rev_EMX1_off5_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCAGCGAGGAGTGACAGCC
 fwd_EMX1_off7_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCACTCCACCTGATCTCGGGG
 rev_EMX1_off7_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCGAGGAGGGAGGGAGCAG
 fwd_EMX1_off8_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNACCACAAATGCCCAAGAGAC
 rev_EMX1_off8_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGACACAGTCAAGGGCCGG
 fwd_EMX1_off9_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCCACCTTTGAGGAGGCCAAA
 rev_EMX1_off9_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTTTCCATCTGAGAAGAGAGTGGT
 fwd_EMX1_off10_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNGTCATACCTTGGCCCTTCT
 rev_EMX1_off10_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTTCCCTAGGCCACACCAG
 fwd_FANCF_off1_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNAACCCACTGAAGAAGCAGGG
 rev_FANCF_off1_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGGTGCTTAATCCGGCTCCAT
 fwd_FANCF_off2_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNTCCAGTGTTTCCATCCCGAA
 rev_FANCF_off2_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCCTCTGACCTCCACAADTCT
 fwd_FANCF_off3_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCTGGGTACAGTTCTGCGTGT
 rev_FANCF_off3_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTTCACTCTGAGCATCGCCAAG
 fwd_FANCF_off4_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNGGTTTAGAGCCAGTGAAGTAGAG
 rev_FANCF_off4_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGCAAGACAAAATCCTCTTTATACTTTG
 fwd_FANCF_off5_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNGGGAGGGGACGGCCTTAC
 rev_FANCF_off5_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGCCTCTGGCGAACATGGC
 fwd_FANCF_off6_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNTCCTGGTTAAGAGCATGGGC
 rev_FANCF_off6_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGATTGAGTCCCCACAGCACA
 fwd_FANCF_off7_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCCAGTGTTTCCCATCCCCAA
 rev_FANCF_off7_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTTGACCTCCACAACCTGGAAAAT
 fwd_FANCF_off8_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNGCTTCCAGACCCACCTGAAG
 rev_FANCF_off8_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTACCGAGGAAAATTGCTTGTCG
 fwd_HEK293_site2_off1_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNGTGTGGAGAGTGAGTAAGCCA
 rev_HEK293_site2_off1_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTACGGTAGGATGATTTGAGGCA
 fwd_HEK293_site2_off2_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNCACAAAGCAGTGTAGCTCAGG
 rev_HEK293_site2_off2_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTTTTTTGGTACTCGAGTGTATTTCAG
 fwd_HEK293_site3_off1_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNTCCCTGTTGACCTGGAGAA
 rev_HEK293_site3_off1_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCACTGTACTTGCCCTGACCA
 fwd_HEK293_site3_off2_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNTTGGTGTGACAGGGAGCAA
 rev_HEK293_site3_off2_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCTGAGATGTGGGCAGAAAGGG
 fwd_HEK293_site3_off3_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNTGAGAGGGGAACAGAAGGGCT
 rev_HEK293_site3_off3_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGTCCAAAGGCCCAAGAACCT
 fwd_HEK293_site3_off4_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNTCCTAGCACTTTGGAAGGTCTG
 rev_HEK293_site3_off4_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGCTCATCTTAATCTGCTCAGCC
 fwd_HEK293_site3_off5_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTNNNNNAAAGGAGCAGCTCTTCTCTGG
 rev_HEK293_site3_off5_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGTCTGCACCATCTCCACAA

fwd_HEK293_site4_off1_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNGGCATGGOTTCTGAGACTCA
 rev_HEK293_site4_off1_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGTCTCCCTTGCACTCCCTGTCTTT
 fwd_HEK293_site4_off2_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNTTTGGCAATGGAGGCATTGG
 rev_HEK293_site4_off2_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGAAGAGGCTGCCDATGAGAG
 fwd_HEK293_site4_off3_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNGGCTGAGGCTCGAATCCTG
 rev_HEK293_site4_off3_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCTGTGGCCTCCATATCCCTG
 fwd_HEK293_site4_off4_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNTTTCCACCAGAACTCAGCCC
 rev_HEK293_site4_off4_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCCTCGTTCTCCACAACAC
 fwd_HEK293_site4_off5_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNCACGGGAAGGACAGGAGAAG
 rev_HEK293_site4_off5_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGCAGGGGAGGGATAAAGCAG
 fwd_HEK293_site4_off6_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNCCACGGGAGATGGCTTATGT
 rev_HEK293_site4_off6_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCACATCCTCACTGTGCCACT
 fwd_HEK293_site4_off7_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNGTCAGTCTCGGCCCTCA
 rev_HEK293_site4_off7_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGCCACTGTAAAGCTCTTGGG
 fwd_HEK293_site4_off8_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNNAGGGTAGAGGGACAGAGCTG
 rev_HEK293_site4_off8_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGGACCCACATAGTCAGTGC
 fwd_HEK293_site4_off9_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNNGCTGTCAGCCCTATCTCCATC
 rev_HEK293_site4_off9_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTTGGGCAATTAGGACAGGGAC
 fwd_HEK293_site4_off10_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNNGCAGCGGAGGAGGTAGATTG
 rev_HEK293_site4_off10_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTCTCAGTACCTGGAGTCCCGA
 fwd_HEK2_ChIP_off1_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNNGACAGGCTCAGGAAAGCTGT
 rev_HEK2_ChIP_off1_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTACACAAGCCTTTCTCCAGGG
 fwd_HEK2_ChIP_off2_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNNAATAGGGGGTGAGACTGGGG
 rev_HEK2_ChIP_off2_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTGCCTCAGACGAGACTTGAGG
 fwd_HEK2_ChIP_off3_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNNGGCCAGCAGGAAAGGAATCT
 rev_HEK2_ChIP_off3_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTTGACTGCACCTGTAGCCATG
 fwd_HEK2_ChIP_off4_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNTCAAGGAAATCACCCTGCC
 rev_HEK2_ChIP_off4_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTAACTTCCTTGGTGTGCAGCT
 fwd_HEK2_ChIP_off5_HTS ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNATGGGCTCAGCTACGTCATG
 rev_HEK2_ChIP_off5_HTS TGGAGTTCAGACGTGTGCTCTTCCGATCTAATAGCAGTGTGGTGGGCAA

fwd_HEK3_ChIP_off1_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNCGCACATCCCTTGTCTCTCT
rev_HEK3_ChIP_off1_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTCTACTGGAGCACACCCCAAG
fwd_HEK3_ChIP_off2_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNTGGGTCACGTAGCTTTGGTC
rev_HEK3_ChIP_off2_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTTGGTGGCCATGTGCAACTAA
fwd_HEK3_ChIP_off3_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNCTACTACGTGCCAGCTCAGG
rev_HEK3_ChIP_off3_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTACCTCCCCTCCTCACTAACC
fwd_HEK3_ChIP_off4_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNGCCTCAGCTCCATTTCTGT
rev_HEK3_ChIP_off4_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTAACCTTTATGGCACCAGGGG
fwd_HEK3_ChIP_off5_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNGAGCTCAGCATTAGCAGGCT
rev_HEK3_ChIP_off5_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTTCCCTGGCTTTCCGATTCCC
fwd_HEK4_ChIP_off1_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNGTGCAATTGGAGGAGGAGCT
rev_HEK4_ChIP_off1_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTCACCAGCTACAGGCAGAACA
fwd_HEK4_ChIP_off3_HTS	ACACTCTTTCCCTACACGACGCTCTTCCGATCTN>NNNCCTACCCCCAACACAGATGG
rev_HEK4_ChIP_off3_HTS	TGGAGTTCAGACGTGTGCTCTTCCGATCTCCACACAACCTCAGGTCCTCC

[00341] Sequences of single-stranded oligonucleotide donor templates (ssODNs) used in HDR studies.

EMX1 sense (SEQ ID NO: 580)

TCATCTGTGCCCTCCCTCCCTGGCCCAGGTGAAGGTGTGGTTCCAGAACCGGAGGACAAAGTACA
AACGGCAGAAGCTGGAGGAGGAAGGGCCTGAGTTTGAGCAGAAGAAGAAGGGCTCCCATCACATC
AACCGGTGGCGCATTGCCACGAAGCAGGCCAATGGGGAGGACATCGATGTCACCTCCAATGACTAG
GGT

EMX1 antisense (SEQ ID NO: 581)

ACCCTAGTCATTGGAGGTGACATCGATGTCTCTCCCATTTGGCCTGCTTCGTGGCAATGCGCCACCG
GTTGATGTGATGGGAGCCCTTCTTCTTCTGCTCAAACCTCAGGCCCTTCTCCTCCAGCTTCTGCCGT
TTGTACTTTGTCTCCGTTCTGGAACACACCTTCACTGCGCCAGGGAGGGAGGGGCACAGATG
A

HEK293 site 3 sense (SEQ ID NO: 582)

CATGCAATTAGTCTATTTCTGCTGCAAGTAAGCATGCATTTGTAGGCTTGATGCTTTTTTCTGCTTCT
CCAGCCCTGGCCTGGGTCAATCCTTGGGGCTTAGACTGAGCACGTGATGGCAGAGGAAAGGAAGC
CCTGCTTCTCCAGAGGGCGTCCGACAGGACAGCTTTTCTAGACAGGGGCTAGTATGTGCAGCTCCT

HEK293 site 3 antisense (SEQ ID NO: 583)

AGGAGCTGCACATACTAGCCCCTGTCTAGGAAAAGCTGTCTGCGACGCCCTCTGGAGGAAGCAGG
GCTTCTTTCTCTGCCATCACGTGCTCAGTCTAAGCCCCAAGGATTGACCCAGGCCAGGGCTGGA
GAAGCAGAAAAAAGCATCAAGCCTACAAATGCATGCTTACTTGCAGCAGAAATAGACTAATTGCATG

HEK site 4 sense (SEQ ID NO: 584)

GGCTGACAAAGGCCGGGCTGGGTGGAAGGAAGGGAGGAAGGGCCGAGGCAGAGGGTCCAAAGCAG
 GATGACAGGCAGGGGCACCGCGGCCCGGTGGCATTGCGGCTGGAGGTGGGGTTAAAGCGG
 AGACTCTGGTGCTGTGTGACTACAGTGGGGGCCCTGCCCTCTCTGAGCCCCGCCCTCCAGGCCTGT
 GTGTGT

HEK site 4 antisense (SEQ ID NO: 585)

ACACACACAGGCCTGGAGGCGGGGGCTCAGAGAGGGCAGGGCCCCACTGTAGTCACACAGCACC
 AGAGTCTCCGCTTTAACCCCCACCTCCAGCCGCAATGCCACCGGGGCGCCGCGGTGCCCTGCCT
 GTCATCCTGCTTTGACCCTCTGCCTCGCCCTTCTCCCTTCCCTTCCACCCAGCCCGGCCTTTGTCA
 GCC

APOE4 sense (SEQ ID NO: 743)

AGCACCGAGGAGCTGCGGGTGGCCCTCGCCTCCCACCTGCGCAAGCTGCGTAAGCGGCTCCTCCG
 CGATGCCGATGACCTGCAGAAGTGCTGGCAGTGTACCAGGCCGGGGCCCGCAGGGCGCCGAG
 CGCGGCCTCAGCGCCATCCGCGAGCGCCTGGGGCCCTGGTGGAACAGGGCCGCGTGCGGGCCG
 CCACTGT

APOE4 antisense (SEQ ID NO: 744)

ACAGTGGCGGCCCGCACGCGGCCCTGTTCCACCAGGGGCCCCAGGCGCTCGCGGATGGCGCTGA
 GGCCGCGCTCGGCGCCCTCGCGGGCCCGGCCTGGTACACTGCCAGGCACCTTCTGCAGGTCATCG
 GCATCGCGGAGGAGCCGCTTACGCAGCTTGCGCAGGTGGGAGGCGAGGGCGACCCGCAGCTCCT
 CGGTGCT

p53 Y163C sense (SEQ ID NO: 745)

ACTCCCCTGCCCTCAACAAGATGTTTTGCCAACTGGCCAAGACCTGCCCTGTGCAGCTGTGGGTTGA
 TTCCACACCCCCGCCCGGCACCCGCGTCCGCGCCATGGCCATCTACAAGCAGTCACAGCACATGAC
 GGAGGTTGTGAGGCGCTGCCCCACCATGAGCGCTGCTCAGATAGCGATGGTGAGCAGCTGGGGC
 TG

p53 Y163C antisense (SEQ ID NO: 746)

CAGCCCCAGCTGCTCACCATCGCTATCTGAGCAGCGCTCATGGTGGGGGCAGCGCCTCACAACTC
 CGTCATGTGCTGTGACTGCTTGTAGATGGCCATGGCGCGGACGCGGGTGCCGGGCGGGGGTGTGG
 AATCAACCCACAGCTGCACAGGGCAGGTCTTGGCCAGTTGGCAAACATCTTGTGAGGGCAGGGG
 AGT

[00342] Deaminase gene gBlocks Gene Fragments

hAID (SEQ ID NO: 586)

CATCCTTGGTACCGAGCTCGGATCCAGCCACCATGGATAGCCTCTTGATGAATAGACGCCAAGTTCT
 GTATCAGTTTAAAACGCTGAGATGGGCAAAGGCCGACGAGAGACATATCTGTGCTATGTCGTTAAG
 CGCAGAGATTACAGCCACCAGTTTCTCTCTCGACTTCCGGCTACCTGCGGAACAAGAATGGTTGCCATG
 TTGAGCTCCTGTTCTGAGGTATATCAGCGACTGGGATTTGGACCCAGGGCGGTGCTATAGGGTGA
 CATGGTTTACCTCCTGGTCACCTTGTATGACTGCGCGCGGCATGTTGCCGATTTTCTGAGAGGGAA
 CCTAACCTGTCTCTGAGGATCTTACCGCGCGACTGTACTTCTGTGAGGACCGGAAAGCCGAACC
 CGAGGGACTGAGACGCCTCCACAGAGCGGGTGTGCAGATTGCCATAATGACCTTTAAGGACTACTT
 CTACTGCTGGAACACCTTCGTGCGAAAATCACGAGCGGACTTTCAAGGCTTGGGAAGGATTGCATGAA
 AACAGCGTCAGGCTTTCCAGGCAGCTTCCCGCATTCTTCTCCCGTTGTACGAGGTTGATGACCTCA
 GAGATGCCTTTAGAACACTGGGACTGTAGGCGGCCGCTCGATTGGTTTGGTGTGGCTCTAA

rAPOBEC1 (mammalian)(SEQ ID NO: 587)

CATCCTTGGTACCGAGCTCGGATCCAGCCACCATGAGCTCAGAGACTGGCCCAGTGGCTGTGGACC
 CCACATTGAGACGGCGGATCGAGCCCATGAGTTTGAGGTATTCTTCGATCCGAGAGAGCTCOGCA
 AGGAGACCTGCCTGCTTTACGAAATTAATTGGGGGGGCCGACTCCATTTGGCGACATACATCACA
 GAACACTAACAAGCACGTCGAAGTCAACTTCATCGAGAAGTTCACGACAGAAAGATATTTCTGTCCG
 AACACAAGGTGCAGCATTACCTGGTTTCTCAGCTGGAGCCCATGCGGCGAATGTAGTAGGGCCATC
 ACTGAATTCCTGTCAAGGTATCCCCACGTCACTCTGTTTATTTACATCGCAAGGCTGTACCACCACGC
 TGACCCCGCAATCGACAAGGCCTGCGGGATTTGATCTCTTCAGGTGTGACTATCCAAATTATGACT
 GAGCAGGAGTCAGGATACTGCTGGAGAACTTTGTGAATTATAGCCCGAGTAATGAAGCCCACTGG
 CCTAGGTATCCCCATCTGTGGGTACGACTGTACGTTCTTGAAGTGTACTGCATCATACTGGGCCTGC
 CTCCTTGTCTCAACATTCTGAGAAGGAAGCAGCCACAGCTGACATTCTTTACCATCGCTCTTCAGTCT
 TGTCAATACCAGCGACTGCCCCACACATTCTCTGGGCCACCGGTTGAAATGAGCGGCCGCTCGA
 TTGGTTTGGTGTGGCTCTAA

pmCDA1 (SEQ ID NO: 588)

CATCCTTGGTACCGAGCTCGGATCCAGCCACCATGACAGACGCTGAATATGTTAGGATCCATGAAAA
 ACTGGATATCTATACATTTAAGAAGCAGTTCTTCAATAACAAAAAGTCAGTATCTCACAGATGCTATGT
 CCTGTTTCAACTCAAGAGAAGAGGAGAAAGGCGGGCCTGTTTCTGGGGGTACGCGGTTAATAAACC
 CCAGTCCGGGACCGAGAGGGGGATTACGCGCCAGATCTTTTCAATTAGGAAGGTTGAAGAGTATCT
 TCGCGACAATCCCGGTCAGTTCACAATTAAGTGGTACAGCTCCTGGAGCCCTTGCCTGATTGCGCC
 GAGAAAATACTCGAATGGTACAACCAGGAGTTGAGAGGCAATGGCCACACTCTCAAGATTTGGGCTT
 GCAAGCTTTACTACGAGAAGAACGCGAGAAATCAGATTGGCTTGTGGAACCTCAGGGACAACGGGG
 TCGGGTTGAATGTTATGGTGTCCGAACATTACCAGTGTGTAGAAAGATCTTCATTAGTCCAGTCA
 AATCAGCTGAACGAGAACAGATGGCTGGAGAAAACACTGAAACGGGCAGAGAAAAGGCGCTCAGAG
 CTGAGTATCATGATCCAGGTCAAAATCCTGCATACAACCAAAGGCCCGGCTGTATAAGCGGCCGCTC
 GATTGGTTTGGTGTGGCTCTAA

haPOBEC3G (SEQ ID NO: 589)

CATCCTTGGTACCGAGCTCGGATCCAGCCACCATGGAGCTGAAGTATCACCCCTGAGATGCGGTTTTT
 CCACTGGTTTTAGTAAAGTGGCGCAAACCTTCATCGGGATCAGGAGTATGAAGTGACCTGGTATATCTCT
 TGGTCTCCCTGCACAAAATGTACACGCGACATGGCCACATTTCTGGCCGAGGATCCAAAAGGTGACG
 CTCACAATCTTTGTGGCCCGCCTGTATTATTTCTGGGACCCGGATTATCAGGAGGCACCTTAGGTCAT
 TGTGCCAAAAGCGCGACGGACCAGGGCGACTATGAAAATCATGAATTATGACGAATTCAGCATTG
 CTGGAGTAAGTTTGTGTACAGCCAGCGGGAGCTGTTGAGCCCTGGAACAATCTTCCCAAGTACTAC
 ATACTGCTTCACATTATGTTGGGGGAGATCCTTCGGCACTCTATGGATCCTCCTACCTTTACGTTTAA
 CTTTAATAATGAGCCTTGGGTTTCGCGGGCGCCATGAAACCTATTTGTGCTACGAGGTCGAGCGGATG
 CATAATGATACGTGGGTCTGCTGAATCAGAGGAGGGGGTTTCTGTGTAACCAGGCTCCACATAAAC
 ATGGATTTCTCGAGGGGGCGGCACGCCGAACTGTGTTTCTTGATGTGATACCTTTCTGGAAGCTCGA
 CCTTGATCAAGATTACAGGGTGACGTGTTTACCTCCTGGTCACCCCTGCTTCAGTTGCGCCCAAGAG
 ATGGCTAAATTTATCAGTAAGAACAAGCATGTGTCCCTCTGTATTTTACAGCCAGAATTTATGATGAC
 CAGGGCCGGTGCCAGGAGGGGCTGCGGACACTCGCTGAGGCGGGCGCGAAGATCAGCATAATGA
 CATACTCGAATTCAAACACTGTTGGGACACTTTTGTGGACCACAGGGCTGCCATTTACGCCGTG
 GGATGGGCTCGACGAACATAGTCAGGATCTCTCAGGCCGGCTGCGAGCCATATTGCAGAACCAGGA
 GAATTAGCGGCCGCTCGATTGGTTTGGTGTGGCTCTAA

rAPOBEC1(*E. Coli*) (SEQ ID NO: 590)

GGCCGGGGATTCTAGAAATAATTTTGTAACTTTAAGAAGGAGATATACCATGGATGTCTTCTGAAA
 CCGGTCCGGTTGCGGTTGACCCGACCCCTGCGTCGTCGATCGAACCAGCAGCAATTCGAAGTTTTCT
 TCGACCGCGTGAAC|GCGTAAAGAAACCTGCCTGCTGTACGAAATCAACTGGGGTGGTCTGACT
 CTATCTGGCGTCACACCTCTCAGAACACCAACAACACGTTGAAGTTAACTTCATCGAAAAATTCACC
 ACCGAACGTTACTTCTGCCCGAACACCCGTTGCTCTATCACCTGGTTCCCTGTCTTGGTCTCCGTGCG
 GTGAATGCTCTCGTGCATCACCGAATTCCTGTCTCGTTACCCGACGTTACCCCTGTTTACTTACAT
 GCGCGTCTGTACCACCACGCGGACCCGCGTAACCGTCAGGGTCTGCGTGACCTGATCTCTTCTGGT
 GTTACCATCCAGATCATGACCGAACAGGAATCTGGTACTGCTGGCGTAACTTCGTTAACTACTCTCC
 GTCTAACGAAGCGCACTGGCCGCGTTACCCGACCTGTGGGTTTCGTCTGTACGTTCTGGAAGTGA
 CTGCATCATCCTGGGTCTGCCGCGGTGCCTGAACATCCTGCGTGTAAACAGCCGACGCTGACCTT
 CTTACCATCGCGCTGCAGTCTTGCCACTACCAGCGTCTGCCGCGCACATCCTGTGGGCGACCGG
 TCTGAAAGGTGGTAGTGGAGGGAGCGGCGGTTCAATGGATAAGAAATAC

[00343] Amino Acid Sequences of NBE1, NBE2, and NBE3.

NBE1 for *E. Coli* expression (His₆-rAPOBEC1-XTEN-dCas9) (SEQ ID NO: 591)

MGSSHHHHHHMSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKTCCLYEINWGGRRHSIWRHTSQNTN
KHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYHHADPRNRQGL
RDLISSGVTIQIMTEQESGYCWRNFVNYSNPSNEAHWPYPHLLWVRLYVLELYCIIILGLPPCLNLRKQPQ
LTFFTIALQSCHYQRLPPHILWATGLKSGSETPGTSESATPESDKKYSIGLAIGTNSVGVAVITDEYKVPK
KFKVLGNTDRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRL
EESFLVEEDKKHERHPHIFGNIVDEVAYHEKYPTIYHLRKKLV DSTDKADLR LIYLALAHMIKFRGHFLIEGDL
NPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRRENLIQAQLPGEKKNGLFGNLIASL
GLTPNFKSNFDLAEDAKLQLSKD TYDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSA
SMIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELL
VKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYVVGPLARGNSRF
AWMTRKSEETITPWNFEVV DKGASAQSFIERMTNFDKNLPNEKVLPHKSLLYEYFTVYNELTKVKYVTE
GMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIKDK
DFLDNEENEDILEDIVLTLTLFEDREMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQS
GKTILDFLKSDGFANRNFQMQLIHDDSLTFKEDIQKAQVSGQDLSHEHIANLAGSPAIKKGILQTVKVVDEL
VKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGKELGSQILKEHPVENTQLONEKLYLYLQ
NGRDMYVDQELDINRLSDYDVAIVPQSFLKDDSIDNKVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQL
LNAKLITQRKFDNLTKAERGGLSELDKAGFIKRQLVETROITKHVAQILDSRMNTKYDENDKLIREVKVITLK
SKLVSDFRKDFQFYKVRINNYHHAHDAYLNAVVGTA LIKKYPKLESEFVYGDYKVDVRKMIKSEQEIG
KATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVL SMPQVNIVKKTEVQT
GGFSKESILPKRNSDKLIARKKDWDPKKGFDSPVAYSVLVAKVEKKGSKKLKSVKELLGITIMERS
FEKNPIDFLEAKGYKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFY LASHYEKL
KGSPEDNEQKQLFVEQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIIHLFTLTNL
GAPAAFKYFDTTIDRKRYTSTKEVL DATLIHQSI TGLYETRIDLSQLGGDSGGSPKKKRKV

NBE1 for Mammalian expression (rAPOBEC1-XTEN-dCas9-NLS) (SEQ ID NO: 592)

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKTCCLYEINWGGRRHSIWRHTSQNTNKHVEVNFIEK
FTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYHHADPRNRQGLRDLISSGVTIQ
IMTEQESGYCWRNFVNYSNPSNEAHWPYPHLLWVRLYVLELYCIIILGLPPCLNLRKQPQLTFFFTIALQSC
HYQRLPPHILWATGLKSGSETPGTSESATPESDKKYSIGLAIGTNSVGVAVITDEYKVPKSKFKVLGNTDR
HSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSSFFHRL EESFLVEEDKK
HERHPHIFGNIVDEVAYHEKYPTIYHLRKKLV DSTDKADLR LIYLALAHMIKFRGHFLIEGDLNPDNSDVKLFI
QLVQTYNQLFEENPINASGVDAKAILSARLSKSRRENLIQAQLPGEKKNGLFGNLIASLGLTPNFKSNFDL
AEDAKLQLSKD TYDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQ
DLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQ
RTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYVVGPLARGNSRF AWMTRKSEETITP
WNFEVV DKGASAQSFIERMTNFDKNLPNEKVLPHKSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQ
KKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIKDKDFLDNEENEDILE
DIVLTLTLFEDREMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFA
NRNFMQLIHDDSLTFKEDIQKAQVSGQDLSHEHIANLAGSPAIKKGILQTVKVVDELVKVMGRHKPENIVI
EMARENQTTQKGQKNSRERMKRIEEGKELGSQILKEHPVENTQLONEKLYLYLQNGRDMYVDQELDI
NRLSDYDVAIVPQSFLKDDSIDNKVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNL
TKAERGGLSELDKAGFIKRQLVETROITKHVAQILDSRMNTKYDENDKLIREVKVITLKS KLVSDFRKDFQF
YKVRINNYHHAHDAYLNAVVGTA LIKKYPKLESEFVYGDYKVDVRKMIKSEQEIGKATAKYFFYSNIM
NFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVL SMPQVNIVKKTEVQTGGFSKESILPKRN
SDKLIARKKDWDPKKGFDSPVAYSVLVAKVEKKGSKKLKSVKELLGITIMERSSEKKNPIDFLEAKG
YKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFY LASHYEKLKGSPEDNEQKQL
FVEQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIIHLFTLTNLGAPAAFKYFDTTI
DRKRYTSTKEVL DATLIHQSI TGLYETRIDLSQLGGDSGGSPKKKRKV

Alternative NBE1 for Mammalian expression with human APOBEC1 (hAPOBEC1-XTEN-dCas9-NLS) (SEQ ID NO: 5737)

MTSEKGPSTGDPTLRRRIEPWEFDVFYDPRELKREACLLYEIKWGMSRKIWRSSGKNTTN
HVEVNFIIKFTSERDFHPSMCSITWFLSWPCWECSQAIREFLSRHPGVTLVIYVARLFW
HMDQQNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMMLY
ALELHCILSLPPCLKISRRWQNHLTFFRLHLQNCHYQTIPPHILLATGLIHPSVAWRGSETP
GTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIGALL
FDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRLEESFLVEEDKK
HERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGD
LNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGEKK
NGLFGNLIASLGLTPNFKSNFDLAEDAQLSKDQYDDDLNLLAQIGDQYADFLAA
KNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQ
SKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTFDNGSIPHQIH
LGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSRF AWMTRKSEETITPW
NFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPKHSLLYEYFTVYNELTKVKYVTEGMR
KPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYH
DLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERLKYAHLFDDKVMKQLKRRRY
TGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQ
GDSLHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGRHHPENIVIAMARENQTTQKGQK
NSRERMKRIEIEGKELGSQILKEHPVENTQLQNEKLYLYLQNGRDMYVDQELDINRLSD
YDVDAIVPQSFLKDDSIDNKVLTRSDKNRGKSDNVPSEEVVKMKMKNYWRQLLNAKLITQ
RKFDNLTKAERGGLSELDKAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVK
VITLKSCLVSDFRKDFQFYKVREINNYHHAHDAYLNAVVGTAIIKYPKLESEFVYGDY
KVYDVRKMIKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIWW
DKGRDFATVRKVLSPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGG
FDSPTVAYSVLVVAKVEKGKSKLKSVKELLGITIMERSSEKPNIDFLEAKGYKEVKKD
LIIKLPKYSLEENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGKSPEDNE
QKQLFVEQHKHYLDEIIEQISEFSKR VILADANLDKVL SAYNKHDKPIREQAENIIHLFTL
TNLGAPAAFKYFDTTIDRKRYTSTKEVLDTLHQHSITGLYETRIDLSQLGGDSGGSPKKK
RKV

NBE2 (rAPOBEC1-XTEN-dCas9-UGI-NLS) (SEQ ID NO: 593)

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRRHSIWRHTSQNTNKHVEVNFIEKF
 TTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYHHADPRNRQGLRDLISSGVTIQ
 IMTEQESGYCWRNFVNYSNPSNEAHWPYPHLLWVRLYVLELYCIILGLPCLNLRKQPOLTFFTIALQSC
 HYQRLPPHILWATGLKSGSETPGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDR
 HSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRLSEESFLVEEDKK
 HERHPIFGNIVDEVAYHEKYPTIYHLRKKLVSDTKADLRILIYLAHAHMIKFRGHFLIEGDLNPDNSDVKLF
 IQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDL
 AEDAKLQLSKDQYDDDLNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQ
 DLTLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQ
 RTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTRIPYVVGPLARGNSRFAMWTRKSEETITP
 WNFEEVVDKGASQAQSFIERMTNFDKNLPNEKVLPHKSHLLYEYFTVYNELTKVKYVTEGMRKPAFLSSEQ
 KKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILE
 DIVLTLTLFEDREMIERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDFGFA
 NRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGILQTVKVVDELVKVMGRHKPENIVI
 EMARENQTTQKGGQNSRERMKRIEIGIKELGSQILKEHPVENTQLQNEKLYLYLQNGRDMYVDQELDI
 NRLSDYDVAIVPQSFLKDDSIDNKVLRSDKNRKGSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNL
 TKAERGGLSELDKAGFIKRQLVETRQITKHVAQILD SRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQF
 YKVRINNYHHAHDAYLNAVVGTAALIKKYPKLESEFVYGDYKVDVRKMIKSEQEI GKATAKYFFYSNIM
 NFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVL SMPQVNIKKTEVQTGGFSKESILPKRN
 SDKLIARKKDWDPKKYGGFDSPTVAYSVLVAKVEKGKSKLKS VKELLGITIMERSSFEKNPIDFLEAKG
 YKEVKKDLIILPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQL
 FVEQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIIHLFTLNLGAPAAFKYFDTTI
 DRKRYTSTKEVL DATLIHQSI TGLYETRIDLSQLGGDSGGSTNLSDIIEKETGKQLVIQESILMLPEEVVEEVIG
 NKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSPKKKRKY

NBE3 (rAPOBEC1-XTEN-Cas9n-UGI-NLS) (SEQ ID NO: 594)

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKRETCCLLYEINWGGRRHSIWRHTSQNTNKHVEVNFIEKF
 TTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYHHADPRNRQGLRDLISSGVTIQ
 IMTEQESGYCWRNFVNYSNPSNEAHWPYPHLLWVRLYVLELYCIILGLPCLNLRKQPOLTFFTIALQSC
 HYQRLPPHILWATGLKSGSETPGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPSSKFKVLGNTDR
 HSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRLSEESFLVEEDKK
 HERHPIFGNIVDEVAYHEKYPTIYHLRKKLVSDTKADLRILIYLAHAHMIKFRGHFLIEGDLNPDNSDVKLF
 IQLVQTYNQLFEENPINASGVDAKAILSARLSKSRLENLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDL
 AEDAKLQLSKDQYDDDLNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQ
 DLTLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQ
 RTFDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTRIPYVVGPLARGNSRFAMWTRKSEETITP
 WNFEEVVDKGASQAQSFIERMTNFDKNLPNEKVLPHKSHLLYEYFTVYNELTKVKYVTEGMRKPAFLSSEQ
 KKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILE
 DIVLTLTLFEDREMIERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDFGFA
 NRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGILQTVKVVDELVKVMGRHKPENIVI
 EMARENQTTQKGGQNSRERMKRIEIGIKELGSQILKEHPVENTQLQNEKLYLYLQNGRDMYVDQELDI
 NRLSDYDVAIVPQSFLKDDSIDNKVLRSDKNRKGSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNL
 TKAERGGLSELDKAGFIKRQLVETRQITKHVAQILD SRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQF
 YKVRINNYHHAHDAYLNAVVGTAALIKKYPKLESEFVYGDYKVDVRKMIKSEQEI GKATAKYFFYSNIM
 NFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVL SMPQVNIKKTEVQTGGFSKESILPKRN
 SDKLIARKKDWDPKKYGGFDSPTVAYSVLVAKVEKGKSKLKS VKELLGITIMERSSFEKNPIDFLEAKG
 YKEVKKDLIILPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQL
 FVEQHKHYLDEIIEQISEFSKRVLADANLDKVL SAYNKHRDKPIREQAENIIHLFTLNLGAPAAFKYFDTTI
 DRKRYTSTKEVL DATLIHQSI TGLYETRIDLSQLGGDSGGSTNLSDIIEKETGKQLVIQESILMLPEEVVEEVIG
 NKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSPKKKRKY

pmCDA1-XTEN-dCas9-UGI (bacteria) (SEQ ID NO: 5742)

MTDAEYVRIHEKLDIYTFKKQFFNKKSVSHRCYVLFELKRRGERRACFWGYAVNKPQS
 GTERGIIHAEIFSIRKVEEYLRDNPQGFTINWYSSWSPCADCAEKILEWYNQELRGNHTL
 KIWACKLYYEKNARNQIGLWNLRDNGVGLNVMVSEHYQCCRKIFIQSSHNQLNENRWL

EKTLKRAEKRRSELSIMIQVKILHTTKSPAVSGSETPGTSESATPESDKKYSIGLAIGTNSV
 GWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRK
 NRICYLQEIFSNEMAKVDDSFHRLSEESFLVEEDKKHERHPIFGNIVDEVAHYEKYPTIYH
 LRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDLNPDNSDVKLFIQLVQTYNQLFE
 ENPINASGVDAKAILSARLSKSRLENLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDLA
 EDAKLQLSKDQYDDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSAS
 MIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILE
 KMDGTEELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFKDNREKIEK
 ILTRIPYYYVGPLARGNSRFAWMTRKSEETITPWNFEEVVDKGASQSFIERMTNFDKNLP
 NEKVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVK
 QLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLT
 LFEDREMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLK
 SDGFANRNFQMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIAKKGILQTVKVV
 DELVKVMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIIEGKELGSQILKEHPVENT
 QLQNEKLYLYLQNGRDMYVDQELDINRLSDYDVAIVPQSFLKDDSIDNKVLTRSDKN
 RGKSDNVPSEEVVKMKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDKAGFIKRQL
 VETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKREINNY
 HHAHDAYLNAVVGTAIIKYPKLESEFVYGDYKVVYDVRKMIKSEQEIGKATAKYFFYS
 NIMNFFKTEITLANGEIRKRLIETNGETGEIVWDKGRDFATVRKVL SMPQVNIVKKTEVQ
 TGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVVAKVEKGKSKKLSV
 KELLGITIMERSSEKNPIDFLEAKGYKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQ
 KGNELALPSKYVNFLYLASHYEKLGKSPEDNEQKQLFVEQHKHYLDEIIEQISEFSKRVIL
 ADANLKVLSAYNKHDKPIREQAENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEV
 LDATLIHQSGITGLYETRIDLSQLGGDSGGSMTNLSDIIEKETGKQLVIQESILMLPEEVEEVI
 GNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKML

pmCDA1-XTEN-nCas9-UGI-NLS (mammalian construct) (SEQ ID NO: 5743)

MTDAEYVRIHEKLDIYTFKKQFFNKKSVSHRCYVLFELKRRGERRACFWGYAVNKPQS
 GTERGIHAEIFSIRKVEEYLRDNPQGQFTINWYSSWSPCADCAEKILEWYNQELRGNHTL
 KIWACKLYYEKNARNQIGLWNLRDNGVGLNVMVSEHYQCCRKIFIQSSHNQLNENRWL
 EKTLKRAEKRRSELSIMIQVKILHTTKSPAVSGSETPGTSESATPESDKKYSIGLAIGTNSV
 GWAVITDEYKVPSSKFKVLGNTDRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRK
 NRICYLQEIFSNEMAKVDDSFHRLSEESFLVEEDKKHERHPIFGNIVDEVAHYEKYPTIYH
 LRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDLNPDNSDVKLFIQLVQTYNQLFE
 ENPINASGVDAKAILSARLSKSRLENLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDLA
 EDAKLQLSKDQYDDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSAS
 MIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILE
 KMDGTEELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFKDNREKIEK
 ILTRIPYYYVGPLARGNSRFAWMTRKSEETITPWNFEEVVDKGASQSFIERMTNFDKNLP
 NEKVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVK
 QLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLT
 LFEDREMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLK
 SDGFANRNFQMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIAKKGILQTVKVV
 DELVKVMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIIEGKELGSQILKEHPVENT
 QLQNEKLYLYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDKN
 RGKSDNVPSEEVVKMKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDKAGFIKRQL

VETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSKLVSDFRKDFQFYK VREINNY
 HHAHDAYLNAVVG TALIKKYPKLESEFVYGDYKVYDVRKMIAKSEQEIGKATAKYFFYS
 NIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVL SMPQVNIVKKTEVQ
 TGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVAYSVLVVAKVEKGKSKKLSV
 KELLGITIMERSSEFEKNPIDFLEAKGYKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQ
 KGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQHKKHYLDEIIEQISEFSKRVIL
 ADANLDKVL SAYNKH RDKPIREQAENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEV
 LDATLIHQ SITGLYETRIDLSQLGGDSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIG
 NKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSPKKKR
 KV

huAPOBEC3G-XTEN-dCas9-UGI (bacteria) (SEQ ID NO: 5744)

MDPPTFTFNFNNEPWVRGRHETYLCYE VERMHNDTWVLLNQRRGFLCNQAPHKHGFLE
 GRHAELCFLDVIPFWKLDLDQDYRVTCFTSWSPCFSCAQEMAKFISK NKHVSLCIFTARIY
 DDQGRQCQEGLRTLAEAGAKISIMTYSEFKHCWDTFVDHQGCPFQPWDGLDEHSQDLSGR
 LRAILQSGSETPGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVP SKKFKVLGNTDR
 HSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDD SFFHRL
 EESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLR LIYLALAHMI
 KFRGHFLIEGDLNPDNSVDKLFQILVQTYNQLFEENPINASGVDAKAILSARLSKSRRL E
 NLIAQLPGEKKNGLFGNLI ALSGLTPNFKSNFDLAEDAQLQSKD TYDDDLDNLLAQIG
 DQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQL
 PEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRT
 FDNGSIPHQIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSRFAWMT
 RKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVL PKHSLLEYEYFTVYNELTK
 VKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDR
 FNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERLKT YAHLFDDKV
 MKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRNF MQLIHDDSLTFKED
 IQKAQVSGQGDSLHEHIANLAGSPA IKKGILQTVKVVDELVKVMGRHKPENIV IEMAREN
 QTTQKGQKNSRERMKRIEEGIKELGSQILKEHPVENTQLQNEKLYLYYLQNGRDMYVDQ
 ELDINRLSDYDVAIVPQSFLKDDSIDNKVLTRSDKNRGKSDNVPSEE VVKMKKNYWRQ
 LLNAKLITQRKFDNLTKAERGG LSELDKAGFIKRQLVETRQITKHVAQILDSRMNTKYDE
 NDKLIREVKVITLKSKLVSDFRKDFQFYK VREINNYHHAHDAYLNAVVG TALIKKYPKLE
 SEFVYGDYKVYDVRKMIAKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETN
 GETGEIVWDKGRDFATVRKVL SMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKD
 WDPKKYGGFDSPTVAYSVLVVAKVEKGKSKKLSVKELLGITIMERSSEFEKNPIDFLEAK
 GYKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEK L
 KGSPEDNEQKQLFVEQHKKHYLDEIIEQISEFSKRVILADANLDKVL SAYNKH RDKPIREQA
 ENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQ SITGLYETRIDLSQLGGD
 SGGSMTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLL
 TSDAPEYKPWALVIQDSNGENKIKML

huAPOBEC3G-XTEN-nCas9-UGI-NLS (mammalian construct) (SEQ ID NO: 5745)

MDPPTFTFNFNNEPWVRGRHETYLCYE VERMHNDTWVLLNQRRGFLCNQAPHKHGFLE
 GRHAELCFLDVIPFWKLDLDQDYRVTCFTSWSPCFSCAQEMAKFISK NKHVSLCIFTARIY
 DDQGRQCQEGLRTLAEAGAKISIMTYSEFKHCWDTFVDHQGCPFQPWDGLDEHSQDLSGR
 LRAILQSGSETPGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVP SKKFKVLGNTDR

HSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRL
 EESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMI
 KFRGHFLIEGDLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRRL
 NLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDLAEDAQLSKDQYADLFLAAKNLS
 DAILSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQSK
 NGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTFDNGSIPH
 QIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSRFAWMT
 RKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPHKSHLLYEYFTVYNEL
 TKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISG
 VEDRFNASLGTYHDLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERLKTYAHL
 FDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRNFMQLIHDDSL
 TFKEDIQKAQVSGQGDSLHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGRHKPENIV
 IEMARENQTTQKGQKNSRERMKRIEEGIKELGSQILKEHPVENTQLQNEKLYLYYLQ
 NGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDKNRGKSDNVPSEE
 VVKMKKNYWRQLLNAKLITQRKFDNLTKAERGGELSELDKAGFIKRQLVETRQITKHVA
 QILDSRMNTKYDENDKLIREVKVITLKSLLVSDFRKDFQFYK VREINNYHHAHDAYLNA
 VVG TALIKKYPKLESEFVYGDYKVYDVRKMIKSEQEIGKATAKYFFYSNIMNFFKTEIT
 LANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVL SMPQVNIVKKTEVQTGGFSKESIL
 PKRNSDKLIARKKD WDPKKYGGFDSPTVAYSVLVVAKVEK GKSKKLSVKELLGITIMER
 SFEKNPIDFLEAKGYKEVKKDLI IKLPKYSLFELENGRKRMLASAGELQKGNELALPSK
 YVNFLYLASHYEKLGSPEDNEQKQLFVEQH KHYLDEIIEQISEFSKR VILADANLDKVL
 SAYNKHRDKPIREQAENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVL DATLIHQ
 SITGLYETRIDLSQLGGD SGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPE
 SDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSPKKKRKV

huAPOBEC3G (D316R_D317R)-XTEN-nCas9-UGI-NLS (mammalian construct) (SEQ ID NO:
 5746)

MDPPTFTFNFNNEPWVRGRHETYLCYE VERMHNDTWVLLNQR RGFLCNQAPHKHGFLE
 GRHAELCFLDVIPFWKLDLDQDYRVTCFTSWSPCFSCAQEMAKFISK NKHVS LCIFTARIY
 RRQGRQCQEGLR TLAEAGAKISIMTYSEFKHCWDTFVDHQGCPFPQWDGLDEHSQDLSGR
 LRAILQSGSETPGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVP SKKFKVLGNTDR
 HSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRL
 EESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMI
 KFRGHFLIEGDLNPDNSDVKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRRL
 NLIAQLPGEKKNGLFGNLIASLGLTPNFKSNFDLAEDAQLSKDQYADLFLAAKNLS
 DAILSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQSK
 NGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTFDNGSIPH
 QIHLGELHAILRRQEDFYFPLKDNREKIEKILTFRIPYYVGPLARGNSRFAWMT
 RKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNEKVLPHKSHLLYEYFTVYNEL
 TKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISG
 VEDRFNASLGTYHDLKIIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERLKTYAHL
 FDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRNFMQLIHDDSL
 TFKEDIQKAQVSGQGDSLHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGRHKPENIV
 IEMARENQTTQKGQKNSRERMKRIEEGIKELGSQILKEHPVENTQLQNEKLYLYYLQ
 NGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDKNRGKSDNVPSEE
 VVKMKKNYWRQLLNAKLITQRKFDNLTKAERGGELSELDKAGFIKRQLVETRQITKHVA
 QILDSRMNTKYDE

NDKLIREVKVITLKSKLVSDFRKDFQFYK VREINNYHHAHDAYLNAV VGTALIKKYPKLE
 SEFVYGDYKVYDVRKMIKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETN
 GETGEIVWDKGRDFATVRKVL SMPQVNIVKKTEVQTGGF SKESILPKRNSDKLIARKKD
 WDPK KYGGFDSPTVAYSVLVVAKVEKGKSKKLKSVKELLGITIMERS SFEKNPIDFLEAK
 GYKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFY LASHYEKL
 KGSPEDNEQKQLFVEQHKHYLDEIIEQISEFSKRVILADANLDK VLSAYNKHRDKPIREQA
 ENIIHLFTLTNLGAPAAF KYFDTTIDRKRYTSTKEVLDATLIHQ SITGLYETRIDLSQLGGD
 SGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLT
 SDAPEYKPWALVIQDSNGENKIKMLSGGSPKKKRKV

[00344] Base Calling Matlab Script

```

WTnuc='GCGGACATGGAGGACGTGCGCGGCCGCCTGGTGCAGTACCGCGGCGAGGTGCAGGCCATGCTCGGC
CAGA
GCACCGAGGAGCTGCGGGTGCGCCTCGCCTCCACCTGCGCAAGCTGCGTAAGCGGCTCCTCCGCGATGCCG
ATGAC
CTGCAGAAGCGCCTGGCAGTGTACCAGGCCGGGCCCCGCGAGGGCGCCGAGCGCGGCTCAGCGCCATCCGC
GAGCG CCTGGGGCCCCCTGGTGGAACAG'(SEQ ID NO: 595);
%cycle through fastq files for different samples files=dir('* .fastq');
for d=1:20
filename=files(d).name;
%read fastq file
[header,seqs,qscore]=fastqread(filename);
seqsLength=length(seqs); % number of sequences seqsFile=
strrep(filename,'.fastq',''); % trims off .fastq
%create a directory with the same name as fastq file ifexist(seqsFile,'dir');
error('Directory already exists. Please rename or move it before moving on. ');
end
mkdir(seqsFile); % make directory
wtLength=length(WTnuc); % length of wildtype sequence
%% aligning back to the wildtype nucleotide sequence
%
% ALN is a matrix of the nucleotide alignment window=1:wtLength;
sBLength=length(seqs); % number of sequences
% counts number of skips nSkips = 0;
ALN= repmat('',[sBLength wtLength]);
% iterate through each sequencing read for i = 1:sBLength
%If you only have forward read fastq files leave as is
%If you have R1 forward and R2 reverse fastq files uncomment the
%next four lines of code and the subsequent end statement
% ifmod(d,2)==0;
% reverse=seqrcomplement(seqs{i});
% [score,alignment,start]=
swalign(reverse,WTnuc,'Alphabet','NT');
% else

[score,alignment,start]=swalign(seqs{i},WTnuc,'Alphabet','NT');
% end

% length of the sequencing read len =
length(alignment(3,:));
% if there is a gap in the alignment , skip = 1 and we will
    
```

```
% throw away the entire read skip = 0;
for j = 1:len
if (alignment(3,j) == '-' || alignment(1,j) == '-') skip = 1;
    break;
end
%in addition if the qscore for any given base in the read is
    %below 31 the nucleotide is turned into an N (fastq qscores that are not letters)
if isletter(qscore{i}(start(1)+j-1)) else
alignment(1,j) = 'N';
    end

end
if skip == 0 && len > 10
ALN(i, start(2):(start(2)+length(alignment)-1)) = alignment(1,:);
    end
end
```

```

% with the alignment matrices we can simply tally up the occurrences of
% each nucleotide at each column in the alignment these
% tallies ignore bases annotated as N
% due to low qscores
TallyNTD=zeros(5,wtLength); for i=1:wtLength

TallyNTD(:,i)=[sum(ALN(:,i)=='A'),sum(ALN(:,i)=='C'),sum(ALN(:,i)=='G'),sum(A
LN(:,i)=='T'),sum(ALN(:,i)=='N')];
end
% we then save these tally matrices in the respective folder for
% further processing

save(strcat(seqsFile,'/TallyNTD'),'TallyNTD'); dlmwrite(strcat(seqsFile,'/TallyNTD.txt'),TallyNTD,'precision',
'%.3f','newline','pc'); end

```

[00345] INDEL Detection Matlab Script

```

WTnuc='GCGGACATGGAGGACGTGCGCGGCCGCCTGGTGCAGTACCGCGGCGAGGTGCAGGCCATGCTCGGC
CAGA
GCACCGAGGAGCTGCGGGTGCGCCTCGCCTCCACCTGCGCAAGCTGCGTAAGCGGCTCCTCCGCGATGCCG
ATGAC
CTGCAGAAGCGCCTGGCAGTGTACCAGGCCGGGGCCCGCGAGGGCGCCGAGCGCGGCCCTCAGCGCCATCCGC
GAGCG CCTGGGGCCCCTGGTGG AACAG'(SEQ ID NO: 595);
%cycle through fastq files for different samples files=dir('* .fastq');
%specify start and width of indel window as well as length of each flank indelstart=154;
width=30; flank=10;

for d=1:3
filename=files(d).name;
%read fastq file
[header,seqs,qscore]=fastqread(filename);
seqsLength=length(seqs); % number of sequences seqsFile
=concat(strrep(filename,'.fastq',''),'_INDELS');
%create a directory with the same name as fastq file+_INDELS ifexist(seqsFile,'dir');
error('Directory already exists. Please rename or move it before moving on. ');
end
mkdir(seqsFile); % make directory
wtLength=length(WTnuc); % length of wildtype sequence sBLength=
length(seqs); % number of sequences

% initialize counters and cell arrays
nSkips = 0; notINDEL=0;
ins={};
dels={}; NumIns=0;
NumDels=0;
% iterate through each sequencing read for i = 1:sBLength
%search for 10BP sequences that should flank both sides of the "INDEL WINDOW"
windowstart=strfind(seqs{i},WTnuc(indelstart-flank:indelstart));

windowend=strfind(seqs{i},WTnuc(indelstart+width:indelstart+width+flank
));
%if the flanks are found proceed
if length(windowstart)==1 && length(windowend)==1
%if the sequence length matches the INDEL window length save as

```

```
%not INDEL
if windowend-windowstart==width+flank notINDEL=notINDEL+1;
    %if the sequence is two or more bases longer than the INDEL
    %window length save as an Insertion
elseif windowend-windowstart>=width+flank+2 NumIns=NumIns+1;
    ins{NumIns}=seqs{i};
    %if the sequence is two or more bases shorter than the INDEL
    %window length save as a Deletion
elseif windowend-windowstart<=width+flank-2 NumDels=NumDels+1;
    dels{NumDels}=seqs{i};
    %keep track of skipped sequences that are either one base
    %shorter or longer than the INDEL window width else
    nSkips=nSkips+1;
end
%keep track of skipped sequences that do not possess matching flank
%sequences else
nSkips=nSkips+1;

        end
end
```

```

fid=fopen(strcat(seqsFile,'/summary.txt'),'wt');
fprintf(fid, 'Skipped reads %i\n not INDEL %i\n Insertions %i\n Deletions
%i\n', [nSkips, notINDEL, NumIns, NumDels]); fclose(fid);
save(strcat(seqsFile,'/nSkips'),'nSkips'); save(strcat(seqsFile,'/notINDEL'),'notINDEL');
save(strcat(seqsFile,'/NumIns'),'NumIns'); save(strcat(seqsFile,'/NumDels'),'NumDels');
save(strcat(seqsFile,'/dels'),'dels');
C = dels;
fid = fopen(strcat(seqsFile,'/dels.txt'),'wt'); fprintf(fid, "%s\n", C{:});
fclose(fid);
save(strcat(seqsFile,'/ins'),'ins'); C = ins;
fid = fopen(strcat(seqsFile,'/ins.txt'),'wt'); fprintf(fid, "%s\n", C{:});
fclose(fid);

end

```

EXAMPLE 5: Cas9 variant sequences

[00346] The disclosure provides Cas9 variants, for example Cas9 proteins from one or more organisms, which may comprise one or more mutations (*e.g.*, to generate dCas9 or Cas9 nickase). In some embodiments, one or more of the amino acid residues, identified below by an asterisk, of a Cas9 protein may be mutated. In some embodiments, the D10 and/or H840 residues of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, are mutated. In some embodiments, the D10 residue of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, is mutated to any amino acid residue, except for D. In some embodiments, the D10 residue of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, is mutated to an A. In some embodiments, the H840 residue of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding residue in any of the amino acid sequences provided in SEQ ID NOs: 11-260, is an H. In some embodiments, the H840 residue of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, is mutated to any amino acid residue, except for H. In some embodiments, the H840 residue of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, is mutated to an A. In some embodiments, the D10 residue of the amino acid sequence provided in SEQ ID NO: 10, or a

corresponding residue in any of the amino acid sequences provided in SEQ ID NOs: 11-260, is a D.

[00347] A number of Cas9 sequences from various species were aligned to determine whether corresponding homologous amino acid residues of D10 and H840 of SEQ ID NO: 10 or SEQ ID NO: 11 can be identified in other Cas9 proteins, allowing the generation of Cas9 variants with corresponding mutations of the homologous amino acid residues. The alignment was carried out using the NCBI Constraint-based Multiple Alignment Tool (COBALT(accessible at st-va.ncbi.nlm.nih.gov/tools/cobalt), with the following parameters. Alignment parameters: Gap penalties -11,-1; End-Gap penalties -5,-1. CDD Parameters: Use RPS BLAST on; Blast E-value 0.003; Find Conserved columns and Recompute on. Query Clustering Parameters: Use query clusters on; Word Size 4; Max cluster distance 0.8; Alphabet Regular.

[00348] An exemplary alignment of four Cas9 sequences is provided below. The Cas9 sequences in the alignment are: Sequence 1 (S1): SEQ ID NO: 11 | WP_010922251 | gi 499224711 | type II CRISPR RNA-guided endonuclease Cas9 [*Streptococcus pyogenes*]; Sequence 2 (S2): SEQ ID NO: 12 | WP_039695303 | gi 746743737 | type II CRISPR RNA-guided endonuclease Cas9 [*Streptococcus gallolyticus*]; Sequence 3 (S3): SEQ ID NO: 13 | WP_045635197 | gi 782887988 | type II CRISPR RNA-guided endonuclease Cas9 [*Streptococcus mitis*]; Sequence 4 (S4): SEQ ID NO: 14 | 5AXW_A | gi 924443546 | *Staphylococcus aureus* Cas9. The HNH domain (bold and underlined) and the RuvC domain (boxed) are identified for each of the four sequences. Amino acid residues 10 and 840 in S1 and the homologous amino acids in the aligned sequences are identified with an asterisk following the respective amino acid residue.

```

S1 1  --MDKK-YSIGLD*IGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLI--GALLFDSG--ETAAEATRLKRTARRRYT 73
S2 1  --MTKKNYSIGLD*IGTNSVGWAVITDDYKVPAKKMKVLGNTDKKYIKKNLL--GALLFDSG--ETAAAEATRLKRTARRRYT 74
S3 1  --M-KKGYSIGLD*IGTNSVGFAVITDDYKVPSKKMKVLGNTDKRFIKKNLI--GALLFDEG--TTAEAARRLKRTARRRYT 73
S4 1  GSHMKRNYILGLD*IGITSVGYGII--DYET-----RDVIDAGVRLFKEANVENNEGRRSKRGARRLKR 61

S1 74  RRKNRICYLQEIFSNEMAKVDDSFHRLEESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRL 153
S2 75  RRKNRLRYLQEIFANEIAKVDESFFQRLDESFLTDDDKTFDSHPIFGNKAEEDAYHQFPTIYHLRKKHLADSSEKADLRL 154
S3 74  RRKNRLRYLQEIFSEEMSKVDSSFFHRLDDSFLIPEDKRESKYPIFATLTEEKEYHKQFPTIYHLRKQLADSKEKTDLRL 153
S4 62  RRRHRIQRVKKLL-----FDYNLLTD-----HSELSGINPYEARVKGLSQKLSEEE 107

S1 154  IYLALAHMIKFRGHFLIEGDLNPDNSDVDKLFIQLVQTYQLFEENPINASGVDAKAILSARLSKSRRLENLIAQLPGEK 233
S2 155  VYLALAHMIKFRGHFLIEGELNAENTDVQKIFADFGVYNRTFDDSHLSEITVDVSILTEKISKSRRLENLIKYYPTEK 234
    
```


S3 154 IYLALAHMIKYRGHFLYEEAFDIKNNDIQKIFNEFISYDNTFEGSSLSGQNAQVEAIFTDKISKSAKRERVLKLPFDEK 233
S4 108 FSAALLHLAKRRG-----VHNVNEVEEDT----- 131

S1 234 KNGLFGNLIALLSLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEIT 313
S2 235 KNTLFGNLIALLALGLQPNFKTNFKLSEDAKLQFSKDTYEEDLEELLGKIGDDYADLFTSAKNLYDAILLSSGILTVDDNST 314
S3 234 STGLFSEFLKLVGNQADFKKHFDLEDKAPLQFSKDTYDEDELENLLGQIGDDFTDLFVSAKKLYDAILLSSGILTVTDPST 313
S4 132 -----GNELS-----TKEQISRN----- 144

S1 314 KAPLSASMIKRYDEHHQDLTLLKALVRQQLPPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKM--DGTEELLV 391
S2 315 KAPLSASMIKRYVEHHEDLEKLKEFIKANKSELYHDIKDKNKNKYAGYIENGVKQDEFYKYLKNILSKIKIDGSDYFLD 394
S3 314 KAPLSASMIERYENHQNDLAALKQFIKNNLPEKYDEVFSDQSKDGYAGYIDGKTTQETFYKYIKNLLSKF--EGTDYFLD 391
S4 145 ----SKALEEKYVAELQ-----LERLKKDG----- 165

S1 392 KLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYFLKDNREKIEKILTFRIPYYVGPLARGNSRFAWMTRKSEE 471
S2 395 KIEREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGDYYPFLKEKQDRIEKILTFRIPYYVGPLVRKDSRFAWAERYRDE 474
S3 392 KIEREDFLRKQRTFDNGSIPHQIHLQEMNAILRRQGEYYPFLKDNKEKIEKILTFRIPYYVGPLARGNRDFAWLTRNSDE 471
S4 166 --EVRGSINRFKTS-----YVKEAKQLLKVKQAYHQLDQSFIDTYIDLLETRRYYEGP--GEGSPFGW-----K 227

S1 472 TITPWNFEVVDKASQAQSFIERMTNFDKNLPNEKVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDL 551
S2 475 KITPWNFDKVIDKEKSAEKFITRMTLNDLYLPEEKVLPKHSVYETYAVYNELTKIKYVNEQGKE--SFFDSNMKQEIFDH 553
S3 472 AIRPWNFEIIVDKASSAEDFINKMTNYDLYLPEEKVLPKHSLLYETFAVYNELTKVKFIAEGLRDYQFLDSGQKKQIVNQ 551
S4 228 DIKEW-----YEMLMGHCTYFPEELRSVKYAYNADLYNALNDLNNLVITRDENEK---LEYEKFQIIEEN 289

S1 552 LFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDR---FNASLGTYHDLKLIKDKDFLDNEENEDILEDIVLTLTLFED 628
S2 554 VFKENRKVTKEKLLNLYLNKEFPYRIKDLIGLDKENKSNFASLGTYHDLKIL-DKAFLDDKVNEEVIEDIIKTLTLFED 632
S3 552 LFKENRKVTTEKDIIHYLHN-VDGYDGIELKGIKQ---FNASLSTYHDLKLIKDKDFMDDAKNEAILENIVHTLTLFED 627
S4 290 VFKQKKKPTLKQIAKEILVNEEDIKGYRVTSTGKPEF---TNLKVYHDIKDITARKEII---ENAELLDQIAKILTIYQS 363

S1 629 REMIEERLKYAHLFDDKVMKQLKR-RRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRNFQQLIHDDSLTFKED 707
S2 633 KDMIHERLQKYSDIFTANQLKKLER-RHYTGWGRLSYKLINGIRNKENKNTILDYLIDDGSANRNFQQLINDDTLFPKQI 711
S3 628 REMIKQRLAQYDLSFDEKVIKALTR-RHYTGWGRLSAKLINGICDKQGTGNTILDYLIDDGKINRNFQQLINDDGLSPKEI 706
S4 364 SEDIQEELTNLSELTQEETEQISNLKGYTGTHNLSLKAINLILDE-----LWHTNDNQIAIFNRLKLP----- 428

S1 708 IQKAQVSGQG[DSLHEHIANLAGSPAIAKKGILQTVKVVDELVKVMGRHKPENIVIEMA]RENQTT-----QK**GQKNSRERM** 781
S2 712 IQKSQVVGDV[DDIEAVVHDLPGSPAIAKKGILQSVKIVDELVKVMG-GNPDNIVIEMA]RENQTT-----NR**GRSQSQORL** 784
S3 707 IQKAQVIGKT[DDVKQVVQELSGSPAIAKKGILQSIKIVDELVKVMG-HAPESIVIEMA]RENQTT-----AR**GKKSQQRY** 779
S4 429 -KKVDLSQQK[EIPTTLVDDFILSPVVKRSFIQSIKVINAIKKYG--LPNDIIELA]REKNSKDAQKMINEM**MOKRNRQTN** 505

S1 782 **KRIEEGIKELGSQIL-----KEHPVENTQLONEKLYLYLQNGRDMYVDQELDINRLSD----YVDH*IVPQSFKDD** 850
S2 785 **KKLQNSLKELGSNILNEEKPSYIEDKVENSHLQNDQLFLYIYIQNGKDMYTGEALDIDHLS-----YDIDH*IIPQAFIKDD** 860
S3 780 **KRIEDSLKILASGL---DSNILKENPTDNNQLQNDRLFLYIYIQNGKDMYTGEALDINQLSS-----YDIDH*IIPQAFIKDD** 852
S4 506 **ERIEEIIIRTTGK-----ENAKYLIEKIKLHDMQEGKCLYSLEAIPLEDLLNPNFNYEVDH*IIPRVSFDN** 570

S1 851 **SIDNKVLTFRSDKNRGSNDVPSSEVVKMKMYRQLLNAKLITQRKFDN-LTKAERG**[GL-SELD-----KAGFIKRQLV] 922
S2 861 **SIDNRVLTSSAKNRGKSDVPSLDIVRARKAEWVRLYKSGLISKRRFDN-LTKAERG**[GL-TEAD-----KAGFIKRQLV] 932
S3 853 **SLDNRVLTSSKDNRGSNDVPSIEVQKRKAFWQQLLDSKLISERKFNN-LTKAERG**[GL-DERD-----KVGFIKRQLV] 924
S4 571 **SFNNKVLVKQEEASKGNRTPFQYLSSSDSKISYETFKKHILNLAQKGRISK**TKKKE[YLLEERDINRFSVQKDFINRNLV] 650

S1	923	ETRQITKHVAQILD SRMNTKYDENDKLIREVKVITLKS KLVSDFRKDFQFYK VREINNYHHAH DAYLNAVVG TALI KKY P	1002
S2	933	ETRQITKHVAQILDARFNTEHDENDKVIRDVKVITLKS NLVSQFRKDFEFYK VREINDYHHAH DAYLNAVVG TALL KKY P	1012
S3	925	ETRQITKHVAQILDARYNTEVNEKDKNR TVKII TLKS NLVSNFRKEFR LYK VREINDYHHAH DAYLNAV VAKAIL KKY P	1004
S4	651	DTRYATRGLMNLRSYFRVN-----NLDVKVKSINGGFTSFLRRKWKFKKERNKGYKHHAE DALIIA-----	712
S1	1003	KLESEFVYGDYKVYDVRKMLIAKSEQ--EIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKG---	1077
S2	1013	KLASEFVYGEYKKYDIRKFTINSSD----KATAKYFFYSNLMNFFKTKVYADGTVFERPIIETNAD-GEI AWNKQ---	1083
S3	1005	KLEPEFVYGEYQKYDLKRYISRSKDPKEVEKATEKYFFYSNLLNFKEEVHYADGTIVKRENI EYSKDTGEI AWNK E---	1081
S4	713	--NADFIFKEWKKLDKAKKVMENQM-----FEEKQAESMPEIETE QEYKEIFITPHQIK	764
S1	1078	-----RDFATVRKVL SMPQVNI VKKTEVQTGGFSKESILPKRNSDKLIARKKD---WDPK KYGGFDSPTVAYSVLV VAKV	1149
S2	1084	-----IDFEKVRKVL SYPQVNI VKKVETQTGGFSKESILPKGSDSKLI PRKTKKVYWDTKKYGGFDSPTVAYS VVADV	1158
S3	1082	-----KDFAI IKVLSLPQVNI VKKREVQTGGFSKESILPKGNSDKLI PRKTKDILLDTTKYGGFDS PVIAYSILLIADI	1156
S4	765	HIKDFKDYKYSHRVDKPNRELINDTLYSTRKDDKGN TLIVNNLNGLYDKDNDKL----KKLIN-KSP----EKLLMYHH	835
S1	1150	EKGKSKKLKSVKELLGITIMERS SFEKNPI-DFLEAKG-----YKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQKG	1223
S2	1159	EKGKAKKLKTVKELVGISIMERSFFEENPV-EFLENKG-----YHNIREDKLIKLPKYSLFEFEGGRRRLASASELQKG	1232
S3	1157	EKGKAKKLKTVKTLVGITIMEKA AFEENPI-TFLENKG-----YHNVRKENILCLPKYSLFELENGRRRLASAKELQKG	1230
S4	836	DPQTYQK LK-----LIMEQYGD EKNPLYKYEETGNYLTKYSKKDNGPVIIKKI KYGNKLN AHL DITDDYPNSRNKV	907
S1	1224	NELALPSKYVNFY LASHYEKLKGS PEDNEQKQLFVEQHKHYLDEIIEQISEFSKR VILADANLDKVL SAYNKH-----	1297
S2	1233	NEMVLPGYLV ELLYHAHRADNF-----NSTEYLN YVSEHKKEFEKVLSCVEDFANLYVDVEKNLSKIRAVADSM-----	1301
S3	1231	NEIVLPVYLT TLLYH SKNVHKL-----DEPGHLEYIQKHRNEFKDLLNLVSEFSQKYVLADANLEKI KSLYADN-----	1299
S4	908	VKLSLKP YRFD-VYLDNGVYKFV-----TVKNLDV IK--KENY YEVNSKAYEEAKLKKI SNQAEFIASFYNNDLIKING	979
S1	1298	RDKPIREQAENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQ SIT-----GLYETRI----DLSQL	1365
S2	1302	DNFSIEEISNSFINLLT LTALGAPADFNFLG EKI PRKRYTSTKECLNATLIHQ SIT-----GLYETRI----DLSKL	1369
S3	1300	EQADIEILANSFINLLT LTALGAPAAFKFFGK D IDRKYTTVSEILNATLIHQ SIT-----GLYETWI----DLSKL	1367
S4	980	ELYRVIGVNNDDLNR IEVNMIDITYR-EYLENMNDKRPPRI IKTIASKT---QSIKKYSTDILGNLYEVKSKKHPQIIKK	1055
S1	1366	GGD	1368
S2	1370	GEE	1372
S3	1368	GED	1370
S4	1056	G--	1056

[00349] The alignment demonstrates that amino acid sequences and amino acid residues that are homologous to a reference Cas9 amino acid sequence or amino acid residue can be identified across Cas9 sequence variants, including, but not limited to Cas9 sequences from different species, by identifying the amino acid sequence or residue that aligns with the reference sequence or the reference residue using alignment programs and algorithms known in the art. This disclosure provides Cas9 variants in which one or more of the amino acid residues identified by an asterisk in SEQ ID NOs: 11-14 (e.g., S1, S2, S3, and S4, respectively) are

mutated as described herein. The residues D10 and H840 in Cas9 of SEQ ID NO: 10 that correspond to the residues identified in SEQ ID NOs: 11-14 by an asterisk are referred to herein as “homologous” or “corresponding” residues. Such homologous residues can be identified by sequence alignment, *e.g.*, as described above, and by identifying the sequence or residue that aligns with the reference sequence or residue. Similarly, mutations in Cas9 sequences that correspond to mutations identified in SEQ ID NO: 10 herein, *e.g.*, mutations of residues 10, and 840 in SEQ ID NO: 10, are referred to herein as “homologous” or “corresponding” mutations. For example, the mutations corresponding to the D10A mutation in SEQ ID NO: 10 or S1 (SEQ ID NO: 11) for the four aligned sequences above are D11A for S2, D10A for S3, and D13A for S4; the corresponding mutations for H840A in SEQ ID NO: 10 or S1 (SEQ ID NO: 11) are H850A for S2, H842A for S3, and H560A for S4.

[00350] A total of 250 Cas9 sequences (SEQ ID NOs: 11-260) from different species were aligned using the same algorithm and alignment parameters outlined above. Amino acid residues homologous to residues 10, and 840 of SEQ ID NO: 10 were identified in the same manner as outlined above. The alignments are provided below. The HNH domain (bold and underlined) and the RuvC domain (boxed) are identified for each of the four sequences. Single residues corresponding to amino acid residues 10, and 840 in SEQ ID NO: 10 are boxed in SEQ ID NO: 11 in the alignments, allowing for the identification of the corresponding amino acid residues in the aligned sequences.

WP_010922251.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 11
WP_039695303.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus gallolyticus]	SEQ ID NO: 12
WP_045635197.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mitis]	SEQ ID NO: 13
5AXW_A	Cas9, Chain A, Crystal Structure [Staphylococcus Aureus]	SEQ ID NO: 14
WP_009880683.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 15
WP_010922251.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 16
WP_011054416.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 17
WP_011284745.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 18
WP_011285506.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 19
WP_011527619.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 20
WP_012560673.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 21
WP_014407541.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 22
WP_020905136.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 23
WP_023080005.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 24
WP_023610282.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 25
WP_030125963.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 26
WP_030126706.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 27
WP_031488318.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 28
WP_032460140.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 29
WP_032461047.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 30
WP_032462016.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 31
WP_032462936.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 32
WP_032464890.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 33
WP_033888930.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 34
WP_038431314.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 35
WP_038432938.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 36
WP_038434062.1	type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus pyogenes]	SEQ ID NO: 37
BAQ51233.1	CRISPR-associated protein, Csn1 family [Streptococcus pyogenes]	SEQ ID NO: 38
KGE60162.1	hypothetical protein MGAS2111_0903 [Streptococcus pyogenes MGAS2111]	SEQ ID NO: 39
KGE60856.1	CRISPR-associated endonuclease protein [Streptococcus pyogenes SS1447]	SEQ ID NO: 40
WP_002989955.1	MULTISPECIES: type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus]	SEQ ID NO: 41
WP_003030002.1	MULTISPECIES: type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus]	SEQ ID NO: 42

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MULTISPECIES: type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus] SEQ ID NO: 43

WP_003065552.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 44

WP_001040076.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 45

WP_001040078.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 46

WP_001040080.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 47

WP_001040081.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 48

WP_001040083.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 49

WP_001040085.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 50

WP_001040087.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 51

WP_001040088.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 52

WP_001040089.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 53

WP_001040090.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 54

WP_001040091.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 55

WP_001040092.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 56

WP_001040094.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 57

WP_001040095.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 58

WP_001040096.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 59

WP_001040097.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 60

WP_001040098.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 61

WP_001040099.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 62

WP_001040100.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 63

WP_001040104.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 64

WP_001040105.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 65

WP_001040106.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 66

WP_001040107.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 67

WP_001040108.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 68

WP_001040109.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 69

WP_001040110.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 70

WP_015058523.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 71

WP_017643650.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 72

WP_017647151.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 73

WP_017648376.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 74

WP_017649527.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 75

WP_017771611.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 76

WP_017771984.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 76

CFQ25032.1 CRISPR-associated protein [Streptococcus agalactiae] SEQ ID NO: 77
 CFV16040.1 CRISPR-associated protein [Streptococcus agalactiae] SEQ ID NO: 78
 KLJ37842.1 CRISPR-associated protein Csn1 [Streptococcus agalactiae] SEQ ID NO: 79
 KLJ72361.1 CRISPR-associated protein Csn1 [Streptococcus agalactiae] SEQ ID NO: 80
 KLL20707.1 CRISPR-associated protein Csn1 [Streptococcus agalactiae] SEQ ID NO: 81
 KLL42645.1 CRISPR-associated protein Csn1 [Streptococcus agalactiae] SEQ ID NO: 82
 WP_047207273.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 83
 WP_047209694.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 84
 WP_050198062.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 85
 WP_050201642.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 86
 WP_050204027.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 87
 WP_050881965.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 88
 WP_050886065.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus agalactiae] SEQ ID NO: 89
 AHN30376.1 CRISPR-associated protein Csn1 [Streptococcus agalactiae 138P] SEQ ID NO: 90
 EAO78426.1 reticulocyte binding protein [Streptococcus agalactiae H36B] SEQ ID NO: 91
 CCW42055.1 CRISPR-associated protein, SAG0894 family [Streptococcus agalactiae ILRI112] SEQ ID NO: 92
 WP_003041502.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus anginosus] SEQ ID NO: 93
 WP_037593752.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus anginosus] SEQ ID NO: 94
 WP_049516684.1 CRISPR-associated protein Csn1 [Streptococcus anginosus] SEQ ID NO: 95
 GAD46167.1 hypothetical protein ANG6_0662 [Streptococcus anginosus T5] SEQ ID NO: 96
 WP_018363470.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus caballi] SEQ ID NO: 97
 WP_003043819.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus canis] SEQ ID NO: 98
 WP_006269658.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus constellatus] SEQ ID NO: 99
 WP_048800889.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus constellatus] SEQ ID NO: 100
 WP_012767106.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus constellatus] SEQ ID NO: 101
 WP_014612333.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus dysgalactiae] SEQ ID NO: 102
 WP_015017095.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus dysgalactiae] SEQ ID NO: 103
 WP_015057649.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus dysgalactiae] SEQ ID NO: 104
 WP_048327215.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus dysgalactiae] SEQ ID NO: 105
 WP_049519324.1 CRISPR-associated protein Csn1 [Streptococcus dysgalactiae] SEQ ID NO: 106
 WP_012515931.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus equi] SEQ ID NO: 107
 WP_021320964.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus equi] SEQ ID NO: 108
 WP_037581760.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus equi] SEQ ID NO: 109
 WP_004232481.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus equinus] SEQ ID NO: 110

WP_009854540.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus gallolyticus] SEQ ID NO: 111
 WP_012962174.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus gallolyticus] SEQ ID NO: 112
 WP_039695303.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus gallolyticus] SEQ ID NO: 113
 WP_014334983.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus infantarius] SEQ ID NO: 114
 WP_003099269.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus iniae] SEQ ID NO: 115
 AHY15608.1 CRISPR-associated protein Csn1 [Streptococcus iniae] SEQ ID NO: 116
 AHY17476.1 CRISPR-associated protein Csn1 [Streptococcus iniae] SEQ ID NO: 117
 ESR09100.1 hypothetical protein IUSA1_08595 [Streptococcus iniae IUSA1] SEQ ID NO: 118
 AGM98575.1 CRISPR-associated protein Cas9/Csn1, subtype II/NMEMI [Streptococcus iniae SF1] SEQ ID NO: 119
 ALF27331.1 CRISPR-associated protein Csn1 [Streptococcus intermedius] SEQ ID NO: 120
 WP_018372492.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus massiliensis] SEQ ID NO: 121
 WP_045618028.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mitis] SEQ ID NO: 122
 WP_045635197.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mitis] SEQ ID NO: 123
 WP_002263549.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 124
 WP_002263887.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 125
 WP_002264920.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 126
 WP_002269043.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 127
 WP_002269448.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 128
 WP_002271977.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 129
 WP_002272766.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 130
 WP_002273241.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 131
 WP_002275430.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 132
 WP_002276448.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 133
 WP_002277050.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 134
 WP_002277364.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 135
 WP_002279025.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 136
 WP_002279859.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 137
 WP_002280230.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 138
 WP_002281696.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 139
 WP_002282247.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 140
 WP_002282906.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 141
 WP_002283846.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 142
 WP_002287255.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 143
 WP_002288990.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus mutans] SEQ ID NO: 144

WP_002289641.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 145
WP_002290427.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 146
WP_002295753.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 147
WP_002296423.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 148
WP_002304487.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 149
WP_002305844.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 150
WP_002307203.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 151
WP_002310390.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 152
WP_002352408.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 153
WP_012997688.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 154
WP_014677909.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 155
WP_019312892.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 156
WP_019313659.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 157
WP_019314093.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 158
WP_019315370.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 159
WP_019803776.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 160
WP_019805234.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 161
WP_024783594.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 162
WP_024784288.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 163
WP_024784666.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 164
WP_024784894.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 165
WP_024786433.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus mutans]	SEQ ID NO: 166
WP_049473442.1	CRISPR-associated protein	Csn1	[Streptococcus mutans]			[Streptococcus mutans]	SEQ ID NO: 167
WP_049474547.1	CRISPR-associated protein	Csn1	[Streptococcus mutans]			[Streptococcus mutans]	SEQ ID NO: 168
EMC03581.1	hypothetical protein	SMU69_09359	[Streptococcus mutans NLML4]			[Streptococcus mutans NLML4]	SEQ ID NO: 169
WP_000428612.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus oralis]	SEQ ID NO: 170
WP_000428613.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus oralis]	SEQ ID NO: 171
WP_049523028.1	CRISPR-associated protein	Csn1	[Streptococcus parasanguinis]			[Streptococcus parasanguinis]	SEQ ID NO: 172
WP_003107102.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus parauberis]	SEQ ID NO: 173
WP_054279288.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus phocae]	SEQ ID NO: 174
WP_049531101.1	CRISPR-associated protein	Csn1	[Streptococcus pseudopneumoniae]			[Streptococcus pseudopneumoniae]	SEQ ID NO: 175
WP_049538452.1	CRISPR-associated protein	Csn1	[Streptococcus pseudopneumoniae]			[Streptococcus pseudopneumoniae]	SEQ ID NO: 176
WP_049549711.1	CRISPR-associated protein	Csn1	[Streptococcus pseudopneumoniae]			[Streptococcus pseudopneumoniae]	SEQ ID NO: 177
WP_0078996501.1	type II	CRISPR	RNA-guided	endonuclease	Cas9	[Streptococcus pseudoporcinus]	SEQ ID NO: 178

EFR44625.1 CRISPR-associated protein, Csn1 family [Streptococcus pseudoporcinus SPIN 20026] SEQ ID NO: 179
 WP_002897477.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus sanguinis] SEQ ID NO: 180
 WP_002906454.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus sanguinis] SEQ ID NO: 181
 WP_009729476.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus sp. F0441] SEQ ID NO: 182
 CQR24647.1 CRISPR-associated protein [Streptococcus sp. FF10] SEQ ID NO: 183
 WP_000066813.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus sp. M334] SEQ ID NO: 184
 WP_009754323.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus sp. taxon 056] SEQ ID NO: 185
 WP_044674937.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus suis] SEQ ID NO: 186
 WP_044676715.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus suis] SEQ ID NO: 187
 WP_044680361.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus suis] SEQ ID NO: 188
 WP_044681799.1 type II CRISPR RNA-guided endonuclease Cas9 [Streptococcus suis] SEQ ID NO: 189
 WP_049533112.1 CRISPR-associated protein Csn1 [Streptococcus suis] SEQ ID NO: 190
 WP_029090905.1 type II CRISPR RNA-guided endonuclease Cas9 [Brochothrix thermosphacta] SEQ ID NO: 191
 WP_006506696.1 type II CRISPR RNA-guided endonuclease Cas9 [Catenibacterium mitsuokai] SEQ ID NO: 192
 AIT42264.1 Cas9hc:NLS:HA [Cloning vector pYB196] SEQ ID NO: 193
 WP_034440723.1 type II CRISPR endonuclease Cas9 [Clostridiales bacterium S5-A11] SEQ ID NO: 194
 AKQ21048.1 Cas9 [CRISPR-mediated gene targeting vector p(bhsp68-Cas9)] SEQ ID NO: 195
 WP_004636532.1 type II CRISPR RNA-guided endonuclease Cas9 [Dolosigranulum pigrum] SEQ ID NO: 196
 WP_002364836.1 MULTISPECIES: type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus] SEQ ID NO: 197
 WP_016631044.1 MULTISPECIES: type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus] SEQ ID NO: 198
 EMS75795.1 hypothetical protein H318_06676 [Enterococcus durans IPLA 655] SEQ ID NO: 199
 WP_002373311.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 200
 WP_002378009.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 201
 WP_002407324.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 202
 WP_002413717.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 203
 WP_010775580.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 204
 WP_010818269.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 205
 WP_010824395.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 206
 WP_016622645.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 207
 WP_033624816.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 208
 WP_033625576.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 209
 WP_033789179.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecalis] SEQ ID NO: 210
 WP_002310644.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 211
 WP_002312694.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 212

WP_002314015.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 213
 WP_002320716.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 214
 WP_002330729.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 215
 WP_002335161.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 216
 WP_002345439.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 217
 WP_034867970.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 218
 WP_047937432.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus faecium] SEQ ID NO: 219
 WP_010720994.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus hirae] SEQ ID NO: 220
 WP_010737004.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus hirae] SEQ ID NO: 221
 WP_034700478.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus hirae] SEQ ID NO: 222
 WP_007209003.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus italicus] SEQ ID NO: 223
 WP_023519017.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus mundtii] SEQ ID NO: 224
 WP_010770040.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus phoeniculicola] SEQ ID NO: 225
 WP_048604708.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus sp. AM1] SEQ ID NO: 226
 WP_010750235.1 type II CRISPR RNA-guided endonuclease Cas9 [Enterococcus villorum] SEQ ID NO: 227
 AII16583.1 Cas9 endonuclease [Expression vector pCas9] SEQ ID NO: 228
 WP_029073316.1 type II CRISPR RNA-guided endonuclease Cas9 [Kandleria vitulina] SEQ ID NO: 229
 WP_031589969.1 type II CRISPR RNA-guided endonuclease Cas9 [Kandleria vitulina] SEQ ID NO: 230
 KDA45870.1 CRISPR-associated protein Cas9/Csn1, subtype II/NMEMI [Lactobacillus animalis] SEQ ID NO: 231
 WP_039099354.1 type II CRISPR RNA-guided endonuclease Cas9 [Lactobacillus curvatus] SEQ ID NO: 232
 AKP02966.1 hypothetical protein ABB45_04605 [Lactobacillus farciminis] SEQ ID NO: 233
 WP_010991369.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria innocua] SEQ ID NO: 234
 WP_033838504.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria innocua] SEQ ID NO: 235
 EHN60060.1 CRISPR-associated protein, Csn1 family [Listeria innocua ATCC 33091] SEQ ID NO: 236
 EFR89594.1 crispr-associated protein, Csn1 family [Listeria innocua FSL S4-378] SEQ ID NO: 237
 WP_038409211.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria ivanovii] SEQ ID NO: 238
 EFR95520.1 crispr-associated protein Csn1 [Listeria ivanovii FSL F6-596] SEQ ID NO: 239
 WP_003723650.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes] SEQ ID NO: 240
 WP_003727705.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes] SEQ ID NO: 241
 WP_003730785.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes] SEQ ID NO: 242
 WP_003733029.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes] SEQ ID NO: 243
 WP_003739838.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes] SEQ ID NO: 244
 WP_014601172.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes] SEQ ID NO: 245
 WP_023548323.1 type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes] SEQ ID NO: 246

WP_031665337.1	type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes]	SEQ ID NO: 247
WP_031669209.1	type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes]	SEQ ID NO: 248
WP_033920898.1	type II CRISPR RNA-guided endonuclease Cas9 [Listeria monocytogenes]	SEQ ID NO: 249
AKI42028.1	CRISPR-associated protein [Listeria monocytogenes]	SEQ ID NO: 250
AKI50529.1	CRISPR-associated protein [Listeria monocytogenes]	SEQ ID NO: 251
EFR83390.1	crispr-associated protein Csn1 [Listeria monocytogenes FSL F2-208]	SEQ ID NO: 252
WP_046323366.1	type II CRISPR RNA-guided endonuclease Cas9 [Listeria seeligeri]	SEQ ID NO: 253
AKE81011.1	Cas9 [Plant multiplex genome editing vector pYLCRISPR/Cas9Pubi-H]	SEQ ID NO: 254
CUO82355.1	Uncharacterized protein conserved in bacteria [Roseburia hominis]	SEQ ID NO: 255
WP_033162887.1	type II CRISPR RNA-guided endonuclease Cas9 [Sharpea azabuensis]	SEQ ID NO: 256
AGZ01981.1	Cas9 endonuclease [synthetic construct]	SEQ ID NO: 257
AKA60242.1	nuclease deficient Cas9 [synthetic construct]	SEQ ID NO: 258
AKS40380.1	Cas9 [Synthetic plasmid pFC330]	SEQ ID NO: 259
4UN5_B	Cas9, Chain B, Crystal structure	SEQ ID NO: 260
WP_010922251	MDKK- YSIGLDIGTNSVGWAVITDEYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_039695303	MTKknYSIGLDIGTNSVGWAVITDDYKVP PAKMKV LVGN TDKKIYIKKNIIGALLFDSGETA --EATRLKRTARRRYT	74
WP_045635197	K-KG-YSIGLDIGTNSVGFVAVITDDYKVP SKKMKV LVGN TDKRFIKKNIIGALLFDEGTTA --EARRLKRTARRRYT	73
5AXW_A	MKRN-YILGLDIGITSVGYGII--DYET-----RDVIDA---GVRLFKEANVenEGRRSKRGARLKR	61
WP_009880683	-----	
WP_010922251	MDKK-YSIGLDIGTNSVGWAVITDEYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_011054416	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKLVGN TDRHIGIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_011284745	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_011285506	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_011527619	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGEIA --EATRLKRTARRRYT	73
WP_012560673	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_014407541	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFGSGETA --EATRLKRTARRRYT	73
WP_020905136	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_023080005	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKLVGN TDRHIGIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_023610282	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKLVGN TDRHIGIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_030125963	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_030126706	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHIGIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_031488318	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73
WP_032460140	MDKK-YSIGLDIGTNSVGWAVITDDYKVP SKKFKVLGN TDRHSIKKNIIGALLFDSGETA --EATRLKRTARRRYT	73

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 KGE60856 -----
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 WP_010750235 1 MNKA-YTLGLDITGNSVGWAVTDDYRLMAKMPVHSMKMEKKIKKNFWGARLFDEGQTA--EERRNKRTARRRLLR 73
 AII16583 1 ADKK-YSIGLDITGNSVGWAVITDEYKVPKFKVLGNTDRHSIKKLI GALLFDSGETA--EATRLKRTARRRYT 112
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 WP_039099354 1 MSRP-YNIGLDITGSSIGWSVWDDQSKLVSVR-----GKYGYGVRLYDEGQTA--AERRSFRTTTRRLK 61
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WP_033838504	1	MKKP-YTIGLDIGTNSVGWAVLTDQYDLVKKRMKIAGDSEKKQIKKNFWGVRLFDEGQTA--ADRRMARTARRIE	73
EHN60060	1	MKKP-YTIGLDIGTNSVGWAVLTDQYDLVKKRMKIAGDSEKKQIKKNFWGVRLFDEGQTA--ADRRMARTARRIE	76
EFR89594		-----	
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AKI50529	1	MKNP-YTIGLDIGTNSVGWAVLTDQYDLVKKRMKVAGNSDKKQIKKNFWGVRLFDDGQTA--VDRRMNRTARRIE	76
EFR83390		-----	
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CUO82355	1	I-VD-YCIGLDLGTSGWAVVDMNHRMKNR-----GKHLWGSRLFSAETA--ATRRSSRSIRRRYN	64
WP_033162887	1	KDIR-YSIGLDIGTNSVGWAVMDEHYELLKKG-----NHHMWGSRLFDAEPA--ATTRASRSIRRRYN	65
AGZ01981	1	ADKK-YSIGLDIGTNSVGWAVITDEYKVPKFKVLTGNTDRHSIKKNLIGALLFDSGETA--EATRLKRTARRRYT	106
AKA60242	1	MDKK-YSIGLAIGTNSVGWAVITDEYKVPKFKVLTGNTDRHSIKKNLIGALLFDSGETA--EATRLKRTARRRYT	73
AKS40380	1	MDKK-YSIGLDIGTNSVGWAVITDEYKVPKFKVLTGNTDRHSIKKNLIGALLFDSGETA--EATRLKRTARRRYT	73
4UN5_B	1	MDKK-YSIGLAIGTNSVGWAVITDEYKVPKFKVLTGNTDRHSIKKNLIGALLFDSGETA--EATRLKRTARRRYT	77

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WP_039695303	75	RRKNRLRYLQEI FANEIAKVDESFFQRLDE-SFLT--DDDKT--F	DSHPIFGNKA-EEDAYHQKFPTIYHLRKHLA	144
WP_045635197	74	RRKNRLRYLQEI FSEEMSKVDSFFHRLDD-SFLI--PEDKR--E	SKYPIFATLT-EEKEYHKQFPTIYHLRQOLA	143
5AXW A	62	RRRHRIQRVKKLLFD-----YNLLTDhSELS	--NPYEARVK-----GLSQKLS	104
WP_009880683		-----	-----	
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WP_032460140	74	RRKNRICYLQEI FSNEMAKVDDSFHRL EE-SFLV--EEDKK--H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLA	143
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WP_032464890	74	RRKNRICYLQEI FSNEMAKVDDSFHRL EE-SFLV--EEDKK--H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	143
WP_033888930		-----	-----	
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WP_038434062	74	RRKNRICYLQEI FSNEMAKVDDSFHRL EE-SFLV--EEDKK--H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLA	143
BAQ51233	1	-----MAKVDDSFHRL EE-SFLV--EEDKK--H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	54
KGE60162		-----	-----	
KGE60856		-----	-----	
WP_002989955	74	RRKNRICYLQEI FSNEMAKVDDSFHRL EE-SFLV--EEDKK--H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	143
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WP_003065552	75	RRKNRLRYLQEI FAEEMTKVDESFFQRLDE-SFLRwdDNKK--L	GRYPIFGNKA-DVVKYHQEFPTIYHLRKHLA	146
WP_001040076	74	RRRNRLRYLQEI FAEEMSKVDDSFHRL EE-SFLV--EEDKR--G	SKYPIFATLQ-EEKYHEKFFPTIYHLRKELA	143

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KLJ72361	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATLQ	-EEKDY	HEK	FSTI	YHLR	KELA	143					
KL20707	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATLQ	-EEKDY	HEK	FSTI	YHLR	KELA	143					
KL42645	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EDDKR	---G	SKYPI	FATM	Q-EEKDY	HEK	FPTI	YHLR	KELA	143					
WP_047207273	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATLQ	-EEKDY	HEK	FSTI	YHLR	KELA	143					
WP_047209694	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATM	Q-EEKDY	HEK	FPTI	YHLR	KELA	143					
WP_050198062	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATLQ	-EEKDY	HEK	FSTI	YHLR	KELA	143					
WP_050201642	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATLQ	-EEKDY	HEK	FSTI	YHLR	KELA	143					
WP_050204027	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EDDKR	---G	SKYPI	FATM	Q-EEKDY	HEK	FPTI	YHLR	KELA	143					
WP_050881965	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATLQ	-EEKDY	HEK	FSTI	YHLR	KELA	143					
WP_050886065	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATLQ	-EEKDY	HEK	FSTI	YHLR	KELA	143					
AHN30376	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATM	Q-EEKDY	HEK	FPTI	YHLR	KELA	143					
EA078426	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	SKYPI	FATLQ	-EEKDY	HEK	FSTI	YHLR	KELA	143					
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WP_003041502	74	RRNRILYLQEI	FAEEMSKVD	DSFFHRL	ED-SFLV	--EEDKR	---G	ERHP	I	FGNIA	-AEVKY	H	DE	FPTI	YHLR	KHLA	143			
WP_037593752	75	RRNRILYLQEI	FTEEMNKVD	ENFFQRL	DD-SFLV	--EEDKQ	---G	SKYPI	FGTL	Q-EEKEY	H	K	F	K	F	TI	YHLR	REELA	144	
WP_049516684	75	RRNRILYLQEI	FTEEMNKVD	ENFFQRL	DD-SFLV	--EEDKR	---G	SRYP	I	FGNIA	-AEVKY	H	D	D	F	F	TI	YHLR	KHLV	144
GAD46167	74	RRNRILYLQEI	FTEEMNKVD	ENFFQRL	DD-SFLV	--EEDKQ	---G	SKYPI	FGTL	Q-EEKEY	H	K	F	K	F	TI	YHLR	REELA	143	
WP_018363470	75	RRNRILYLQEI	FTEEMNKVD	ENFFQRL	DD-SFLV	--EEDKQ	---G	SKYPI	FGTL	Q-EEKEY	H	K	F	K	F	TI	YHLR	REELA	143	
WP_003043819	74	RRNRILYLQEI	FANEMAKL	DDSFQRL	EE-SFLT	--DNDKN	---F	DSHP	I	FGNKA	-EEDAY	H	Q	K	F	TI	YHLR	KHLA	144	
WP_006269658	74	RRNRILYLQEI	FTEEMNKVD	ENFFQRL	DD-SFLT	--EEDKQ	---N	ERHP	I	FGNIA	-AEVKY	H	D	D	F	F	TI	YHLR	RRHLA	143
WP_048800889	74	RRNRILYLQEI	FTEEMNKVD	ENFFQRL	DD-SFLT	--EEDKR	---G	EHP	I	FGNIA	-AEVKY	H	D	D	F	F	TI	YHLR	RRHLA	143
WP_012767106	74	RRNRILYLQEI	FSSEMSKVD	DSFFHRL	EE-SFLV	--EEDKQ	---H	SKYPI	FGTL	Q-EEKEY	Y	K	E	F	E	F	TI	YHLR	RKRLA	143
WP_014612333	74	RRNRILYLQEI	FSSEMSKVD	DSFFHRL	EE-SFLV	--EEDKQ	---H	ERHP	I	FGNIV	-DEVAY	H	E	K	Y	P	TI	YHLR	RKRLA	143
WP_015017095	74	RRNRILYLQEI	FSSEMSKVD	DSFFHRL	EE-SFLV	--EEDKQ	---H	ERHP	I	FGNIV	-DEVAY	H	E	K	Y	P	TI	YHLR	RKRLA	143
WP_015057649	74	RRNRILYLQEI	FSSEMSKVD	DSFFHRL	EE-SFLV	--EEDKQ	---H	ERHP	I	FGNIV	-DEVAY	H	E	K	Y	P	TI	YHLR	RKRLA	143
WP_048327215	74	RRNRILYLQEI	FSSEMSKVD	DSFFHRL	EE-SFLV	--EEDKQ	---H	ERHP	I	FGNIV	-DEVAY	H	E	K	Y	P	TI	YHLR	RKRLA	143
WP_049519324	74	RRNRILYLQEI	FSSEMSKVD	DSFFHRL	EE-SFLV	--EEDKQ	---H	ERHP	I	FGNIV	-DEVAY	H	E	K	Y	P	TI	YHLR	RKRLA	143
WP_012515931	74	RRNRILYLQEI	FTEEMAKV	DDGFFQRL	ED-SFYV	--LEDKE	---G	ERHP	I	FGNIV	-DEVAY	H	E	K	Y	P	TI	YHLR	RKRLA	143
WP_021320964	74	RRNRILYLQEI	FTEEMAKV	DDGFFQRL	ED-SFYV	--LEDKE	---G	NKHP	I	FANLA	-DEVAY	H	K	K	Y	P	TI	YHLR	RKELV	143
WP_037581760	74	RRNRILYLQEI	FTEEMAKV	DDGFFQRL	ED-SFYV	--LEDKE	---G	NKHP	I	FANLA	-DEVAY	H	K	K	Y	P	TI	YHLR	RKELV	143
WP_004232481	74	RRNRILYLQEI	FAKEMAKV	DESFFQRL	EE-SFLT	--DDDKT	---F	DSHP	I	FGNKA	-EEDT	Y	H	Q	E	F	TI	YHLR	KHLA	143
WP_009854540	75	RRNRILYLQEI	FAEEMTKV	DESFFYRL	DE-SFLT	--TDEKD	---F	ERHP	I	FGNKA	-EEDAY	H	Q	K	F	TI	YHLR	NYLA	144	
WP_012962174	75	RRNRILYLQEI	FAEEMAKV	DESFFYRL	DE-SFLT	--TDDKD	---F	ERHP	I	FGNKA	-DEIK	Y	H	Q	E	F	TI	YHLR	KHLA	144

WP_039695303	75	RRKNRLRYLQEI	FANEIAKVDESFQRLDE-SFLT--DDDKT---	DSHPIFGNKA-EEDAYHQKFPTIYHLRKHHLA	144
WP_014334983	74	RRKNRLRYLQEI	FAKEMTKVDESFQRLDE-SFLT--DDDKT---	DSHPIFGNKA-EEDAYHQKFPTIYHLRKHHLA	143
WP_003099269	74	RRKYRIKELQKI	FSSEMNELDIAFFPRLSE-SFLV--SDDKE---	ENHPIFGNLK-DEITYHNDYPTIYHLRQTLA	143
AHY15608	74	RRKYRIKELQKI	FSSEMNELDIAFFPRLSE-SFLV--SDDKE---	ENHPIFGNLK-DEITYHNDYPTIYHLRQTLA	143
AHY17476	74	RRKYRIKELQKI	FSSEMNELDIAFFPRLSE-SFLV--SDDKE---	ENHPIFGNLK-DEITYHNDYPTIYHLRQTLA	143
ESR09100					
AGM98575	74	RRKYRIKELQKI	FSSEMNELDIAFFPRLSE-SFLV--SDDKE---	ENHPIFGNLK-DEITYHNDYPTIYHLRQTLA	143
ALF27331	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_018372492	74	RRNRIRYLQHI	FAEEMNRADENFFHRLKE-SFFV--EEDKT---	SKYPIFGTLE-EEKYHKNYPTIYHLRKTILA	143
WP_045618028	75	RRKNRLRYLQEI	FTEEMSKVDISFFHRLDD-SFLV--PEDKR---	SKYPIFATLE-EEKEYHKNFPTIYHLRKHHLA	144
WP_045635197	74	RRKNRLRYLQEI	FSEEMSKVDDSFHRLDD-SFLI--PEDKR---	SKYPIFATLT-EEKEYHQKFPTIYHLRQQLA	143
WP_002263549	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002263887	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLT--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002264920	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLT--DDDKN---	DSYPIFGNKA-EEDAYHQKFPTIYHLRKHHLA	143
WP_002269043	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002269448	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002271977	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002272766	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002273241	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002275430	74	RRNRILYLQEI	FAEEMSKVDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002276448	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002277050	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLT--DDDKN---	DSHPIFGNKA-EEDAYHQKFPTIYHLRKHHLA	143
WP_002277364	74	RRNRILYLQEI	FAEEMSKVDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002279025	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002279859	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLT--DDDKN---	DSHPIFGNKA-EEDAYHQKFPTIYHLRKHHLA	143
WP_002280230	74	RRNRILYLQEI	FAEEMSKVDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002281696	74	RRNRILYLQEI	FAEEMSKVDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002282247	74	RRNRILYLQEI	FAEEMSKVDDSFHRLDE-SFLT--DDDKN---	DSHPIFGNKA-EEDAYHQKFPTIYHLRKHHLA	143
WP_002282906	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002283846	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002287255	74	RRNRILYLQEI	FAEEMSKVDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002288990	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002289641	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143
WP_002290427	74	RRNRILYLQEI	FSEEMGVKDDSFHRLDE-SFLV--TEDKR---	ERHPIFGNLE-EEVKYHENFPTIYHLRQYLA	143

RRNRILYLQEI	FSEEMGVND	FFHRL	ED-SFLV	--TEDKR	---	74	WP_002295753	ERHPI	FGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_002296423	ERHPI	FGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FAEEMQVDES	FFQRL	DD-SFLV	--EEDKR	---	74	WP_002304487	SRYP	IFGTLK	-EEKY	HKFEFKT	YHLR	REKLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_002305844	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_002307203	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_002310390	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_002352408	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FAEEMSKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_012997688	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_014677909	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FAEEMSKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_019312892	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_019313659	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_019314093	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_019315370	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_019803776	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_019805234	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_024783594	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FAEEMSKVDD	FFHRL	ED-SFLT	--DDDKN	---	74	WP_024784288	DSHP	IFGNKA	-EEDAY	HQKFP	TIYHLR	KKHLA	143
RRNRILYLQEI	FAEEMSKVDD	FFHRL	ED-SFLT	--TEDKR	---	74	WP_024784666	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_024784894	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FAEEMKVDD	FFHRL	ED-SFLT	--DDDKN	---	74	WP_024786433	DSHP	IFGNKA	-EEDAY	HQKFP	TIYHLR	KKHLA	143
RRNRILYLQEI	FAEEMSKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_049473442	ERHP	IFGNLE	-EEVKY	YENFPTI	YHLRQ	YLA	143
RRNRILYLQEI	FSEEMKVDD	FFHRL	ED-SFLV	--TEDKR	---	74	WP_049474547	ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	143
EMC03581						67		ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	136
WP_000428612						75		ERHP	IFGNLE	-EEVKY	HENFPTI	YHLRQ	YLA	144
WP_000428613						75		SKYP	IFATLI	-EEKE	YHKQFP	TIYHLR	KKQLA	144
WP_049523028						74		SKYP	IFATLA	-EEKE	YHKQFP	TIYHLR	KKQLA	144
WP_003107102						43		SKYP	IFGTLE	-EEKE	YHKQFP	TIYHLR	KKILA	143
WP_054279288						76		DRHP	IFGNIK	-DEV	YHKNYPTI	YHLR	KKLA	112
WP_049531101						75		EAHP	IFGTLQ	-EEKAY	HDNYPTI	YHLR	KKALA	145
WP_049538452						75		SKYP	IFATLT	-EEKE	YHKQFP	TIYHLR	KKQLA	144
WP_049549711						75		SKYP	IFATLA	-EEKE	YHKNYPTI	YHLR	KKQLA	144
WP_007896501						76		SKYP	IFATLV	-EEKE	YHKQFP	TIYHLR	KKQLA	144
EFR44625						28		DKHP	IFGNSK	-EERAY	HKTYPTI	YHLR	KKDLA	145
WP_002897477						74		DKHP	IFGNSK	-EERAY	HKTYPTI	YHLR	KKDLA	97
								SKYP	IFATLQ	-EEKE	YHKQFP	TIYHLR	KKQLA	143

WP_002906454	74	RRKNRLRYLQEI FSEESI KLDSFFHRLDD-SFLV--PEDKR---	143
WP_009729476	75	RRKNRLRYLQEI FSEIIGKVDSSFFHRLDD-SFLI--PEDKR---	144
CQR24647	74	RRNRILYLQDI FSPELNQVDESFLHRLDD-SFLVa--EDKR---	143
WP_000066813	75	RRKNRLRYLQEI FSQEI SKVDSSFFHRLDD-FFLV--PEDKR---	144
WP_009754323	75	RRKNRLRYLQEI FAEEMSKVDSSFFHRLDD-SFLV--PEDKS---	144
WP_044674937	74	RRNRILYLQEI FAEEINKIDDSFFQRLDD-SFLIv--EDKQ---	143
WP_044676715	74	RRNRILYLQEI FAEEINKIDDSFFQRLDD-SFLIv--EDKQ---	143
WP_044680361	74	RRNRILYLQEI FAEEINKIDDSFFQRLDD-SFLIv--EDKQ---	143
WP_044681799	74	RRNRILYLQEI FAEEINKIDDSFFQRLDD-SFLIv--EDKQ---	143
WP_049533112	74	RRNRILYLQEI FAEEINKVDENFFQRLDD-SFLV--DEDKR---	143
WP_029090905	28	HRKFRLLLEDMEKEI LSKDPSFFIRLKE-AFLSpkDEQK---F	100
WP_006506696	61	KRRERIRLLRAI LQDMVLEKDP TFFIRLEHTSFLD--EEDKakylG	139
AIT42264	74	RRKNRICYLQEI FSNEMAKVDSSFFHRLDE-SFLV--EEDKK--H	143
WP_034440723	73	RRFRIRLQKI FDKSMGEVDSNFFHRLDE-SFLV--EEDKE---Y	142
AKQ21048	74	RRKNRICYLQEI FSNEMAKVDSSFFHRLDE-SFLV--EEDKK--H	143
WP_004636532	74	RRNRILYLQDI FQOPMLAIDENFFHRLDD-SFFV--PDDKS---Y	143
WP_002364836	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_016631044	25	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	94
EMS75795		-----	
WP_002373311	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_002378009	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_002407324	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_002413717	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_010775580	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_010818269	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_010824395	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_016622645	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_033624816	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_033625576	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_033789179	74	RRNRILYLQAF FEEAMTLDENFFARLQE-SFLV--PEDKK---W	143
WP_002310644	74	RRQRILELQKI FAPEILKIDEHFFARLNE-SFLV--LDEKK---Q	143
WP_002312694	74	RRQRILELQKI FAPEILKIDEHFFARLNE-SFLV--PDEKK---Q	143
WP_002314015	74	RRQRILELQKI FAPEILKIDEHFFARLNE-SFLV--LDEKK---Q	143
WP_002320716	74	RRQRILELQKI FAPEILKIDEHFFARLNE-SFLV--LDEKK---Q	143
		SKYPI FATLE-EEKEYHKKFFPTIYHLRKKHLA	143
		SKYPI FATLA-EEKKYHKQFFPTIYHLRQQLA	144
		ERHVI FGNIA-DEVKYHKKEFPTIYHLRKKHLA	143
		SKYPI FATLV-EEKEYHKKFFPTIYHLRKKHLA	144
		SKYPI FATLA-EEKEYHKKFFPTIYHLRKKHLA	144
		SKHPI FGTLQ-EEKKYHKQFFPTIYHLRQQLA	143
		SKHPI FGTLQ-EEKEYHKQFFPTIYHLRQQLA	143
		SKHPI FGTLQ-EEKEYHKQFFPTIYHLRQQLA	143
		SKHPI FGTLQ-EEKKYHKQFFPTIYHLRQQLA	143
		ERHPI FGNIA-AEVKYHDDFFPTIYHLRKKHLA	143
		---LFNDKdyTDADYYEQYKTIYHLRYDLI	100
		DNYNLFIDEDfNDYTYHKYPTIYHLRKAIC	139
		ERHPI FGNIV-DEVAYHEKYPTIYHLRKKLV	143
		SKYPI F SNEK-EDKNYYDKYPTIYHLRKKDLA	142
		ERHPI FGNIV-DEVAYHEKYPTIYHLRKKLV	143
		DRHPI FGSLE-EEVAYHNTYPTIYHLRKKHLA	143
		HRHPI FAKLE-DEVAYHETYPTIYHLRKKLA	143
		HRHPI FAKLE-DEVAYHETYPTIYHLRKKLA	143

		HRHPI FAKLE-DEVAYHETYPTIYHLRKKLA	143
		HRHPI FAKLE-DEVAYHETYPTIYHLRKKLA	143
		HRHPI FAKLE-DEVAYHETYPTIYHLRKKLA	143
		HRHPI FAKLE-DEVAYHETYPTIYHLRKKLA	143
		HRHPI FAKLE-DEVAYHETYPTIYHLRKKLA	143
		HRHPI FAKLE-DEVAYHETYPTIYHLRKKLA	143
		SRHPVFATIK-QEKSYHQTYPTIYHLRQALA	143
		SRHPVFATIK-QEKSYHQTYPTIYHLRQALA	143
		SRHPVFATIK-QEKSYHQTYPTIYHLRQALA	143
		SRHPVFATIK-QEKSYHQTYPTIYHLRQALA	143

WP_002330729	74	RRRQRIILELQKI	FAPEILKIDEHFFARLNE-SFLV--LDEKK---	Q	SRHPVFATIK-QEKSYHQTYPTIYHLRQALA	143
WP_002335161	74	RRRQRIILELQKI	FAPEILKIDEHFFARLNE-SFLV--LDEKK---	Q	SRHPVFATIK-QEKSYHQTYPTIYHLRQALA	143
WP_002345439	74	RRRQRIILELQKI	FAPEILKIDEHFFARLNE-SFLV--LDEKK---	Q	SRHPVFATIK-QEKSYHQTYPTIYHLRQALA	143
WP_034867970	74	RRKYRLSKLQDL	FAEELCKQDDCFVRLNE-SFLV--PEEKQ---	Y	KPASIFFTLE-EEKEYYQKTYPTIYHLRQKLV	143
WP_047937432	74	RRRQRIILELQKI	FAPEILKIDEHFFARLNE-SFLV--LDEKK---	Q	SRHPVFATIK-QEKSYHQTYPTIYHLRQALA	143
WP_010720994	74	RRKYRLSKLQDL	FAEELCKQDDCFVRLNE-SFLV--PEEKQ---	Y	KPASIFFTLE-EEKEYYQKTYPTIYHLRQKLV	143
WP_010737004	74	RRKYRLSKLQDL	FAEELCKQDDCFVRLNE-SFLV--PEEKQ---	Y	KPASIFFTLE-EEKEYYQKTYPTIYHLRQKLV	143
WP_034700478	74	RRKYRLSKLQDL	FAEELCKQDDCFVRLNE-SFLV--PEEKQ---	Y	KPASIFFTLE-EEKEYYQKTYPTIYHLRQKLV	143
WP_007209003	74	RRKNRICYLQEI	FQPEMHLDNFFYRLNE-SFLVa--DDAK---	Y	DKHPIFGTLD-EEIHFEHQFPTIYHLRKYLA	143
WP_023519017	74	RRRQRLALQDI	FAEEIHKKDPNFARLEE-GDRV--EADKR--F		AKFPVFATLS-EEKNYHRQYPTIYHLRHDIA	143
WP_010770040	74	RRRNRI	CRQLQDLFTEEMNQVDANFFHRLQE-SFLV--PDEKE--F		ERHAI FGKME-EEVSYREFPTIYHLRKHLA	143
WP_048604708	74	RRRQRI	SYLQTFQEEEMNRIDPNFFNRLDE-SFLI--EEDKL--S		ERHPIFGTIE-EEVAYHKNYATIYHLRKELA	143
WP_010750235	74	RRKYRI	LELQKIFSEELKKDSSHFFARLDE-SFLI--PEDKQ---	Y	ARFPIFPTLL-EEKAYYQNYPTIYHLRQKLA	143
AI116583	113	RRKNRICYLQEI	FSNEMAKVDDSFHRLNE-SFLV--EEDKK--H		ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	182
WP_029073316	66	KRRERIRLLRGIMED	MVLDVDPFFIRLANVSFLD--QEDKKdy1K		SNYNLFIDKDFNDKTYDYKTYPTIYHLRKHLC	144
WP_031589969	66	KRRERIRLLREIMED	MVLDVDPFFIRLANVSFLD--QEDKKdy1K		SNYNLFIDKDFNDKTYDYKTYPTIYHLRKHLC	144
KDA45870	75	RRKNRLRYLQEI	FAPALAKVDPFFYRLNE-SSLVa--EDKK---	Y	DVYPIFGKRE-EELLYHDTHKTYPTIYHLRSELA	144
WP_039099354	62	RRKWRGLLREI	FEPYITPVDDFFLRKKQ-SNLS--PKDQR--K		-QTSLFNDRT--DRAFYDDTYPTIYHLRYKLM	132
AKP02966	65	RRKNRINWLNEI	FSEELANTDPSFLIRLQN-SWVSKDPDRK--R		DKYNLFIDNPTDKEYYREFPTIYHLRKELI	137
WP_010991369	74	RRNRISYLQGI	FAEEMSKTDANFFCRLSD-SFYV--DNEKR--N		SRHPFFATIE-EEVEYHKNYPTIYHLREELV	143
WP_033838504	74	RRNRISYLQGI	FAEEMSKTDANFFCRLSD-SFYV--DNEKR--N		SRHPFFATIE-EEVEYHKNYPTIYHLREELV	143
EHN60060	77	RRNRISYLQGI	FAEEMSKTDANFFCRLSD-SFYV--DNEKR--N		SRHPFFATIE-EEVEYHKNYPTIYHLREELV	146
EFR89594					-----	
WP_038409211	74	RRNRRIAYLQEI	FAAEMAENVANFFYRLDE-SFYI--ESEKR--H		SRHPFFATIE-EEVAYHEEYKTYPTIYHLREKLV	143
EFR95520					-----	
WP_003723650	74	RRNRISYLQEI	FAVEMANIDANFFCRLND-SFYV--DSEKR--N		SRHPFFATIE-EEVAYHDNYRTIYHLREKLV	143
WP_003727705	74	RRNRISYLQEI	FAVEMANIDANFFCRLND-SFYV--DSEKR--N		SRHPFFATIE-EEVAYHKNYRTIYHLREELV	143
WP_003730785	74	RRNRISYLQEI	FAVEMANIDANFFCRLND-SFYV--DSEKR--N		SRHPFFATIE-EEVAYHKNYRTIYHLREELV	143
WP_003733029	74	RRNRISYLQEI	FAIQMNEVDNFFNRLKE-SFYA--ESDKK---	Y	NRHPFFGTVE-EEVAYYKDFPTIYHLRKELI	143
WP_003739838	74	RRNRISYLQEI	FALEMANIDANFFCRLND-SFYV--DSEKR--N		SRHPFFATIE-EEVAYHKNYRTIYHLREELV	143
WP_014601172	74	RRNRISYLQEI	FAVEMANIDANFFCRLND-SFYV--DSEKR--N		SRHPFFATIE-EEVAYHKNYRTIYHLREELV	143
WP_023548323	74	RRNRISYLQEI	FAVEMANIDANFFCRLND-SFYV--DSEKR--N		SRHPFFATIE-EEVAYHKNYRTIYHLREELV	143
WP_031665337	74	RRNRISYLQEI	FAVEMANIDANFFCRLND-SFYV--DSEKR--N		SRHPFFATIE-EEVAYHKNYRTIYHLREELV	143
WP_031669209	74	RRNRISYLQEI	FAIQMNEVDNFFNRLKE-SFYA--ESDKK---	Y	NRHPFFGTVE-EEVAYYKDFPTIYHLRKELI	143

WP_033920898	74	RRNRISYLQEI FAVEMANIDANFFCRLND-SFYV--DSEKR---N	SRHPFFATIE-EEVAYHKNYRTIYHLREELV	143
AKI42028	77	RRNRISYLQEI FAVEMANIDANFFCRLND-SFYV--DSEKR---N	SRHPFFATIE-EEVAYHKNYRTIYHLREELV	146
AKI50529	77	RRNRISYLQEI FAVEMANIDANFFCRLND-SFYV--DSEKR---N	SRHPFFATIE-EEVAYHKNYRTIYHLREELV	146
EFR83390		-----	-----	
WP_046323366	74	RRNRISYLQEI FTAEMFEVDANFFYRLED-SFYI--ESEKR---Q	SRHPFFATIE-EEVAYHENYRTIYHLREKLV	143
AKE81011	90	RRKNRICYLQEI FSNEMAKVDDSFHRLEE-SFLV--EEDKK---H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	159
CUO82355	65	KRRERIRLLRAILQDMVLEKDPFFIRLEHtSFLD--EEDKakylG	DYNLFIDEDfNDYTYHHKYPTIYHLRkALC	143
WP_033162887	66	KRRERIRLLRDLLGDMVMEVDPTFFIRLLNvSFLD--EEDKQknlG	DYNLFIEKdfNDKTYDKYPTIYHLRkELC	144
AGZ01981	107	RRKNRICYLQEI FSNEMAKVDDSFHRLEE-SFLV--EEDKK---H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	176
AKA60242	74	RRKNRICYLQEI FSNEMAKVDDSFHRLEE-SFLV--EEDKK---H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	143
AKS40380	74	RRKNRICYLQEI FSNEMAKVDDSFHRLEE-SFLV--EEDKK---H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	143
4UN5_B	78	RRKNRICYLQEI FSNEMAKVDDSFHRLEE-SFLV--EEDKK---H	ERHPIFGNIV-DEVAYHEKYPTIYHLRKKLV	147

WP_010922251	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_039695303	145	DSSEKADLRLVYLALAHMIKFRGHFLIEGE-LNAENTDVQKI--FADFVGVYVRT--FDDS-H	LSEITVDVA--SI	212
WP_045635197	144	DSKEKTDLRLIYLALAHMIKYRGHFLYEEA-FDIKNNDIQKI--FNEFISIDYNT--FEGS-S	LSGQNAQVE--AI	211
5AXW A	105	EEEFSA-----ALLHLAKRRG--VHNV-----NEVE-----EDT--GN--	-----E-----	134
WP_009880683		-----	-----	
WP_010922251	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_011054416	144	DSTDKVDLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_011284745	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_011285506	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_011527619	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_012560673	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_014407541	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_020905136	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_023080005	144	DSTDKVDLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_023610282	144	DSTDKVDLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_030125963	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_030126706	144	DSTDKVDLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_031488318	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_032460140	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_032461047	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_032462016	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGG-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_032462936	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INANGVDAK--AI	211
WP_032464890	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_033888930	1	-----PDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_038431314	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	36
WP_038432938	144	DSTDKVDLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
WP_038434062	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
BAQ51233	55	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASRVDAK--AI	211
KGE60162		-----	-----	
KGE60856		-----	-----	
WP_002989955	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSDV ^D KL--FIQLVQTYNQL--FEEN--	INASGVDAK--AI	211
WP_003030002	144	DISQKADLRLVYLALAHMIKFRGHFLIEGQ-LKAENTNVQAL--FKDFVEYDKT--VEES-H	LSEMTVDAL--SI	211
WP_003065552	147	DSSEKADLRLVYLALAHMIKFRGHFLIEGE-LNAENTDVQKI--FADFVGVYVRT--FDDS-H	LSEITVDA--SI	214
WP_001040076	144	DKKEKADLRLVYLALAHMIIKFRGHFLIEDDrFDVVRTDIQKQ--YQAFLEIFDIT--FENN-D	LLSQDVDVE--AI	212

WP_001040078	144	DKQEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_001040080	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNIT--FENN-D	LLSQNVVDE--AI	212
WP_001040081	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTI--FENN-D	LLSQNVVDE--AI	212
WP_001040083	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040085	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040087	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040088	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040089	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040090	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040091	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040092	144	DKKEKADLRLVYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDTS--FENN-H	LLSQNVVDE--AI	212
WP_001040094	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVVDE--AI	212
WP_001040095	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVVDE--AI	212
WP_001040096	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVVDE--AI	212
WP_001040097	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVVDE--AI	212
WP_001040098	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVVDE--AI	212
WP_001040099	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVVDE--AI	212
WP_001040100	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVVDE--AI	212
WP_001040104	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040105	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_001040106	144	DKKEKANLRLVYIALAHIIKFRGHFLIEDDsFDVRNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_001040107	144	DKKEKADLRLVYIALAHIIKFRGHFLIEDDsFDVRNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_001040108	144	DKKEKADLRLVYIALAHIIKFRGHFLIEDDsFDVRNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_001040109	144	DKKEKANLRLVYIALAHIIKFRGHFLIEDDsFDVRNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_001040110	144	DKKEKANLRLVYIALAHIIKFRGHFLIEDDsFDVRNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_015058523	144	DKKEKADLRLVYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDTS--FENN-H	LLSQNVVDE--AI	212
WP_017643650	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDrFDVRNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVVDE--AI	212
WP_017647151	144	DKKEKADLRLFYIALAHIIKFRGHFLIEDDsFDVRNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_017648376	144	DKKEKADLRLFYIALAHIIKFRGHFLIEDDsFDVRNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_017649527	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
WP_017771611	144	DKKEKADLRLVYIALAHIIKFRGHFLIEDDsFDVRNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNVVDE--AI	212
WP_017771984	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
CFQ25032	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212
CFV16040	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVRNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVVDE--AI	212

KLJ37842	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
KLJ72361	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
KL220707	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
KL42645	144	DKKEKANLRLVYIALAHIIKFRGHFLIEDDsFDVrNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNI dVE--GI	212
WP_047207273	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
WP_047209694	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDIQKQ--YQAFLEIFDFTT--FENN-D	LLSQNVdVE--AI	212
WP_050198062	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
WP_050201642	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
WP_050204027	144	DKKEKANLRLVYIALAHIIKFRGHFLIEDDsFDVrNTDIQRQ--YQAFLEIFDFTT--FENN-H	LLSQNI dVE--GI	212
WP_050881965	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
WP_050886065	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
AHN30376	144	DKKEKADLRLVYIALAHIIKFRGHFLIEDDrFDVrNTDIQKQ--YQAFLEIFDTS--FENN-H	LLSQNVdVE--AI	212
EAO78426	144	DKKEKADLRLIYIALAHIIKFRGHFLIEDDsFDVrNTDISKQ--YQDFLEIFNTT--FENN-D	LLSQNVdVE--AI	212
CCW42055	144	DKKEKADLRLVYIALAHIIKFRGHFLIEDDrFDVrNTDIQKQ--YQAFLEIFDFTT--FENN-H	LLSQNVdVE--AI	212
WP_003041502	144	DISQKADLRLVYIALAHMIKFRGHFLIEGQ-LKAENTNVQAL--FKDFVEVYDKT--VEES-H	LSEITVDAL--SI	211
WP_0037593752	145	NSKEKADLRLVYIALAHMIKFRGHFLIEGQ-LKAENTDVQAL--FKDFVEEYDKT--IEES-H	LSEITVDAL--SI	212
WP_049516684	145	DISQKADLRLVYIALAHMIKFRGHFLIEGQ-LKAENTNVQAL--FKDFVEVYDKT--VEES-H	LSEMTVDAL--SI	212
GAD46167	144	NSKEKADLRLVYIALAHMIKFRGHFLIEGQ-LKAENTDVQAL--FKDFVEEYDKT--IEES-H	LSEITVDAL--SI	211
WP_018363470	145	DSTEKADLRLVYIALAHMIKFRGHFLIEGE-LNAENTDVQKL--FTDFVGVYDRT--FDDS-H	LSEITVDA--SI	212
WP_003043819	144	DSPEKADLRLIYIALAHIIKFRGHFLIEGK-LNAENSDVAKL--FYQLIQTYNQL--FEES--	LDEIEVDK--GI	211
WP_006269658	144	DTSKKADLRLVYIALAHMIKFRGHFLIEGD-LKAENTDVQAL--FKDFVEEYDKT--IEES-H	LSEITVDAL--SI	211
WP_048800889	144	DSTGKADLRLVYIALAHMIKFRGHFLIEGQ-LKAENTDVQTL--FNDFVEVYDKT--IEES-H	LAEITVDAL--SI	211
WP_012767106	144	DSTDKADLRLIYIALAHMIKFRGHFLIEGD-LNPDNSDMDKL--FIQLVQTYNQL--FEEN--	INASRVdAK--AI	211
WP_014612333	144	DSTDKADLRLIYIALAHMIKFRGHFLIEGD-LNPDNSDVQKL--FIQLVQTYNQL--FEEK--	INASGVdAK--AI	211
WP_015017095	144	DSTDKADLRLIYIALAHMIKFRGHFLIEGD-LNPDNSDMDKL--FIQLVQTYNQL--FEEN--	INASRVdAK--AI	211
WP_015057649	144	DSTDKADLRLIYIALAHMIKFRGHFLIEGD-LNPDNSDMDKL--FIQLVQTYNQL--FEEN--	INASRVdAK--AI	211
WP_048327215	144	DSTDKADLRLIYIALAHMIKFRGHFLIEGD-LNPDNSDMDKL--FIQLVQTYNQL--FEEN--	INASRVdAK--AI	211
WP_049519324	144	DSTDKADLRLIYIALAHMIKFRGHFLIEGD-LNPDNSDVQKL--FIQLVQTYNQL--FEEN--	INASRVdAK--AI	211
WP_012515931	144	DNPQKADLRLIYI AVAHIIKFRGHFLIEGT-LSSKNNLQKS--FDHLVDTYNLL--FEEQ--	LLTEGINAK--EL	211
WP_021320964	144	DNPQKADLRLIYI AVAHIIKFRGHFLIEGT-LSSKNNLQKS--FDHLVDTYNLL--FEEQ--	LLTEGINAK--EL	211
WP_037581760	144	DNPQKADLRLIYI AVAHIIKFRGHFLIEGT-LSSKNNLQKS--FDHLVDTYNLL--FEEQ--	LLTEGINAK--EL	211
WP_004232481	144	DSPEKADLRLVYIALAHMIKFRGHFLIEGQ-LNAENTDVQKI--FADFVGVYDRT--FDDS-H	LSEITVDA--SI	211
WP_009854540	145	DSSEKADLRLVYIALAHMIKYRGHFLIEGK-LNAENTDVQKL--FTDFVGVYDRT--FDDS-H	LSEITVDVA--ST	212
WP_012962174	145	DSHEKADLRLIYIALAHMIKFRGHFLIEGE-LNAENTDVQKL--FEAFVEVYDRT--FDDSS-N	LSEITVDAS--SI	212

WP_039695303	145	DSSEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKI--FADFVGVYVRT--FDDS-H	DSSEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKI--FADFVGVYVRT--FDDS-H	LSEITVDVA---SI	211
WP_014334983	144	DSQEKADLRVYLALAHMIKYRGHFLIEGE-LNAENTDVQKL--FNVFVETDKI--VDES-H	DSQEKADLRVYLALAHMIKYRGHFLIEGE-LNAENTDVQKL--FNVFVETDKI--VDES-H	LSEIEVDAS---SI	211
WP_003099269	144	DSDQKADLRVYLALAHMIKFRGHFLIEGN-LDSENTDVHVL--FLNLVNIYNNL--FEED--	DSDQKADLRVYLALAHMIKFRGHFLIEGN-LDSENTDVHVL--FLNLVNIYNNL--FEED--	VETASIDAE---KI	211
AHY15608	144	DSDQKADLRVYLALAHMIKFRGHFLIEGN-LDSENTDVHVL--FLNLVNIYNNL--FEED--	DSDQKADLRVYLALAHMIKFRGHFLIEGN-LDSENTDVHVL--FLNLVNIYNNL--FEED--	VETASIDAE---KI	211
AHY17476	144	DSDQKADLRVYLALAHMIKFRGHFLIEGN-LDSENTDVHVL--FLNLVNIYNNL--FEED--	DSDQKADLRVYLALAHMIKFRGHFLIEGN-LDSENTDVHVL--FLNLVNIYNNL--FEED--	VETASIDAE---KI	211
ESR09100					
AGM98575	144	DSDQKADLRVYLALAHMIKFRGHFLIEGN-LDSENTDVHVL--FLNLVNIYNNL--FEED--	DSDQKADLRVYLALAHMIKFRGHFLIEGN-LDSENTDVHVL--FLNLVNIYNNL--FEED--	VETASIDAE---KI	211
ALF27331	144	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_018372492	144	DTPDKMDIRLYLALAHMIKYRGHFLIEGD-LDIENIGIQDS--FKSFIEEYNTQ--FGTK--	DTPDKMDIRLYLALAHMIKYRGHFLIEGD-LDIENIGIQDS--FKSFIEEYNTQ--FGTK--	-LDSTTKVE---AI	209
WP_045618028	145	DSKEKADFRLIYLALAHMIKYRGHFLIEES-FDIKNNDIQKI--FNEFISIYDNT--FEGS-S	DSKEKADFRLIYLALAHMIKYRGHFLIEES-FDIKNNDIQKI--FNEFISIYDNT--FEGS-S	LNGQNAQVE---AI	212
WP_045635197	144	DSKEKTDLRVYLALAHMIKYRGHFLIEEA-FDIKNNDIQKI--FNEFISIYDNT--FEGS-S	DSKEKTDLRVYLALAHMIKYRGHFLIEEA-FDIKNNDIQKI--FNEFISIYDNT--FEGS-S	LSGQNAQVE---AI	211
WP_002263549	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002263887	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002264920	144	DSTEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	DSTEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	LSEITVDAS---SI	211
WP_002269043	144	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002269448	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002271977	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002272766	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002273241	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002275430	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002276448	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002277050	144	DSTEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	DSTEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	LSEITVDAS---SI	211
WP_002277364	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002279025	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002279859	144	DSTEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	DSTEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	LSEITVDAS---SI	211
WP_002280230	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002281696	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002282247	144	DSTEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	DSTEKADLRVYLALAHMIKFRGHFLIEGE-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	LSEITVDAS---SI	211
WP_002282906	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002283846	144	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002287255	144	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKVDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002288990	144	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002289641	144	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002290427	144	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	DNPEKTDLRVYLALAHMIKFRGHFLIEGK-FDTRNDVQRL--FQEFFLAVYDNT--FENS-S	LQEQNVQVE---EI	211

WP_002295753	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002296423	144	DNPEKTDLRLVYLALAHIIKFGGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002304487	144	NSTEKADLRLVYLSLAHMIKFRGHFLIEGQ-LKAENTNVQAL--FKDFVEVYDKT--VEES-H	LSEMTVDAL---SI	211
WP_002305844	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002307203	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQKL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002310390	144	DNPEKTDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_002352408	144	DNPEKTDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_012997688	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_014677909	144	DNPEKTDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_019312892	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_019313659	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_019314093	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQKL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_019315370	144	DNPEKTDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_019803776	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_019805234	144	DNPEKTDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_024783594	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_024784288	144	DSTEKADLRLVYLALAHMIKFRGHFLIEGK-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	LSEITVDAS---SI	211
WP_024784666	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_024784894	144	DNPEKTDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_024786433	144	DSTEKADLRLVYLALAHMIKFRGHFLIEGK-LNAENTDVQKL--FADFVGVYDRT--FDDS-H	LSEITVDAS---SI	211
WP_049473442	144	DNPEKVDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
WP_049474547	144	DNPEKTDLRLVYLALAHIIKFRGHFLIEGK-FDTRNNDVQRL--FQEFLLAVYDNT--FENS-S	LQEQNVQVE---EI	211
EMC03581	137			
WP_000428612	145	DSKEKTDLRLIYLALAHMIKYRGHFLYEDT-FDIKNNDIQKI--FNEFISIYDNT--FEGN-S	LSGQNVQVE---AI	212
WP_000428613	145	DSKEKTDLRLIYLALAHMIKYRGHFLYEDT-FDIKNNDIQKI--FSEFISIYDNT--FEGS-S	LSGQNAQVE---AI	212
WP_049523028	144	DSKEKVDLRLIYLALAHIIKFRGHFLYEDS-FDIKNNDIQKI--FNEFTIYDNT--FEES-S	LSKQNAQVE---EI	211
WP_003107102	113	DSDEKADLRLIYLALAHIIKFRGHFLIEGD-LDSQNTDVNAL--FLKLVDTYNLM--FEDD--	IDTQITIDAT---VI	180
WP_054279288	146	DNTEKADLRLIYLALAHIIKFRGHFLIEGA-LSANNTDVQKL--VHALVDAYNIM--FEED--	LDIEAIDVK---AI	213
WP_049531101	145	DSKEKADLRLIYLTLAHMIKYRGHFLYEEES-FDIKNNDIQKI--FNEFISIYDNT--FEGS-S	LSGQNAQVE---AI	212
WP_049538452	145	DSKEKADLRLIYLALAHMIKYRGHFLYEEA-FDIKNNDIQKI--FNEFISIYDNT--FEGS-S	LSGQNEQVE---AI	212
WP_049549711	145	DSKEKADLRLIYLVLAHMIKYRGHFLYEEA-FDIKNNDIQKI--FNEFISIYDNT--FEGS-S	LSGQNAQVE---TI	212
WP_007896501	146	DRDQKADLRLIYLALSHIIKFRGHFLIEGK-LNSENTDVQKL--FIALVTVYNLL--FEEE--	IAGETCDAK---AL	213
EFR44625	98	DRDQKADLRLIYLALSHIIKFRGHFLIEGK-LNSENTDVQKL--FIALVTVYNLL--FEEE--	IAGETCDAK---AL	165
WP_002897477	144	DSKEKSDVRLIYLALAHMIKYRGHFLYEEET-FDIKNNDIQKI--FNEFINIYDNT--FEGS-S	LSGQNAQVE---AI	211

WP_002906454	144	DSKEKTDLRLIYLALAHMIKYRGHFLYEES-FDIKNNDIQKI--FNEFISIDYNT--FEGS-S	LSGQNAQVE---AI	211
WP_009729476	145	DSKEKTDLRLIYLALAHMIKYRGHFLYEES-FDIKNNDIQKI--FNEFISIDYNT--FEGN-S	LSGQNVQVE---AI	212
CQR24647	144	DSSEKADLRLIYLALAHMIKYRGHFLIIEG-FIDIRNMNSQNL--FKEFLLAFDGI--QVDC-Y	LASKHTDIS---GI	211
WP_000066813	145	DSKEKTDLRLIYLALAHMIKYRGHFLYEES-FDIKNNDIQKI--FSEFISIDYNT--FEGK-S	LSGQNAQVE---AI	212
WP_009754323	145	DSKEKADLRLIYLALAHMIKYRGHFLYEES-FDIKNNDIQKI--FNEFINIYDNT--FEGS-S	LSGQNAQVE---AI	212
WP_044674937	144	DSSQKADIRLIYLALAHMIKYRGHFLFEGD-LKSENKDVQHL--FNDFVEMFDKT--VEGS-Y	LSENLPNVA---DV	211
WP_044676715	144	DSSQKADIRLIYLALAHMIKYRGHFLFEGD-LKSENKDVQHL--FNDFVEMFDKT--VEGS-Y	LSENLPNVA---DV	211
WP_044680361	144	DSSQKADIRLIYLALAHMIKYRGHFLFEGD-LKSENKDVQHL--FNDFVEMFDKT--VEGS-Y	LSENLPNVA---DV	211
WP_044681799	144	DSSQKADIRLIYLALAHMIKYRGHFLFEGD-LKSENKDVQHL--FNDFVEMFDKT--VEGS-Y	LSENLPNVA---DV	211
WP_049533112	144	DISQKADLRLIYLALAHMIKFRGHFLIEGQ-LKAENTNVQAL--FKDFVEVYDKT--VEES-H	LSEMTVDAL---SI	211
WP_029090905	101	SQHRQFDIREVYLAIHHLIKYRGHFIYEDQTFITDGNQLQHH--IKAIITMINST1--NR--	IIPETIDINVFEKI	171
WP_006506696	140	ESTEKADPRLIYLALAHMIKYRGHFLIIEGQ-FNMDASNIEDK--LSDIFTQFTSFnnIPEdD	--KKNLEIL---EI	210
AIT42264	144	DSTDKADLRLIYLALAHMIKFRGHFLIIEGD-LNPDNSDVKL--FIQLVQTYNQL--FEEN--	INASGVDAK---AI	211
WP_034440723	143	DSNQKADLRLIYLALAHMIKYRGHFLIIEGD-LKMDGISISES--FQEFIDSYNEVcaLEDE-N	NDELLTQIE---NI	217
AKQ21048	144	DSTDKADLRLIYLALAHMIKFRGHFLIIEGD-LNPDNSDVKL--FIQLVQTYNQL--FEEN--	INASGVDAK---AI	211
WP_004636532	144	DNPEKADLRLIYLALAHMIKYRGHFLIIEGE-LNNTENTSISET--FEQFLDTYSDI--FKEQ--	LVGDISKVE---EI	210
WP_002364836	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_016631044	95	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENTSVKDQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	168
EMS75795				
WP_002373311	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENTSVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_002378009	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_002407324	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_002413717	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_010775580	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_010818269	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_010824395	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENTSVKDQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_016622645	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEK--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_033624816	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKDQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_033625576	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_033789179	144	DSSEQADLRLIYLALAHMIKYRGHFLIIEGK-LSTENISVKEQ--FQQFMIYYNQT--FVNGes	PLPESVLIIE---EE	217
WP_002310644	144	DSSEKADIRLIVYLAHMAHLLKYRGHFLIIEGE-LNTENSSVTEI--FRQFLSTYNQO--FSEA-D	KLDEAVDCS---FV	216
WP_002312694	144	DSSEKADIRLIVYLAHMAHLLKYRGHFLIIEGE-LNTENSSVTEI--FRQFLSTYNQO--FSEA-G	KLDEAVDCS---FV	216
WP_002314015	144	DSSEKADIRLIVYLAHMAHLLKYRGHFLIIEGE-LNTENSSVTEI--FRQFLSTYNQO--FSEA-D	KLDEAVDCS---FV	216
WP_002320716	144	DSSEKADIRLIVYLAHMAHLLKYRGHFLIIEGE-LNTENSSVTEI--FRQFLSTYNQO--FSEA-D	KLDEAVDCS---FV	216

WP_002330729	144	DSSEKADIRLVYLAMAHLLKYRGHFLIEGE-LNTENSSVTET--FRQFLSTYNQO--FSEA-D	KLDEAVDCS---FV	216
WP_002335161	144	DSSEKADIRLVYLAMAHLLKYRGHFLIEGE-LNTENSSVTET--FRQFLSTYNQO--FSEA-D	KLDEAVDCS---FV	216
WP_002345439	144	DSSEKADIRLVYLAMAHLLKYRGHFLIEGE-LNTENSSVTET--FRQFLSTYNQO--FSEA-D	KLDEAVDCS---FV	216
WP_034867970	144	DSTEKEDRLVYLALAHLLKYRGHFLFEGD-LDTENTSIIEES--FRVFLQYKQ--SDQP--	-LIVHQPVL---TI	209
WP_047937432	144	DSSEKADIRLVYLAMAHLLKYRGHFLIEGE-LNTENSSVTET--FRQFLSTYNQO--FSEA-D	KLDEAVDCS---FV	216
WP_010720994	144	DSTEKEDRLVYLALAHLLKYRGHFLFEGD-LDTENTSIIEES--FRVFLQYKQ--SDQP--	-LIVHQPVL---TI	209
WP_010737004	144	DSTEKEDRLVYLALAHLLKYRGHFLFEGD-LDTENTSIIEES--FRVFLQYKQ--SDQP--	-LIVHQPVL---TI	209
WP_034700478	144	DSTEKEDRLVYLALAHLLKYRGHFLFEGD-LDTENTSIIEES--FRVFLQYKQ--SDQP--	-LIVHQPVL---TI	209
WP_007209003	144	DGDEKADRLVYLALAHLLKYRGHFLIEGE-LNTENSSVTET--FRVFLQYKQ--SDQP--	FIDESIDFS---EV	214
WP_023519017	144	NSKEQADIRLVYLALAHLLKYRGHFLFEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	-IGLDLDDAV---PI	209
WP_010770040	144	DTSEQADRLVYLALAHLLKYRGHFLIEGE-LNTENSSVTET--FRVFLQAYQOQ--FPEP--	PLAVPDNIE---EL	212
WP_048604708	144	DAEEKADRLVYLALAHLLKYRGHFLIEGE-LNTENSSVTET--FRVFLQAYQOQ--FPEP--	PVDETSIG---SI	208
WP_010750235	144	DSTEKADIRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	-IIFYKDIP---LI	209
AI116583	183	DSTDKADRLVYLALAHLLKYRGHFLIEGE-LNTENSSVTET--FRVFLQAYQOQ--FPEP--	INASGVDAK---AI	250
WP_029073316	145	ESKEKEDPRLIYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	--KKIDEVL---NV	215
WP_031589969	145	ESKEKEDPRLIYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	--KKIDEVL---NV	215
KDA45870	145	NDRPADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	----KLDVA---DI	209
WP_039099354	133	TEKRQFDIREIYLAMHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	-TDKLAEVK---AL	206
AKP02966	138	INKNKADIRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	-FPDNCWWhi sDI	208
WP_010991369	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KLEDNKDVA---KI	217
WP_033838504	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KLEDNKDVA---KI	217
EHN60060	147	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KLEDNKDVA---KI	220
EFR89594		-----	-----	
WP_038409211	144	NSSDKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	RLEENKEVA---EI	217
EFR95520		-----	-----	
WP_003723650	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KVEENIEVA---NI	217
WP_003727705	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KVEENTEVA---SI	217
WP_003730785	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KVEENTEVA---SI	217
WP_003733029	144	DSQKKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KTEKNQEVA---QI	217
WP_003739838	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KMEENTTVA---DI	217
WP_014601172	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KVEENIEVA---NI	217
WP_023548323	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KVEENIEVA---NI	217
WP_031665337	144	NSSEKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KVEENIEVA---NI	217
WP_031669209	144	DSQKKADRLVYLALAHLLKYRGHFLIEGE-LDTENTSVTEN--YQFLQAYQOQ--FPEP--	KTEKNQEVA---QI	217

WP_033920898	144	NSSEKADLRLVYLALAHIIKYRGNFLIEGA-LDTKNTSVDGV--YEQFIQTYNQV--FMSNiE	KVEENIEVA---NI	217
AKI42028	147	NSSEKADLRLVYLALAHIIKYRGNFLIEGA-LDTKNTSVDGV--YEQFIQTYNQV--FMSNiE	KVEENIEVA---NI	220
AKI50529	147	NSSEKADLRLVYLALAHIIKYRGNFLIEGA-LDTKNTSVDGV--YEQFIQTYNQV--FMSNiE	KVEENIEVA---NI	220
EFR83390		-----	-----	
WP_046323366	144	NSSDKADLRLVYLALAHIIKYRGNFLIEGK-LDTKNTSVDEV--FKQFIKTYNQV--FASDiE	RIENNEVA---KI	217
AKE81011	160	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSVDVKL--FIQLVQTYNQV--FEEN--	INASGVDAK---AI	227
CUO82355	144	ESTEKADPRLIYLALHHIVKYRGNFLYEGQkFNMDASNIEDK--LSDVFTQFADFnnIPYEdD	--KKNLEIL---EI	214
WP_033162887	145	ENKEKADPRLIYLALHHIVKYRGNFLYEGQsFTMDNSDI EER--LNSAI EKFM SIneFDNRiV	--SDINSMI---AV	215
AGZ01981	177	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSVDVKL--FIQLVQTYNQV--FEEN--	INASGVDAK---AI	244
AKA60242	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSVDVKL--FIQLVQTYNQV--FEEN--	INASGVDAK---AI	211
AKS40380	144	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSVDVKL--FIQLVQTYNQV--FEEN--	INASGVDAK---AI	211
4UN5_B	148	DSTDKADLRLIYLALAHMIKFRGHFLIEGD-LNPDNSVDVKL--FIQLVQTYNQV--FEEN--	INASGVDAK---AI	215

WP_010922251	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_039695303	213	LTEK-ISKSRRLLENLIKY-Y-PT	EKKNTLFGNLIALALGLQPNFKTNF--KLSED-A--KLQ--FSKDTYEEDLEE	278
WP_045635197	212	FTDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLIIVGNQADFKKHF--DLEDK-A--PLQ--FSKDTYDEDLEN	277
5AXW A	135	LSTK-----EQISRN-S-K	-----LEEKyVa--ELQ-----	157
WP_009880683		-----	-----	
WP_010922251	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_011054416	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_011284745	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_011285506	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_011527619	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_012560673	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_014407541	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_020905136	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_023080005	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_023610282	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_030125963	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_030126706	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_031488318	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_032460140	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_032461047	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_032462016	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_032462936	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_032464890	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_033888930	37	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	102
WP_038431314	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_038432938	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_038434062	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
BAQ51233	123	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	188
KGE60162		-----	-----	
KGE60856		-----	-----	
WP_002989955	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A--KLQ--LSKDTYDDDLN	277
WP_003030002	212	LTEK-VSKSRRLLENLIAH-Y-PA	EKKNTLFGNLIALSGLQPNFKTNF--QLSED-A--KLQ--FSKDTYEEDLEG	277
WP_003065552	215	LTEK-ISKSRRLLENLIKY-Y-PT	EKKNGLFGNLIALSGLQPNFKTNF--KLSED-A--KLQ--FSKDSYEEDLGE	280
WP_001040076	213	LTDK-ISKSAKKDRILIAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKHF--NLEDK-T--PLQ--FAKDSYDEDLEN	278

KLJ37842	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
KLJ72361	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
KL20707	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
KL242645	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
WP_047207273	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
WP_047209694	213	LTDK-ISKSAKKDRILAR-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
WP_050198062	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
WP_050201642	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
WP_050204027	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
WP_050881965	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
WP_050886065	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
AHN30376	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
EA078426	213	LTDK-ISKSAKKDRILAQ-Y-PN	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
CCW42055	213	LTDK-ISKSAKKDRILAQ-Y-PD	QKSTGIFAEFLKLIIVGNQADFKKYF--NLEDK-T---	PLQ--FAKDSYEDLEEN	278
WP_003041502	212	LTEK-VSKSRRLLENLIAH-Y-PA	EKKNTLFGNLIALSGLQPNFKTNF--QLESD-A--	KLQ--FSKDTYEEDLEG	277
WP_0037593752	213	LTEK-VSKSRRLLENLIAH-Y-PT	EKKNTLFGNLIALSGLQPNFKTNF--QLESD-A--	KLQ--FSKDTYEEDLEE	278
WP_049516684	213	LTEK-VSKSRRLLENLVEC-Y-PT	EKKNTLFGNLIALSGLQPNFKTNF--QLESD-A--	KLQ--FSKDTYEEDLEG	277
GAD46167	212	LTEK-VSKSRRLLENLIAH-Y-PT	EKKNTLFGNLIALSGLQPNFKTNF--QLESD-A--	KLQ--FSKDTYEEDLEE	278
WP_018363470	213	LTEK-VSKSRRLLENLIAH-Y-PT	EKKNTLFGNLIALSGLQPNFKTNF--QLESD-A--	KLQ--FSKDTYEEDLEE	278
WP_003043819	212	LTEK-VSKSRRLLENLINN-Y-PK	EKKNTLFGNLIALSGLQPNFKTNF--QLESD-A--	KLQ--FSKDTYEEDLEE	277
WP_006269658	212	LTEK-VSKSRRLLENLIAH-Y-PT	EKKNTLFGNLIALSGLQPNFKTNF--QLESD-A--	KLQ--FSKDTYEEDLEG	277
WP_048800889	212	LTEK-VSKSRRLLENLIVK-Y-PT	EKKNTLFGNLIALSGLQPNFKTNF--QLESD-A--	KLQ--FSKDTYEEDLEE	277
WP_012767106	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKRNGLFGNLIALSGLQPNFKSNF--DLAED-A--	KLQ--LSKDTYDDDLN	277
WP_014612333	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKRNGLFGNLIALSGLQPNFKSNF--DLAED-A--	KLQ--LSKDTYDDDLN	277
WP_015017095	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKRNGLFGNLIALSGLQPNFKSNF--DLAED-A--	KLQ--LSKDTYDDDLN	277
WP_015057649	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKRNGLFGNLIALSGLQPNFKSNF--DLAED-A--	KLQ--LSKDTYDDDLN	277
WP_048327215	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKRNGLFGNLIALSGLQPNFKSNF--DLAED-A--	KLQ--LSKDTYDDDLN	277
WP_049519324	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKRNGLFGNLIALSGLQPNFKSNF--DLAED-A--	KLQ--LSKDTYDDDLN	277
WP_012515931	212	LSAA-LSKSRRLLENLISL-I-PG	QKKTGIFGNIIALSGLQPNFKANF--GLSKD-V--	KLQ--LAKDTYADDLDS	277
WP_021320964	212	LSAA-LSKSRRLLENLISL-I-PG	QKKTGIFGNIIALSGLQPNFKANF--GLSKD-V--	KLQ--LAKDTYADDLDS	277
WP_037581760	212	LSAA-LSKSRRLLENLISL-I-PG	QKKTGIFGNIIALSGLQPNFKANF--GLSKD-V--	KLQ--LAKDTYADDLDS	277
WP_004232481	212	LTEK-ISKSRRLLENLIKQ-Y-PT	EKKNTLFGNLIALSGLQPNFKTNF--KLESD-A--	KLQ--FSKDTYEEDLEE	277
WP_009854540	213	LTEK-ISKSRRLLENLIKQ-Y-PT	EKKNTLFGNLIALSGLQPNFKTNF--KLESD-A--	KLQ--FSKDTYEEDLEE	278
WP_012962174	213	LTEK-FSKSRRLLENLIKQ-Y-PT	EKKNTLFGNLIALSGLQPNFKTSF--KLESD-A--	KLQ--FSKDTYEEDLEE	278

WP_039695303	213	LTEK-ISKRRLENLIKY-Y-PT	EKKNTLFGNLIALALGLQPNFKTNF--KLS	277
WP_014334983	212	LTEK-VSKRRLENLIKQ-Y-PT	EKKNTLFGNLIALALGLQPNFKTNF--KLS	277
WP_003099269	212	LTSK-TSKRRLENLIAE-I-PN	QKRNMFLGNLVSALGLTPNFKTNF--ELLE	277
AHY15608	212	LTSK-TSKRRLENLIAE-I-PN	QKRNMFLGNLVSALGLTPNFKTNF--ELLE	277
AHY17476	212	LTSK-TSKRRLENLIAE-I-PN	QKRNMFLGNLVSALGLTPNFKTNF--ELLE	277
ESR09100		-----	-----	
AGM98575	212	LTSK-TSKRRLENLIAE-I-PN	QKRNMFLGNLVSALGLTPNFKTNF--ELLE	277
ALF27331	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_018372492	210	FTEN-SSKAKRVEITLGL-F-PD	ETAAGNLDKFLKMLGNQADFKKVF--DLEE	275
WP_045618028	213	FTDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHF--DLEE	278
WP_045635197	212	FTDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHF--DLEE	277
WP_002263549	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002263887	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002264920	212	LTEK-ISKRRLEKLINN-Y-PK	EKKNTLFRNLVALSLGLQPNFKTNF--KLS	277
WP_002269043	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002269448	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002271977	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002272766	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002273241	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002275430	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002276448	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002277050	212	LTEK-ISKRRLEKLINN-Y-PK	EKKNTLFGNLIALSLGLQPNFKTNF--KLS	277
WP_002277364	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002279025	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002279859	212	LTEK-ISKRRLEKLINN-Y-PK	EKKNTLFGNLIALSLGLQPNFKTNF--KLS	277
WP_002280230	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002281696	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002282247	212	LTEK-ISKRRLEKLINN-Y-PK	EKKNTLFGNLIALSLGLQPNFKTNF--KLS	277
WP_002282906	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002283846	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002287255	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002288990	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002289641	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277
WP_002290427	212	LTDK-ISKSAKKDRVLKL-F-PN	EKSNGRFAEFLKLI VGNQADFKKHF--ELE	277

WP_002295753	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_002296423	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDIYEEEEEV	277
WP_002304487	212	LTEK-VSKRRLENLVECY-PT	EKKNTLFGNLI ALSGLQPNFKTF--QISED-A---KLQ--FSKDTYEEEDLEG	277
WP_002305844	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_002307203	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--LSKDTYEEEEEV	277
WP_002310390	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDIYEEEEEV	277
WP_002352408	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDIYEEEEEV	277
WP_012997688	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_014677909	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDIYEEEEEV	277
WP_019312892	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_019313659	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_019314093	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--LSKDTYEEEEEV	277
WP_019315370	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_019803776	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-V---PLQ--FSKDTYEEEEEV	277
WP_019805234	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDIYEEEEEV	277
WP_024783594	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_024784288	212	LTEK-ISKRRLEKLNIN-Y-PK	EKKNTLFGNLI ALSGLQPNFKTF--KLESD-A---KLQ--FSKDTYEEEDLEE	277
WP_024784666	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_024784894	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
WP_024786433	212	LTEK-ISKRRLEKLNIN-Y-PK	EKKNTLFGNLI ALSGLQPNFKTF--KLESD-A---KLQ--FSKDTYEEEDLEE	277
WP_049473442	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEDLEE	277
WP_049474547	212	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	277
EMC03581	205	LTDK-ISKSAKKDRVCLKL-F-PN	EKSNRFAEFLKLI VGNQADFKKHF--ELEEK-A---PLQ--FSKDTYEEEEEV	270
WP_000428612	213	FTDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHF--DLEEK-A---PLQ--FSRDTYDEEDLEN	278
WP_000428613	213	FTDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHF--DLGEK-A---PLQ--FSKDTYDEEDLEN	278
WP_049523028	212	FTDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHF--DLEEK-A---PLQ--FSKDTYEEEDLES	277
WP_003107102	181	LTEK-MSKRRLENLIAK-I-PN	QKKNTLFGNLI SLSGLTPNFKANF--ELSEDA---KLQ--ISKESFEEDLDN	246
WP_054279288	214	LTEK-ISKTRRLENLISN-I-PG	QKKNGLFGNLI ALSGLTPNFKSHF--NLPEDA---KLQ--LAKDTYDEEELNN	279
WP_049531101	213	FTDK-ISKTKRERVLKL-F-PD	QKSTGLFSEFLKLI VGNQADFKKHF--DLEEK-A---PLQ--FSKDTYDEEDLEN	278
WP_049538452	213	FSDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHF--DLEEK-A---PLQ--FSKDTYDEEDLEN	278
WP_049549711	213	FTDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHF--DLGEK-A---PLQ--FSKDTYDEEDLEN	278
WP_007896501	214	LTAK-TSKSKRLESLISE-F-PG	QKKNGLFGNLI ALSGLRPNFKSNF--GLSEDA---KLQ--ITKDTYEEEELDN	279
EFR44625	166	LTAK-TSKSKRLESLISE-F-PG	QKKNGLFGNLI ALSGLRPNFKSNF--GLSEDA---KLQ--ITKDTYEEEELDN	231
WP_002897477	212	FTDK-ISKSAKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHF--DLEEK-A---PLQ--FSKDTYDEEELN	277

WP_002906454	212	FTDK-ISKSTKRERVLKL-F-SD	EKSTGLFSEFLKLI VGNQADFKKHf--DLEEK-A---PLQ--FSKDTYDEDLEN	277
WP_009729476	213	FTDK-ISKSARERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHf--DLEEK-A---PLQ--FSRDTYDEDLEN	278
CQR24647	212	ITAK-ISKSRKVEAVLEQ-F-PD	QKNSFFGNMVSFLGMPNFKSNf--ELDED-A---KLQ--FSRDSYDEDLEN	277
WP_000066813	213	FTDK-ISKSTKRERVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHf--DLEEK-A---PLQ--FSKDTYDEDLEN	278
WP_009754323	213	FTGK-ISKSVKREHVLKL-F-PD	EKSTGLFSEFLKLI VGNQADFKKHf--DLEEK-A---SLQ--FSKDTYDEDLEN	278
WP_044674937	212	LVEK-VKSRRLLENILHY-F-PN	EKKNGLFGNFI ALALGLQPNFKTNf--ELAED-A---KIQ--FSKETEEEDLEE	277
WP_044676715	212	LVEK-VKSRRLLENILHY-F-PN	EKKNGLFGNFI TALGLQPNFKTNf--ELAED-A---KIQ--FSKETEEEDLEE	277
WP_044680361	212	LVEK-VKSRRLLENILHY-F-PN	EKKNGLFGNFI ALALGLQPNFKTNf--ELAED-A---KIQ--FSKETEEEDLEE	277
WP_044681799	212	LVEK-VKSRRLLENILHY-F-PN	EKKNGLFGNFI ALALGLQPNFKTNf--ELAED-A---KIQ--FSKETEEEDLEE	277
WP_049533112	212	LTEK-VKSRRLLENILIAH-Y-PA	EKKNLTFGNLIALSLGLQPNFKTNf--QLSED-A---KLQ--FSKDTYEEEDLEG	277
WP_029090905	172	LlDRmMNRSSKVKFLIEL--TG	KQDKPLLKELFNLI VGLKAKPASIFe--QENlativETM-nMSTEQVQLDLIT	243
WP_006506696	211	LKKP-LSKKAKVDEVMTL-IaPE	KDYKSAFKELAVTGIAGNKMNVTKMIlcEPIKQ-Gds-EIKlkFSDSNYDDQFSE	283
AIT42264	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLI ALSLGLTPNFKSNf--DLAED-A---KLQ--LSKDTYDDDLDN	277
WP_034440723	218	FKQD-ISRSKKLDAQIAL-F-QG	-KRQSLFGIFLTLIVGNKANFQKIF--NLEDD---iKLD--LKEEDYDENLEE	283
AKQ21048	212	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLI ALSLGLTPNFKSNf--DLAED-A---KLQ--LSKDTYDDDLDN	277
WP_004636532	211	LSSK-QSRSRKHEQIMAL-F-PN	ENKLNFGFRFMMLIVGNTSNFKPVf--DLDEE-Y---KlK--LSDETYEEDLDT	276
WP_002364836	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_016631044	169	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	234
EMS75795	1	-----	-----MDEE-A---KIQ--LSKESYEEEEELES	20
WP_002373311	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_002378009	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_002407324	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_002413717	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_010775580	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KIKItYASESYEEDLEG	285
WP_010818269	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_010824395	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_016622645	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_033624816	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_033625576	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_033789179	218	LTEK-ASRTKKSEKVLQQ-F-PQ	EKANGLFGQFLKLMVGNKADFKKVf--GLEEE-A---KI--tYASESYEEDLEG	283
WP_002310644	217	FTEK-MSKTKKAEtLLKY-F-PH	EKSNGLSQFIKLMVGNQGNFKNVf--GL-EE-A---KLQ--FSKETEEEDLEE	281
WP_002312694	217	FTEK-MSKTKKAEtLLKY-F-PH	EKSNGLSQFIKLMVGNQGNFKNVf--GL-EEeA---KLQ--FSKETEEEDLEE	282
WP_002314015	217	FTEK-MSKTKKAEtLLKY-F-PH	EKSNGLSQFIKLMVGNQGNFKNVf--GL-EEeA---KLQ--FSKETEEEDLEE	282
WP_002320716	217	FTEK-MSKTKKAEtLLKY-F-PH	EKSNGLSQFIKLMVGNQGNFKNVf--GL-EEeA---KLQ--FSKETEEEDLEE	282

WP_002330729	217	FTEK-MSKTKKKAETLLKY-F-PH	EKSNGLSQFIKLMVGNQGNFKNVF--GL-EE-A---KLQ--FSKETEYEDLEE	281
WP_002335161	217	FTEK-MSKTKKKAETLLKY-F-PH	EKSNGLSQFIKLMVGNQGNFKNVF--GL-EEA---KLQ--FSKETEYEDLEE	282
WP_002345439	217	FTEK-MSKTKKKAETLLKY-F-PH	EKSNGLSQFIKLMVGNQGNFKNVF--GL-EEA---KLQ--FSKETEYEDLEE	282
WP_034867970	210	LTDK-LSKTKKVEEILKY-Y-PT	EKINFFAQCLKLI VGNQANFKRIF--DLEAE-V---KLQ--FSKETEYEDLES	275
WP_047937432	217	FTEK-MSKTKKKAETLLKY-F-PH	EKSNGLSQFIKLMVGNQGNFKNVF--GL-EEA---KLQ--FSKETEYEDLEE	282
WP_010720994	210	LTDK-LSKTKKVEEILKY-Y-PT	EKINFFAQCLKLI VGNQANFKRIF--DLEAE-V---KLQ--FSKETEYEDLES	275
WP_010737004	210	LTDK-LSKTKKVEEILKY-Y-PT	EKINFFAQCLKLI VGNQANFKRIF--DLEAE-V---KLQ--FSKETEYEDLES	275
WP_034700478	210	LTDK-LSKTKKVEEILKY-Y-PT	EKINFFAQCLKLI VGNQANFKRIF--DLEAE-V---KLQ--FSKETEYEDLES	275
WP_007209003	215	LTQQ-LSKSERADNVLKL-F-PD	EKGTGIFAQFIKLI VGNQGNFKKVF--QLEED---qKLQ--LSTDDYEENIEN	280
WP_023519017	210	LTER-LSKAKRVEKVLAY-Y-PS	EKSTGNFAQFIKLMVGNQANFKKTF--DLEEE-M---KLN--FTRDCYEEDINE	275
WP_010770040	213	FSEK-VSRARKVEAILSV-Y-SE	EKSTGTLAQFIKLMVGNQGRFKKTF--DLEED-G---IIQ--IPKEEYEELEET	278
WP_048604708	209	FADK-VSRAKKAEGLAL-F-PD	EKRNGTFDQFIKMI VGNQGNFKKTF--ELEED-A---KLQ--FSKEEYDESLEA	274
WP_010750235	210	LTDK-LSKSKKVEKILQY-Y-PK	EKTGCLAQFIKLI VGNQGNFKQAF--HLDEE-V---KIQ--ISKETEYEDLEK	275
AII16583	251	LSAR-LSKSRRLLENLIAQ-L-PG	EKKNGLFGNLI ALSLGLTPNFKSNF--DLAED-A---KLQ--LSKDTYDDDLDN	316
WP_029073316	216	LKEP-LSKHKADKAFAL-FdTT	KDNKAA YKELCAALAGNKNFVTKMLkeAELHD-EdekDI sfkFSDATFDDAFVE	289
WP_031589969	216	LKEP-LSKHKAEKAFAL-FdTT	KDNKAA YKELCAALAGNKNFVTKMLkeAELHD-EdekDI sfkFSDATFDDAFVE	289
KDA45870	210	FKDNtFSKTKKSEELKL---SG	-KKQLAHQLFKMMVGNMGSFKKVL--GTDEE---hKLS--FGKDTYEDDLND	275
WP_039099354	207	LIDNhQASNRQOALLLiYtPS	KONKAIATELLKAILGLKAKFNVL T--GIEAEdvktTLT--FNAENFDEEMVK	285
AKP02966	209	LIGR-GNATQKSNILNN-F--T	KETKLLKEVINLILGNVAHLNTIFktSLTKDeE---KLS--FSGKDIESKLDD	278
WP_010991369	218	LVEK-VTRKEKLERILKL-Y-PG	EKSAGMFAQFISLIVGSKGNFQKPF--DLIEK-S---DIE--CAKDSYEEDLES	283
WP_033838504	218	LVEK-VTRKEKLERILKL-Y-PG	EKSAGMFAQFISLIVGSKGNFQKPF--DLIEK-S---DIE--CAKDSYEEDLES	283
EHN60060	221	LVEK-VTRKEKLERILKL-Y-PG	EKSAGMFAQFISLIVGSKGNFQKPF--DLIEK-S---DIE--CAKDSYEEDLES	286
EFR89594	1	-----LKL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKPF--DLIEK-S---DIE--CAKDSYEEDLES	52
WP_038409211	218	LSEK-LTRREKLDKILKL-Y-TG	EKSTGMFAQFISLIVGSKGNFQKPF--DLIEK-S---DIE--CAKDSYEEDLES	283
EFR95520		-----	-----	
WP_003723650	218	LAGE-FTRREKFERILQL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--DLIEK-T---DIE--CAKDSYEEDLET	283
WP_003727705	218	LAGE-FTRREKFERILRL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--NLVEK-T---DIE--CAKDSYEEDLEA	283
WP_003730785	218	LAGE-FTRREKFERILRL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--NLVEK-T---DIE--CAKDSYEEDLEA	283
WP_003733029	218	LAEK-FTRKDKLDKILSL-Y-PG	EKTG VFAQFVNIIVGSTGKFKKHF--NLHEK-K---DIN--CAEDTYDLDLES	283
WP_003739838	218	LAGE-FTRKEKLERILQL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--DLVEK-T---DIE--CAKDSYEEDLEA	283
WP_014601172	218	LAGE-FTRREKFERILQL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--DLIEK-T---DIE--CAKDSYEEDLEA	283
WP_023548323	218	LAGE-FTRREKFERILQL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--DLIEK-T---DIE--CAKDSYEEDLET	283
WP_031665337	218	LAGE-FTRREKFERILQL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--DLIEK-T---DIE--CAKDSYEEDLET	283
WP_031669209	218	LAEK-FTRKDKLDKILSL-Y-PG	EKTG VFAQFVNIIVGSTGKFKKHF--NLHEK-K---DIN--CAEDTYDLDLES	283

WP_033920898	218	LARK-FTRREKFERILQL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--DLIEK-T---DIE--CAKDSYEEDLET	283
AKI42028	221	LAGK-FTRREKFERILQL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--DLIEK-T---DIE--CAKDSYEEDLEA	286
AKI50529	221	LARK-FTRREKFERILQL-Y-PG	EKSTGMFAQFISLIVGSKGNFQKVF--DLIEK-T---DIE--CAKDSYEEDLET	286
EFR83390		-----	-----	
WP_046323366	218	FSEK-LTKREKLDKILNL-Y-PN	EKSTDLFAQFISLIIIGSKGNFKKFF--NLTEK-T---DIE--CAKDSYEEDLEV	283
AKE81011	228	LSAR-LSKRRLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A---KLQ--LSKDTYDDDDLDN	293
CUO82355	215	LKKP-LSKKAKVDEVMAL-IsPE	KEFKSAYKELVTGIAGNKMVTKMILcESIKQ-Gds-EIKlkFSDSNYDDQFSE	287
WP_033162887	216	LSKI-YQRSKKADLLKI-MnPT	KEEKAAAYKEFTKALVGLKFNISKMILaQEVKK-Gdt-DIVl eFSNANYDSTIDE	288
AGZ01981	245	LSAR-LSKRRLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A---KLQ--LSKDTYDDDDLDN	310
AKA60242	212	LSAR-LSKRRLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A---KLQ--LSKDTYDDDDLDN	277
AKS40380	212	LSAR-LSKRRLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A---KLQ--LSKDTYDDDDLDN	277
4UN5_B	216	LSAR-LSKRRLENLIAQ-L-PG	EKKNGLFGNLIALSGLTPNFKSNF--DLAED-A---KLQ--LSKDTYDDDDLDN	281

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WP_002345439	283	LLLEKIGDDYIDL	FVQAKNVYDAVLL	SEILSDSTKNT	TRAKLSAGMIRRY	DAHAKEDLVLL	KRFVKEN-LPKKYRAFFGDNSV	361	
WP_034867970	276	LLLEKIGDEYLDI	FLQAKKVDAILL	SEIISSTVKHT	KAKLSSGMVER	YERHKADLAKFKQ	FVKEN-VPQKATVFFKDTTK	354	
WP_047937432	283	LLLEKIGDDYIDL	FVQAKNVYDAVLL	SEILSDSTKNT	TRAKLSAGMIRRY	DAHAKEDLVLL	KRFVKEN-LPKKYRAFFGDNSV	361	
WP_010720994	276	LLLEKIGDEYLDI	FLQAKKVDAILL	SEIISSTVKHT	QAKLSSGMVER	YERHKADLAKFKQ	FVKEN-VPQKATVFFKDTTK	354	
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WP_048604708	275	LLGEIGDEYAD	VFAAKNVYNA	VELSGILVTVDN	STKAKLSASMI	KRYEDHKTDLK	LKFEFIRKN-LPEKYHEI	FNDKNT	353
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WP_033838504	284	LLALIIGDEYAE	LFWAAKNAYS	AVLSSIIITVAE	TETNAKLSASMI	ERFDTHEEDL	GELKAFIKLH-LPKHYEEI	FSNTEK	362
EHN60060	287	LLALIIGDEYAE	LFWAAKNAYS	AVLSSIIITVAE	TETNAKLSASMI	ERFDTHEEDL	GELKAFIKLH-LPKHYEEI	FSNTEK	365
EFR89594	53	LLALIIGDEYAE	LFWAAKNAYS	AVLSSIIITVAE	TETNAKLSASMI	ERFDTHEEDL	GELKAFIKLH-LPKHYEEI	FSNTEK	131
WP_038409211	284	LLAKIGDEYAEI	FVAAKSTYN	AVLNSIIITVTD	TETKAKLSASMI	ERFDKHAKDLKRL	KAFFKMQ-LPEKFN	EVFNDIEK	362
EFR95520		-----							
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WP_003727705	284	LLAIIGDEYAE	LFWAAKNTYN	AVLSSIIITVTD	TETNAKLSASMI	ERF	DAHEKDLVELKAFIKLN-LPKQYQEI	FNNAAI	362
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WP_003733029	284	LLAIIGDEFAE	VFWAAKNAYNA	VLSNIIITVTD	STTRAKLSASLI	ERFENHKKDLKMKR	FVRTY-LPEKYDEI	FDDTEK	362
WP_003739838	284	LLAIIGDEYAE	LFWAAKNTYN	AVLSSIIITVTD	TETNAKLSASMI	ERF	DAHEKDLSELKAFIKLH-LPKQYEEI	FSNVAI	362
WP_014601172	284	LLAIIGDEYAE	LFWAAKNTYN	AVLSSIIITVTD	TETNAKLSASMI	ERF	DAHEKDLVELKAFIKLN-LPKQYQEI	FNNAAI	362
WP_023548323	284	LLAIIGDEYAE	LFWAAKNTYN	AVLSSIIITVTD	TETNAKLSASMI	ERF	DAHEKDLVELKAFIKLN-LPKQYEEI	FSNAAI	362
WP_031665337	284	LLAIIGDEYAE	LFWAAKNTYN	AVLSSIIITVND	TETNAKLSASMI	ERF	DAHEKDLVELKAFIKLN-LPKQYEEI	FSNAAI	362
WP_031669209	284	LLAIIGDEFAE	VFWAAKNAYNA	VLSNIIITVTD	STTRAKLSASLI	ERFENHKKDLKMKR	FVRTY-LPEKYDEI	FDDTEK	362

WP_033920898	284	LLAII	IGDEYAELFVA	AKNTYNA	VWLVSSII	ITVTD	TETNAKLSAS	MIERF	DAHEKDL	VELKAFI	IKLN-LPKQ	YEEI	FSNA	AI	362
AKI42028	287	LLAII	IGDEYAELFVA	AKNTYNA	VWLVSSII	ITVTA	TETNAKLSAS	MIERF	DAHEKDL	GELKAFI	IKLH-LPKQ	YQEI	FNNA	AI	365
AKI50529	287	LLAII	IGDEYAELFVA	AKNTYNA	VWLVSSII	ITVTD	TETNAKLSAS	MIERF	DAHEKDL	VELKAFI	IKLN-LPKQ	YEEI	FSNA	AI	365
EFR83390															
WP_046323366	284	LLARV	GDEYAEI	FVA	AKNAYNA	VWLVSSII	ITVSN	TETKAKLSAS	MIERF	DKDKDL	KRMKAF	FFVR-LPEN	FNEV	FN	362
AKE81011	294	LLAQI	GDQYADL	FLAA	KNLSDAI	LLSDI	LRVNT	EITKAPL	SASMI	KRYDE	HHQDLTLL	KALVRQ	-LPEKY	KEI	372
CUO82355	288	VENDL	GE-YVE	FIDSL	HNIYSW	VELQ	TIMGATH	D-NASI	SEAMV	SRYNK	HHEDLQ	LLKCKI	KDN-VPK	KYFDM	364
WP_033162887	289	LQSE	LGE-YIE	FIEML	HNIYSW	VELQ	AILGATH	D-NPSI	SAAMV	ERYEE	HKDLR	VLKKV	IREE-LP	DKYNE	365
AGZ01981	311	LLAQI	GDQYADL	FLAA	KNLSDAI	LLSDI	LRVNT	EITKAPL	SASMI	KRYDE	HHQDLTLL	KALVRQ	-LPEKY	KEI	389
AKA60242	278	LLAQI	GDQYADL	FLAA	KNLSDAI	LLSDI	LRVNT	EITKAPL	SASMI	KRYDE	HHQDLTLL	KALVRQ	-LPEKY	KEI	356
AKS40380	278	LLAQI	GDQYADL	FLAA	KNLSDAI	LLSDI	LRVNT	EITKAPL	SASMI	KRYDE	HHQDLTLL	KALVRQ	-LPEKY	KEI	356
4UN5_B	282	LLAQI	GDQYADL	FLAA	KNLSDAI	LLSDI	LRVNT	EITKAPL	SASMI	KRYDE	HHQDLTLL	KALVRQ	-LPEKY	KEI	360

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 WP_045635197 357 --DGYAG YIDG K TTQETFYKYIKNLLSK-F--EGTDYFL--DKIEREDFLRKQRTFDNNGSIPHQIHLQEM 419
 5AXW_A 169 -----G SINR - -----TSDYVK-----EA 183
 WP_009880683 41 --NGYAG YIDG G ASQEEFYKFIKPILEK-M--DGTEELLA--KLNREDLLRKQRTFDNNGSIPHQIHLGEL 103
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WP_002282247	357	--NGYAG	YIEN	G	VKQDEFYKYLKNTLSK-I--TGSDFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002282906	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002283846	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002287255	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002288990	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002289641	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002290427	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419

WP_002295753	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002296423	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002304487	357	--NGYAG	YVGA	D	ATEEEFYKYVKGILNK-V--EGADVWL--DKIDREDFLRKQRTFDNGSIPHQIHLQEM	429
WP_002305844	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002307203	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002310390	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_002352408	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_012997688	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_014677909	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_019312892	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_019313659	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_019314093	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_019315370	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGNGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_019803776	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_019805234	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_024783594	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_024784288	357	--NGYAG	YIEN	G	VKQDEFYKYLKNTLSK-I--TGSDFL--DQIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_024784666	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_024784894	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_024786433	357	--NGYAG	YIEN	G	VKQDEFYKYLKNTLSK-I--AGSDYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_049473442	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_049474547	357	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
EMC03581	350	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_000428612	358	--DGYAG	YIDG	K	TNQEAFYKYLKGLLNK-I--EGSGYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_000428613	358	--DGYAG	YIDG	K	TTQESFYKYIKNLLSK-F--EGADYFL--EKIEREDFLRKQRTFDNGSIPHQIHLQEM	420
WP_049523028	357	--DGYAG	YIDG	K	TTQEGFYKYIKNLLSK-F--EGTDYFL--EKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_003107102	326	--NGYAG	YING	K	TSQEDFYKYIKPILSK-L--KGAESLIs--KLEREDFLRKQRTFDNGSIPHQIHLNEL	388
WP_054279288	359	--DGYAG	YISG	K	TSQEAFFYKIPILET-L--DGAEDFLt--KINREDFLRKQRTFDNGSIPHQIHLGEL	421
WP_049531101	358	--EGYAG	YIDS	K	TTQEAFFYKIKNLLSK-I--DGADYLL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	420
WP_049538452	358	--DGYAG	YVDG	K	TTQEAFFYKIKNLLSK-F--EGADYFL--EKIEREDFLRKQRTFDNGSIPHQIHLQEM	420
WP_049549711	358	--DGYAG	YIDG	K	TTQEAFFYKIKNLLSK-F--EGTDYFL--EKIEREDFLRKQRTFDNGSIPHQIHLQEM	420
WP_007896501	359	--NGYAG	YIEG	K	VSQEDFYRYIKPILSR-L--KGGDEFLa--KIDRDDFLRKQRTFDNGSIPHQIHLKEL	421
EFR44625	311	--NGYAG	YIEG	K	VSQEDFYRYIKPILSR-L--KGGDEFLa--KIDRDDFLRKQRTFDNGSIPHQIHLKEL	373
WP_002897477	357	--DGYAG	YIDG	K	TTQEAFFYKYIKNLLSK-F--EGADYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	419

WP_002906454	357	--DGYAG	FIDG	K	TTQEAFFYKIKNLLSK-L--EGADYFL--NKIEREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_009729476	358	--DGYAG	YIDG	K	TTQETFYKIKNLLSK-F--EGADYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	420
CQR24647	358	--NGYAG	YIDG	K	TNQEDFYKILNLLQK-V--DGGDYFI--EKIEREDFLRKQRTFDNGSIPHQVHLDEM	420
WP_000066813	358	--DGYAG	YIDG	K	TTQEAFFYKIKNLLSK-F--EGADYFL--DKIEREDFLKKQRTFDNGSIPHQIHLQEM	420
WP_009754323	358	--DGYAG	YIDG	K	TTQEAFFYKIKNLLSK-F--EGADYFL--DKIEREDFLRKQRTFDNGSIPHQIHLQEM	420
WP_044674937	357	--DGYAG	YIEG	K	TTQENFYRIFIKKAIEK-I--EGSDYFI--DKIDREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_044676715	357	--DGYAG	YIEG	K	TTQENFYRIFIKKAIEK-I--EGSNYFI--DKIDREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_044680361	357	--DGYAG	YIEG	K	TTQENFYRIFIKKAIEK-I--EGSNYFI--DKIDREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_044681799	357	--DGYAG	YIEG	K	TTQENFYRIFIKKAIEK-I--EGSDYFI--DKIDREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_049533112	357	--KGYAG	YIEN	G	VKQDEFYKYLKGIILQ-I--NGSGDFL--DKIDREDFLRKQRTFDNGSIPHQIHLQEM	419
WP_029090905	316	fyTDYIG	YEES	K	SKEERLFKHIELLLAKENv1TtVEHAL1eKNITFASLLPLQRSSRNAV1PYQVHEKEL	403
WP_006506696	361	ksKGYYN	YINR	K	APVDEFYKYVKKIEK-VdtPEAKQIln--DIELENFLLKQNSRTNGSVPYQMQLDEM	429
AIT42264	357	--NGYAG	YIDG	G	ASQEEFYKFIKPILEK-M--DGTEELLV--KLNREDLLRKQRTFDNGSIPHQIHLGEL	419
WP_034440723	363	--NGYAG	YIDG	K	TSQEDFYKFKVKAQLKG--eENGEYFL--EAIENENFLRKQRTFDNGSIPHQIHLQEL	425
AKQ21048	357	--NGYAG	YIDG	G	ASQEEFYKFIKPILEK-M--DGTEELLV--KLNREDLLRKQRTFDNGSIPHQIHLGEL	419
WP_004636532	356	--NGYAG	YIDG	K	TNQEDFYKIEKVMKT-IksDKDYFL--DKIDREDLLRKQRTFDNGSIPHQIHLQEM	420
WP_002364836	363	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_016631044	314	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	378
EMS75795	100	--NGYAG	YIDG	K	TTQEDFYKFLKKEKLN-I--AGSERFM--EKVDQENFLLKQRTTANGVIPHQVHLTEL	162
WP_002373311	363	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_002378009	363	--DGYAG	YITH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_002407324	363	--DGYAG	YITH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_002413717	363	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_010775580	365	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	429
WP_010818269	363	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_010824395	363	--DGYAG	YITH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_016622645	363	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_033624816	363	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_033625576	363	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_033789179	363	--DGYAG	YIAH	A	VSQLKFYQYVKKIIQD-I--AGAAYFL--EKIAQENFLRKQRTFDNGSIPHQIHLAEL	427
WP_002310644	361	--NGYAG	YIEG	H	ATQEDFYKFKVKKELTG-I--RGSEVFL--TKIEQENFLRKQRTFDNGSIPHQIHLTEL	423
WP_002312694	362	--NGYAG	YIEG	H	ATQEAFFYKFKVKKELTG-I--RGSEVFL--TKIEQENFLRKQRTFDNGSIPHQIHLSEL	424
WP_002314015	362	--NGYAG	YIEG	H	ATQEDFYKFKVKKELTG-I--RGSEVFL--TKIEQENFLRKQRTFDNGSIPHQIHLTEL	424
WP_002320716	362	--NGYAG	YIEG	H	ATQEDFYKFKVKKELTG-I--RGSEVFL--TKIEQENFLRKQRTFDNGSIPHQIHLTEL	424

WP_002330729	361	--NGYAG	YIEG	H	ATQEDFYKFKKELTG-I--RGSEVFL--TKIEQENFLRKQRTFDNGVIPHQIHLTEL	423
WP_002335161	362	--NGYAG	YIEG	H	ATQEDFYKFKKELTG-I--RGSEVFL--TKIEQENFLRKQRTFDNGVIPHQIHLTEL	424
WP_002345439	362	--NGYAG	YIEG	H	ATQEDFYKFKKELTG-I--RGSEVFL--TKIEQENFLRKQRTFDNGVIPHQIHLTEL	424
WP_034867970	355	--NGYAG	YIKG	K	TTQEEFYKFKKELSG-V--VGSEPFLL--EKIDQETFLKQRTYTNNGVIPHQVHLLIEL	417
WP_047937432	362	--NGYAG	YIEG	H	ATQEDFYKFKKELTG-I--RGSEVFL--TKIEQENFLRKQRTFDNGVIPHQIHLTEL	424
WP_010720994	355	--NGYAG	YIKG	K	TTQEEFYKFKKELSG-V--VGSEPFLL--EKIDQETFLKQRTYTNNGVIPHQVHLLIEL	417
WP_010737004	355	--NGYAG	YIKG	K	TTQEEFYKFKKELSG-V--VGSEPFLL--EKIDQETFLKQRTYTNNGVIPHQVHLLIEL	417
WP_034700478	355	--NGYAG	YIKG	K	TTQEEFYKFKKELSG-V--VGSEPFLL--EKIDQETFLKQRTYTNNGVIPHQVHLLIEL	417
WP_007209003	359	--NGYAG	YIDG	K	TKEEFYKYLKTTLVQ--KSGYQYFI--EKIEQENFLRKQRIYDNGVIPHQVHAEEL	421
WP_023519017	355	--NGYAG	YVKG	K	ATQEDFYKFLRTELALG-L--EESQSIM--EKIDLEIYLLKQRTFANGVIPHQIHLVEM	417
WP_010770040	358	--SGYAG	YVEN	S	VTQAEFYKYIKKAIK-V--PGAAYFL--EKIEQETFLDKQRTFNNNGVIPHQIHLTEL	422
WP_048604708	354	--DGYAG	YIDN	S	TSQEFYKYITNLIK-I--DGAAYFL--KKIENEDFLRKQRTFDNGIIPHQIHLTEL	418
WP_010750235	355	--DGYAG	YIDG	K	TTQADFYKFLKELTG-V--PGSEPM--AKIDQENFLKQRTPTNGVIPHQVHLTEF	417
AII16583	396	--NGYAG	YIDG	G	ASQEEFYKFIKPILEK-M--DGTEELLV--KLNREDLLRKQRTFDNGSIPHQIHLGEL	458
WP_029073316	367	kKNNYCN	YINH	K	TPVDFYKYIKKLIK-I--DDPDVKTILN--KIELESFMLKQNSRTNGAVPYQMLDEL	435
WP_031589969	367	kKNNYCN	YINH	K	TPVDFYKYIKKLIK-I--DDPDVKTILN--KIELESFMLKQNSRTNGAVPYQMLDEL	435
KDA45870	354	-iSGYAG	YIDG	K	VSEEDFYKYTKKTLKG-I--PETEILQ--KIDANNYLKQRTKDNNGVIPHQVHLKEL	417
WP_039099354	360	-----	YVDG	K	--SKEDFYGDI--TKALKNpdpPivSEIKK--LIELDQFMPKQRTKDNNGVIPHQVHLKEL	425
AKP02966	349	--QAYDD	YINK	K	---KELYTSLKFLKValp--TNLAKEAe--EKISKGTYLKPRNSENSEGVVYQLNKIEM	415
WP_010991369	363	--HGYAG	YIDG	-	TKQADFYKYMKTLEN-I--EGADYFI--AKIEKENFLRKQRTFDNGAIPHQLHLEEL	425
WP_033838504	363	--HGYAG	YIDG	-	TKQADFYKYMKTLEN-I--EGADYFI--AKIEKENFLRKQRTFDNGAIPHQLHLEEL	425
EHN60060	366	--HGYAG	YIDG	-	TKQADFYKYMKTLEN-I--EGADYFI--AKIEKENFLRKQRTFDNGAIPHQLHLEEL	428
EFR89594	132	--HGYAG	YIDG	-	TKQADFYKYMKTLEN-I--EGADYFI--AKIEKENFLRKQRTFDNGAIPHQLHLEEL	194
WP_038409211	363	--DGYAG	YIDG	-	TTQEKFYKYMKMLAN-I--DGADYFI--DQIEEENFLRKQRTFDNGIIPHQLHLEEL	425
EFR95520	1	-----	----	-	-----MCKMLAN-I--DGADYFI--DQIEEENFLRKQRTFDNGIIPHQLHLEEL	44
WP_003723650	363	--DGYAG	YIDG	-	TKQVDFYKYLKTTLEN-I--EGSDYFI--AKIEEENFLRKQRTFDNGAIPHQLHLEEL	425
WP_003727705	363	--DGYAG	YIDG	-	TKQVDFYKYLKTTLEN-I--EGADYFI--TKIEEENFLRKQRTFDNGVIPHQLHLEEL	425
WP_003730785	363	--DGYAG	YIDG	-	TKQVDFYKYLKTTLEN-I--EGADYFI--TKIEEENFLRKQRTFDNGVIPHQLHLEEL	425
WP_003733029	363	--HGYAG	YISG	-	TKQADFYKYMKTLEN-I--EGADYFI--AKIEEENFLRKQRTFDNGVIPHQLHLEEL	425
WP_003739838	363	--DGYAG	YIDG	-	TKQVDFYKYLKTTLEN-I--EGADYFI--AKIEEENFLRKQRTFDNGAIPHQLHLEEL	425
WP_014601172	363	--DGYAG	YIDG	-	TKQVDFYKYLKTTLEN-I--EGADYFI--AKIEEENFLRKQRTFDNGAIPHQLHLEEL	425
WP_023548323	363	--DGYAG	YIDG	-	TKQVDFYKYLKTTLEN-I--EGADYFI--AKIEEENFLRKQRTFDNGVIPHQLHLEEL	425
WP_031665337	363	--DGYAG	YIDG	-	TKQVDFYKYLKTTLEN-I--EGSDYFI--AKIEEENFLRKQRTFDNGAIPHQLHLEEL	425
WP_031669209	363	--HGYAG	YISG	-	TKQADFYKYMKTLEN-I--EGADYFI--AKIEEENFLRKQRTFDNGVIPHQLHLEEL	425

WP_033920898	363	--DGYAG	YIDG	-	TKQVDFYKYLKTILEN-I--EGADYFI--AKIEEENFLRKQRTFDNGAIPHQLHLEEL	425
AKI42028	366	--DGYAG	YIDG	-	TKQVDFYKYLKTILEN-I--EGADYFI--AKIEEENFLRKQRTFDNGAIPHQLHLEEL	428
AKI50529	366	--DGYAG	YIDG	-	TKQVDFYKYLKTILEN-I--EGADYFI--AKIEEENFLRKQRTFDNGAIPHQLHLEEL	428
EFR83390		-----	----	-	-----	
WP_046323366	363	--DGYAG	YIEG	-	TKQEAIFYKMKMLEH-V--EGADYFI--NQIEEENFLRKQRTFDNGAIPHQLHLEEL	425
AKE81011	373	--NGYAG	YIDG	G	ASQEEFYKFIKPILEK-M--DGTEELLV--KLNREDLLRKQRTFDNGSIPHQIHLGEL	435
CUO82355	365	kVKGYYN	YINR	K	APVDEFYKFKKCIK-VdtPEAKQIlh--DIELENFLLKQNSRTNGSVPYQMQLDEM	433
WP_033162887	366	klHNYLG	YIKY	D	TPVEEFYKIKGLLAK-VdtDEAREIle--RIDLEKFMKQNSRTNGSIPYQMQLDEM	434
AGZ01981	390	--NGYAG	YIDG	G	ASQEEFYKFIKPILEK-M--DGTEELLV--KLNREDLLRKQRTFDNGSIPHQIHLGEL	452
AKA60242	357	--NGYAG	YIDG	G	ASQEEFYKFIKPILEK-M--DGTEELLV--KLNREDLLRKQRTFDNGSIPHQIHLGEL	419
AKS40380	357	--NGYAG	YIDG	G	ASQEEFYKFIKPILEK-M--DGTEELLV--KLNREDLLRKQRTFDNGSIPHQIHLGEL	419
4UN5_B	361	--NGYAG	YIDG	G	ASQEEFYKFIKPILEK-M--DGTEELLV--KLNREDLLRKQRTFDNGSIPHQIHLGEL	423

WP_010922251	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_039695303	423	HAILRRQGDYFFLKE--KQD	RIEKILTFRIPYYVGPL	VRKD--SRFAWAEY--RSDEKITPWNFDKVIDKEK	489
WP_045635197	420	NAILRRQGEYFFLKD--NKE	KIEKILTFRIPYYVGPL	ARGN--RDFAWLTR--NSDEAIRPWNFEIIVDKAS	486
5AXW A	184	KQLLKVQKAYHQLDQSF1--D	TYIDLLETRRTYYEGPG	--Eg-SPFGWKDI-----	229
WP_009880683	104	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	170
WP_010922251	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_011054416	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_011284745	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_011285506	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_011527619	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_012560673	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_014407541	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_020905136	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_023080005	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_023610282	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_030125963	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_030126706	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_031488318	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_032460140	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_032461047	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_032462016	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_032462936	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_032464890	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_033888930	245	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	311
WP_038431314	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_038432938	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_038434062	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
BAQ51233	331	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	397
KGE60162		-----	-----	-----	
KGE60856		-----	-----	-----	
WP_002989955	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_003030002	420	HAILRRQEEHYFFLKE--NQD	KIEKILTFRIPYYVGPL	ARKG--SRFAWAEY--KADEKITPWNFDDILDKEK	486
WP_003065552	423	HAILRRQGDYFFLKE--NQD	RIEKILTFRIPYYVGPL	ARKD--SRFSAWAEY--HSDEKITPWNFDKVIDKEK	489
WP_001040076	421	RAIIRRQSEYFFLKE--NLD	RIEKILTFRIPYYVGPL	AREK--SDFAWMTR--KTDDSI RPNFNFEDLVDKEK	487

KLJ37842	421	KAIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
KLJ72361	421	KAIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
KL20707	421	KAIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
KL242645	421	KAIIRRQSEYYPFLKE--NLD	RIEKILTFRIPYYVGPL	AREK--SDFAWMTR--KTDDSI	487
WP_047207273	421	KAIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
WP_047209694	421	RAIIRRQSEYYPFLKE--NLD	RIEKILTFRIPYYVGPL	AREK--SDFAWMTR--KTDDSI	487
WP_050198062	421	KAIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
WP_050201642	421	KAIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
WP_050204027	421	KAIIRRQSEYYPFLKE--NLD	RIEKILTFRIPYYVGPL	AREK--SDFAWMTR--KTDDSI	487
WP_050881965	421	KAIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
WP_050886065	421	KDIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
AHN30376	421	KAIIRRQSEYYPFLKE--NQD	KIEKILTFRIPYYVGPL	ARGN--SDFAWMTR--KTDDSI	487
EAO78426	421	KAIIRRQSEYYPFLKE--NQD	RIEKILTFRIPYYIGPL	AREK--SDFAWMTR--KTDDSI	487
CCW42055	421	RAIIRRQSEYYPFLKE--NLD	RIEKILTFRIPYYVGPL	AREK--SDFAWMTR--KTDDSI	487
WP_003041502	420	HAILRRQGEHYPPFLKE--NQD	KIEKILTFRIPYYVGPL	ARKG--SRFAWAEY--KADEKIT	486
WP_037593752	421	HAILRRQGEHYPPFLKE--NQD	KIEKILTFRIPYYVGPL	ARKG--SRFAWAEY--KADEKIT	487
WP_049516684	421	HAILRRQGEHYPPFLKE--NQD	KIEKILTFRIPYYVGPL	ARKG--SRFAWAEY--KADEKIT	487
GAD46167	420	HAILRRQGEHYPPFLKE--NQD	KIEKILTFRIPYYVGPL	ARKG--SRFAWAEY--KADEKIT	486
WP_018363470	421	HAILRRQGEHYPPFLKE--NQD	KIEKILTFRIPYYVGPL	ARKG--SRFAWAEY--KADEKIT	486
WP_003043819	430	HAILRRQGEHYPPFLKE--NQD	EIEKILTFRIPYYVGPL	ARKD--SRFAWAEY--KADEKIT	487
WP_006269658	420	HAILRRQGEHYPPFLKE--NRE	KIEKILTFRIPYYVGPL	ARGN--SRFAWAEY--KSEEAIT	496
WP_048800889	420	HAILRRQGEHYPPFLKE--NQD	KIEKILTFRIPYYVGPL	ARKG--SRFAWAEY--KADEKIT	486
WP_012767106	420	HAILRRQEDFYPPFLKD--NRE	KIEKILTFRIPYYVGPL	VRKG--SRFAWAEY--KADEKIT	486
WP_014612333	420	HAILRRQEDFYPPFLKD--NRE	KIEKILTFRIPYYVGPL	ARGN--SRFAWAEY--KSEETIT	486
WP_015017095	420	HAILRRQEDFYPPFLKD--NRE	KIEKILTFRIPYYVGPL	ARGN--SRFAWAEY--KSEETIT	486
WP_015057649	420	HAILRRQEDFYPPFLKD--NRE	KIEKILTFRIPYYVGPL	ARGN--SRFAWAEY--KSEETIT	486
WP_048327215	420	HAILRRQEDFYPPFLKD--NRE	KIEKILTFRIPYYVGPL	ARGN--SRFAWAEY--KSEETIT	486
WP_049519324	420	HAILRRQEDFYPPFLKD--NRE	KIEKILTFRIPYYVGPL	ARGN--SRFAWAEY--KSEETIT	486
WP_012515931	420	HAILRRQEVFFFLKD--NRK	KIESLLTFRIPYYVGPL	ARG-h-SRFAWVKR--KFDGAI	486
WP_021320964	420	HAILRRQEVFFFLKD--NRK	KIESLLTFRIPYYVGPL	ARG-h-SRFAWVKR--KFDGAI	486
WP_037581760	420	HAILRRQEVFFFLKD--NRK	KIESLLTFRIPYYVGPL	ARG-h-SRFAWVKR--KFDGAI	486
WP_004232481	420	RTILRRQGEYYPFLKE--NQA	KIEKILTFRIPYYVGPL	ARKN--SRFAWAKY--HSDEPIT	486
WP_009854540	421	HAILRRQGDYYPFLKE--KQD	RIEKILTFRIPYYVGPL	VRKD--SRFAWAEY--RSDEKIT	487
WP_012962174	421	HAILRRQGEHYAFLKE--NQA	KIEKILTFRIPYYVGPL	ARKN--SRFAWAEY--HSDEKIT	487

WP_039695303 423 HAILRRQGDYYPFLKE--KQD RIEKILTFRIPYYVGPL 489
 WP_014334983 420 HSILRRQGDYYPFLKE--NQA KIEKILTFRIPYYVGPL 486
 WP_003099269 420 KAIIRRQEKFYFPLKE--NOK KIEKILTFKIPYYVGPL 486
 AHY15608 420 KAIIRRQEKFYFPLKE--NOK KIEKILTFKIPYYVGPL 486
 AHY17476 420 KAIIRRQEKFYFPLKE--NOK KIEKILTFKIPYYVGPL 486
 ESR09100 -----
 AGM98575 420 KAIIRRQEKFYFPLKE--NOK KIEKILTFKIPYYVGPL 486
 ALF27331 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_018372492 433 QAIILNQSKYYPLAE--NKE KIEKILTFRIPYYVGPL 499
 WP_045618028 421 NAIIRRQGEHYFPLQE--NKE KIEKILTFRIPYYVGPL 487
 WP_045635197 420 NAIIRRQGEYFPLKD--NKE KIEKILTFRIPYYVGPL 486
 WP_002263549 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002263887 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002264920 420 HAILRRQGDYYPFLKE--NOK RIEKILTFRIPYYVGPL 486
 WP_002269043 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002269448 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002271977 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002272766 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002273241 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002275430 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002276448 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002277050 420 HAILRRQGDYYPFLKE--NOK RIEKILTFRIPYYVGPL 486
 WP_002277364 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002279025 420 RAIIRRQSEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002279859 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002280230 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002281696 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002282247 420 HAILRRQGDYYPFLKE--NOK RIEKILTFRIPYYVGPL 486
 WP_002282906 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002283846 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002287255 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002288990 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002289641 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486
 WP_002290427 420 RAIIRRQAEFYFPLAD--NQD RIEKILTFRIPYYVGPL 486

WP_002295753	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_002296423	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_002304487	430	HAILRRQGEHYFFLKE--NQD	KIEKILTFRIPYYVGPL	VRKG--SRFAWAEY--KADEKITPWNFDDILDKEK	496
WP_002305844	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_002307203	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_002310390	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_002352408	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_012997688	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_014677909	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_019312892	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_019313659	420	RAIIRRQAEFYFFLAD--NQD	RIEKLLTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_019314093	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_019315370	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_019803776	420	RAIIRRQAEFYFFLAD--NQD	RIEKLLTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_019805234	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_024783594	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_024784288	420	HAILRRQGDYFFLKE--NQD	RIEKILTFRIPYYVGPL	ARKN--SRFAWAEY--HSDEAVTPWNFDQVIDKES	486
WP_024784666	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_024784894	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ASGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_024786433	420	HAILRRQGDYFFLKE--NQD	RIEKILTFRIPYYVGPL	ARKN--SRFAWAEY--HSDEAVTPWNFDQVIDKES	486
WP_049473442	420	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
WP_049474547	420	RAIIRRQAEFYFFLAD--NQD	RIEKLLTFRIPYYVGPL	ASGK--SDFAWLSR--KSADKITPWNFDEIVDKES	486
EMC03581	413	RAIIRRQAEFYFFLAD--NQD	RIEKILTFRIPYYVGPL	ARGK--SDFAWLSR--KSADKITPWNFDEIVDKES	479
WP_000428612	421	NAILRRQGEHYFFLKE--NKE	KIEKILTFRIPYYVGPL	ARGN--RDFAWLTR--NSDQAIRPWNFEEIVDKAS	487
WP_000428613	421	NAILRRQGEHYFFLKD--NKE	KIEKILTFRIPYYVGPL	ARGN--RDFAWLTR--NSDEAIRPWNFEEIVDKAS	487
WP_049523028	420	NAILRHQGEYFFLKE--NKD	KIEQILTFRIPYYVGPL	ARGN--SDFAWLSR--NSDEAIRPWNFEEIVDKSS	486
WP_003107102	389	KSIIRRQEKYFFLKD--KQV	RIEKILTFRIPYYVGPL	ANG-n-SSFAWVKR--RSNESITPWNFEEVVEQEA	455
WP_054279288	422	QAILERQAYFFLKD--NQE	KIEKILTFRIPYYIGPL	ARG-n-SRFAWLTR--TSDQKITPWNFDEMVDQEA	488
WP_049531101	421	NAILRRQGEHYFFLKE--NRE	KIEKILTFRIPYYVGPL	ARGN--RDFAWLTR--NSDQAIRPWNFEEIVDKAS	487
WP_049538452	421	NAILRRQGEHYFFLKE--NKE	KIEKILTFRIPYYVGPL	ARGN--RDFAWLTR--NSDQAIRPWNFEEIVDKAS	487
WP_049549711	421	NAILRRQGEHYFFLKE--NKE	KIEKILTFRIPYYVGPL	ARGN--RDFAWLTR--NSDQAIRPWNFEEIVDKAS	487
WP_007896501	422	HAILRRQEKYFFLAE--QKE	KIEQLLCFRIPYYVGPL	AKGgn-SSFAWVKR--RSDEPITPWNFKDVVDEEA	489
EFR44625	374	HAILRRQEKYFFLAE--QKE	KIEQLLCFRIPYYVGPL	AKGgn-SSFAWVKR--RSDEPITPWNFKDVVDEEA	441
WP_002897477	420	NAILRRQGEHYFFLKE--NRE	KIEKILTFRIPYYVGPL	ARDN--RDFSWLTR--NSDEPIRPNWNFEEIVDKAR	486

WP_002906454	420	NAILRRQGEHYLFLKE--NRE	KIEKILAFRIPIYYVGPL	ARGN--RDFAWLTR--NSDQAIRPWNFEVVDKAS	486
WP_009729476	421	NAILRRQGEHYFFLKE--NKE	KIEKILTFRIPIYYVGPL	ARGN--RDFAWLTR--NSDQAIRPWNFEVVDKAS	487
CQR24647	421	KAILRRQGEFYFFLKE--NAE	KIQIILTFKIPYYVGPL	ARGN--SRFAWASY--NSNEKMTPNFDNVIDKTS	487
WP_000066813	421	NAIIRRQGEHYFFLQE--NKE	KIEKILTFRIPIYYVGPL	ARGN--GDFAWLTR--NSDQAIRPWNFEVVDQAS	487
WP_009754323	421	NAILRRQGEHYPLLKE--NKE	KIEKILTFRIPIYYVGPL	ARGN--RDFAWLTR--NSDQAIRPWNFEVVDKAS	487
WP_044674937	420	HAIIRRQAEFFPFLVE--NQD	KIEKILTFRIPIYYVGPL	ARGK--SEFAWLNR--KSDEKIRPWNFDEMVDKET	486
WP_044676715	420	HAIIRRQAEFFPFLVE--NQD	KIEKILTFRIPIYYVGPL	ARGK--SEFAWLNR--KSDEKIRPWNFDEMVDKET	486
WP_044680361	420	HAIIRRQAEFFPFLVE--NQD	KIEKILTFRIPIYYVGPL	ARGK--SEFAWLNR--KSDEKIRPWNFDEMVDKET	486
WP_044681799	420	HAIIRRQAEFFPFLVE--NQD	KIEKILTFRIPIYYVGPL	ARGK--SEFAWLNR--KSDEKIRPWNFDEMVDKET	486
WP_049533112	420	HAILRRQEEHYFFLKE--NQD	KIEKILTFRIPIYYVGPL	ARGK--SRFAWAEY--KADEKITPWNFDDIILDK	486
WP_029090905	404	VAILLENQATYFFLLE--QKD	NIHKLTLTFRIPIYYVGPL	ADQKd--SEFAMVVR--KQAGKITPWFNFEEMVDIDA	471
WP_006506696	430	IKIIDNQAEEYPIILKE--KRE	QLLSILTLTFRIPIYYVGPL	ETSEh--AWIKRlegKENQRILPWNQDIIVDVA	498
AIT42264	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPIYYVGPL	ARG-n--SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_034440723	426	TAVLDQEQEKHYFFLKE--NRD	KIISLLTLTFRIPIYYVGPL	AKGE--SRFAWLER--sNSEEKIKPWNFKIIVDIDK	493
AKQ21048	420	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPIYYVGPL	ARG-n--SRFAWMTR--KSEETITPWNFEVVDKGA	486
WP_004636532	421	QAILDRQSQYFFLAE--NRD	KIESLVTFRIPYYVGPL	TVSDq--SEFAMMER--QSDEPIRPWNFDEIVNKR	488
WP_002364836	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_016631044	379	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	446
EMS75795	163	KAIIERQKPYPSLEE--ARD	KMIRLLTFRIPIYYVGPL	AQGEetSSFAWLER--KTPEKVTPNATEVIDYSA	231
WP_002373311	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_002378009	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_002407324	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_002413717	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_010775580	430	QAIHRQAAYFFLKE--NQK	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	497
WP_010818269	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_010824395	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_016622645	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_033624816	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_033625576	428	QAIHRQAAYFFLKE--NQE	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_033789179	428	QAIHRQAAYFFLKE--NQK	KIEQLVTFRIPYYVGPL	SKGDa--STFAWLKR--QSEEPiRPWNLQETVDDLQ	495
WP_002310644	424	RAIIANQKKHYFFLKE--EQE	KLESLLTFKIPYYVGPL	AKKQenSPFAWLIR--KSEEKIKPWNLPEIIVDMEG	492
WP_002312694	425	RAIIANQKKHYFFLKE--EQE	KLESLLTFKIPYYVGPL	AKKQenSPFAWLIR--KSEEKIKPWNLPEIIVDMEG	493
WP_002314015	425	RAIIANQKKHYFFLKE--EQE	KLESLLTFKIPYYVGPL	AKKQenSPFAWLIR--KSEEKIKPWNLPEIIVDMEG	493
WP_002320716	425	RAIIANQKKHYFFLKE--EQE	KLESLLTFKIPYYVGPL	AKKQenSPFAWLIR--KSEEKIKPWNLPEIIVDMEG	493

WP_002330729	424	RAIIANQKHHYFFLKE--EQE	KLESLLTFKIPYYVGPL	AKKQenSPFAWLIR--KSEEKIKPWNLP	492
WP_002335161	425	RAIIANQKHHYFFLKE--EQE	KLESLLTFKIPYYVGPL	AKKQenSPFAWLIR--KSEEKIKPWNLP	493
WP_002345439	425	RAIIANQKHHYFFLKE--EQE	KLESLLTFKIPYYVGPL	AKKQenSPFAWLIR--KSEEKIKPWNLP	493
WP_034867970	418	KAIIDQKQKHHYFFLEE--AGP	KIIALFKFRIPYYVGPL	AKEQeasSFAWIER--KTAEKINPWN	486
WP_047937432	425	RAIIANQKHHYFFLKE--EQE	KLESLLTFKIPYYVGPL	AKKQenSPFAWLIR--KSEEKIKPWNLP	493
WP_010720994	418	KAIIDQKQKHHYFFLEE--AGP	KIIALFKFRIPYYVGPL	AKEQeasSFAWIER--KTAEKINPWN	486
WP_010737004	418	KAIIDQKQKHHYFFLEE--AGP	KIIALFKFRIPYYVGPL	AKEQeasSFAWIER--KTAEKINPWN	486
WP_034700478	418	KAIIDQKQKHHYFFLEE--AGP	KIIALFKFRIPYYVGPL	AKEQeasSFAWIER--KTAEKINPWN	486
WP_007209003	422	RAILRKQEKYYFFLKE--NHE	KIEQIFKVRIPYYVGPL	AKHNeqSRFAWNI--KSDEPIRPN	490
WP_023519017	418	REIMDRQKRFYFFLKG--AQG	KIEKLLTFRIPYYVGPL	AQEGq-SPFAWIKR--KSPSQITP	485
WP_010770040	423	EAIIQKQATYYFFLAD--NKE	EMKQLVTFRIPYYVGPL	ADGN--SPFAWLER--ISSEPIR	489
WP_048604708	419	KAILHHQAMYYFFLQE--KFS	NFVDLLTFRIPYYVGPL	ANGN--SRFSWLSR--KSDEPIR	485
WP_010750235	418	KAIIDQKQKYYFFLEK--SKE	KMIQLLTFRIPYYVGPL	AQDKetSSFAWLER--KTTEKIK	486
AII16583	459	HAILRRQEDFYFFLKD--NRE	KIEKILTFRIPYYVGPL	ARG-n-SRFAMMTR--KSEETIT	525
WP_029073316	436	NKILENQSYYSDLKD--NED	KIRSILTFRIPYYVGPL	ITKDr--QFDWIIKkegKENERIL	506
WP_031589969	436	NKILENQSYYSDLKD--NED	KIRSILTFRIPYYVGPL	ITKDr--QFDWIIKkegKENERIL	506
KDA45870	418	VAIVENQKYYFFLRE--NKD	KFEKILNFRIPYYVGPL	ARGN--SKFAWLTR--a-GEK	484
WP_039099354	426	DRIENQKYYPWLAE--LNP	KLDELVAFRVYYVGPL	QQSSsdAKFAWMIR--KAEGQIT	509
AKP02966	416	EKIIDNQSYYFFLKE--NKE	KLLSILSFRIPYYVGPL	-QSSekNPFAMMER--KSNGHAR	483
WP_010991369	426	EAILHQQAKYYFFLKE--NYD	KIKSLVTFRIPYYVGPL	ANGQ--SEFAWLTR--KADGEIR	492
WP_033838504	426	EAILHQQAKYYFFLKE--NYD	KIKSLVTFRIPYYVGPL	ANGQ--SEFAWLTR--KADGEIR	492
EHN60060	429	EAILHQQAKYYFFLKE--NYD	KIKSLVTFRIPYYVGPL	ANGQ--SEFAWLTR--KADGEIR	495
EFR89594	195	EAILHQQAKYYFFLKE--NYD	KIKSLVTFRIPYYVGPL	ANGQ--SEFAWLTR--KADGEIR	261
WP_038409211	426	EAILHQQAKYYFFLRK--DYE	KIRSLVTFRIPYFIGPL	ANGQ--SDFAWLTR--KADGEIR	492
EFR95520	45	EAILHQQAKYYFFLRK--DYE	KIRSLVTFRIPYFIGPL	ANGQ--SDFAWLTR--KADGEIR	111
WP_003723650	426	EAILHQQAKYYFFLKE--DYD	KIKSLVTFRIPYFVGPL	ANGQ--SEFAWLTR--KADGEIR	492
WP_003727705	426	EAILHQQAKYYFFLRE--GYD	KIKSLVTFRIPYFVGPL	ANGQ--SEFAWLTR--KDDGEIR	492
WP_003730785	426	EAILHQQAKYYFFLRE--GYD	KIKSLVTFRIPYFVGPL	ANGQ--SEFAWLTR--KDDGEIR	492
WP_003733029	426	EAILHQQAKYYFFLRE--DYE	KIKSLVTFRIPYFVGPL	ANGQ--SEFAWLTR--KDDGEIR	492
WP_003739838	426	EAILHQQAKYYFFLKE--AYD	KIKSLVTFRIPYFVGPL	ANGQ--SDFAWLTR--KADGEIR	492
WP_014601172	426	EAILHQQAKYYFFLRE--DYE	KIKSLVTFRIPYFVGPL	ANGQ--SEFAWLTR--KADGEIR	492
WP_023548323	426	EAILHQQAKYYFFLRE--DYE	KIKSLVTFRIPYFVGPL	ANGQ--SEFAWLTR--KADGEIR	492
WP_031665337	426	EAILHQQAKYYTFLKE--DYD	KIKSLVTFRIPYFVGPL	ANGQ--SEFAWLTR--KADGEIR	492
WP_031669209	426	EAILHQQAKYYFFLRE--DYE	KIKSLVTFRIPYFVGPL	ANGQ--SEFAWLTR--KADGEIR	492

WP_033920898	426	EAIHQQAKYYPFLRE--DYE	KIKSLVTFRIPIYFVGPL	AKGQ--SEFAWLTR--KADGEIRPWNIEEKVDFGK	492
AKI42028	429	EAIHQQAKYYPFLRE--DYE	KIKSLVTFRIPIYFVGPL	AKGQ--SEFAWLTR--KADGEIRPWNIEEKVDFGK	495
AKI50529	429	EAIHQQAKYYPFLRE--DYE	KIKSLVTFRIPIYFVGPL	AKGQ--SEFAWLTR--KADGEIRPWNIEEKVDFGK	495
EFR83390		-----	-----	-----	
WP_046323366	426	EAILHQQAKYYPFLKV--DYE	KIKSLVTFRIPIYFVGPL	ANGQ--SEFSWLTR--KADGEIRPWNIEEKVDFGK	492
AKE81011	436	HAILRRQEDFYFLKD--NRE	KIEKILTFRIPIYFVGPL	ARG-n-SRFAMMTR--KSEETITPWNFEVVDKGA	502
CUO82355	434	IKIIDNQAKYYPVLKE--KRE	QLLSILTFRIPIYFVGPL	ETSEh---AWIKRlegKENQRILPWNYYQDTVDVDA	502
WP_033162887	435	IQIIDNQSVYYPQLKE--NRD	KLISILEFRIPIYFVGPL	AHSE---FAWIKKfedKQKERILPWNYYDQIVDIDA	503
AGZ01981	453	HAILRRQEDFYFLKD--NRE	KIEKILTFRIPIYFVGPL	ARG-n-SRFAMMTR--KSEETITPWNFEVVDKGA	519
AKA60242	420	HAILRRQEDFYFLKD--NRE	KIEKILTFRIPIYFVGPL	ARG-n-SRFAMMTR--KSEETITPWNFEVVDKGA	486
AKS40380	420	HAILRRQEDFYFLKD--NRE	KIEKILTFRIPIYFVGPL	ARG-n-SRFAMMTR--KSEETITPWNFEVVDKGA	486
4UN5_B	424	HAILRRQEDFYFLKD--NRE	KIEKILTFRIPIYFVGPL	ARG-n-SRFAMMTR--KSEETITPWNFEVVDKGA	490

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 WP_039695303 490 SAEKFI TRMTLNDLYLPEEKVLPKSHVYETAYAVNELTKIKYVN--EQGKES-FFDSNMKQEIFDHVFK--ENR-KVTK 563
 WP_045635197 487 SAEDFINKMTNDLYLPEEKVLPKSHLLYETFAVYNELTKVKFIA--EGLRDYqFLDSGQKKQIVNQLFK--ENR-KVTE 561
 5AXW A 230 --KEWYEMLMGHCTYFPEELRSVKYAYNADLYNALNDLNLNLAITR--DENEKLeY--KFQI IENVFK--QKK-KPTL 299
 WP_009880683 171 SAQSFIERMTNFDKKNLPNEKVLPHKSHLLYEFYFTVYNELTKVKYVT--EGMRKPaFLSGEQKKAIVDLLFK--TNR-KVTV 245
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 BAQ51233 398 SAQSFIERMTNFDKKNLPNEKVLPHKSHLLYEFYFTVYNELTKVKYVT--EGMRKPaFLSGEQKKAIVDLLFK--TNR-KVTV 472
 KGE60162 -----
 KGE60856 -----
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 WP_003065552 490 SAEKFI TRMTLNDLYLPEEKVLPKSHVYETAYAVNELTKIKYVN--EQGKDS-FFDSNMKQEIFDHVFK--ENR-KVTK 563
 WP_001040076 488 SAEAFIHRMTNNDLYLPEEKVLPKSHLLYEFYFTVYNELTKVRFIA--EGFKDFqFLNRKQKETIFNSLFK--EKR-KVTE 562

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WP_014334983	487	SAEKFI TRMTLNDLYLPEEKVLPKHSVYETFTVYNELTKIKYVN--EQGESF--FFDANMKQEI FDHVFK--ENR-KVTK	560
WP_003099269	487	SARAFIERMTNFDTYLPPEEKVLPKHSPLYEMFMVYNELTKVKYQT--EGMKRPVFLSSEDKKEI VNL LFK--KER-KVTV	561
AHY15608	487	SARAFIERMTNFDTYLPPEEKVLPKHSPLYEMFMVYNELTKVKYQT--EGMKRPVFLSSEDKKEI VNL LFK--KER-KVTV	561
AHY17476	487	SARAFIERMTNFDTYLPPEEKVLPKHSPLYEMFMVYNELTKVKYQT--EGMKRPVFLSSEDKKEI VNL LFK--KER-KVTV	561
ESR09100		-----	
AGM98575	487	SARAFIERMTNFDTYLPPEEKVLPKHSPLYEMFMVYNELTKVKYQT--EGMKRPVFLSSEDKKEI VNL LFK--KER-KVTV	561
ALF27331	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_018372492	500	SASRFIERMTLHDLVLPDEKVLPRHSLIYEKYTVFNELTKVRFPT--EGGKEV--YFSKTDKENI FDSL FK--RYR-KVTK	573
WP_045618028	488	SAEDFINKMTNYDLYLPPEEKVLPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSGQKQI VTL QLFK--EKR-KVTE	562
WP_045635197	487	SAEDFINKMTNYDLYLPPEEKVLPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSGQKQI VNL QLFK--ENR-KVTE	561
WP_002263549	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002263887	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002264920	487	SVEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002269043	487	SVEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002269448	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002271977	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002272766	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002273241	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002275430	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002276448	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002277050	487	SAQAFIEHMTNNDLYLPNEKVLPRHSLLYEKFTVYNELTKIKYVT--EIGEAK--FFDANL KQEI FDGLFK--HER-KVTK	560
WP_002277364	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002279025	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002279859	487	SVEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002280230	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002281696	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002282247	487	SAQAFIEHMTNNDLYLPNEKVLPRHSLLYEKFTVYNELTKIKYVT--EIGEAK--FFDANL KQEI FDGLFK--HER-KVTK	560
WP_002282906	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002283846	487	SVEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002287255	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002288990	487	SVEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002289641	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560
WP_002290427	487	SAEAFINRMTNYDLYLPNQKVLPRHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK	560

WP_002295753 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_002296423 487 SVEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_002304487 497 SAEKFI TRMTLNDLYLPEEKVLPKHSLLYETFTVYNELTKVKYVN--EQGEAK--FFDANMKQEI FDHVFK--ENR-KVTK 570
 WP_002305844 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_002307203 487 SVEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_002310390 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_002352408 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_012997688 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_014677909 487 SVEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_019312892 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_019313659 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_019314093 487 SVEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_019315370 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_019803776 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_019805234 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_024783594 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_024784288 487 SAQAFIEHMTNNDLYLPEEKVLPKHSLLYEKFTVYNELTKVKYVT--EIGEAK--FFDANLQEI FDGLFK--HER-KVTK 560
 WP_024784666 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_024784894 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_024786433 487 SAQAFIEHMTNNDLYLPEEKVLPKHSLLYEKFTVYNELTKVKYVT--EIGEAK--FFDANLQEI FDGLFK--HER-KVTK 560
 WP_049473442 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 WP_049474547 487 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 560
 EMC03581 480 SAEAFINRMTNIDLKPKHSLLYEKFTVYNELTKVKYKT--EQGKTA--FFDANMKQEI FDGVFK--VYR-KVTK 553
 WP_000428612 488 SAESFINKMTNIDLKPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSRQKKDIFYTLFKaeDKR-KVTE 564
 WP_000428613 488 SAEDFINKMTNIDLKPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSGQKKQIVTQLFK--EKR-KVTE 562
 WP_049523028 487 SAEDFIHRMTNIDLKPKHSLLYETFTVYNELTKVKYIA--EGMKDYqFLDSGQKKQIVNQLFK--EKR-KVTE 561
 WP_003107102 456 SAKVFIERTNFDTYLPPEEKVLPKHSLLYEMFTVYNELTKVKYQA--EGMRKPeFLSSEEKIEIVSNLFLK--TER-KVTV 530
 WP_054279288 489 SAQAFIERMTNFDEYLPQEKVLPKHSLTYEYFTVYNELTKVKYVT--EGMTKPeFLSAGQKEQIVELLEFK--KYR-KVTV 563
 WP_049531101 488 SAEAFINKMTNIDLKPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSGQKKQIVNQLFK--EKR-KVTE 562
 WP_049538452 488 SAEDFINKMTNIDLKPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSGQKKQIVNQLFK--EKR-KVTE 562
 WP_049549711 488 SAEDFINKMTNIDLKPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSGQKKQIVNQLFK--EKR-KVTE 562
 WP_007896501 490 SAQAFIEGTMNDYTLPEEKVLPKHSLPEYEMFTVYNELTKVKYIA--ENMTKPI YLSAEQKEAIDHLLFK--QTR-KVTV 564
 EFR44625 442 SAQAFIEGTMNDYTLPEEKVLPKHSLPEYEMFTVYNELTKVKYIA--ENMTKPI YLSAEQKEAIDHLLFK--QTR-KVTV 516
 WP_002897477 487 SAEDFIHRMTNIDLKPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSGQKKQIVNQLFK--EKR-KVTE 561

WP_002906454 487 SAEDFINKMTNYDLYLPEEKVLPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSDGQKKQIVNQLFK--DKR-KVTE 561
 WP_009729476 488 SAEDFINKMTNYDLYLPEEKVLPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFLDSDGQKKQIVTQLFK--EKR-KVTE 562
 CQR24647 488 SAQAFIERMTNNDLYLPDQKVLPKHSLLYQKFAVYNELTKIKYVT--ETGEAR-LFDVFLKKEIFDGLFK--KER-KVTK 561
 WP_000066813 488 SAEDFINKMTNYDLYLPEEKVLPKHSLLYETFAVYNELTKVKFIA--EGLTRYqFLDKKQKKDIFDTFFKaeNKR-KVTE 564
 WP_009754323 488 SAESFINKMTNYDLYLPEEKVLPKHSLLYETFAVYNELTKVKFIA--EGLRDYqFFDSGQKKQIVNQLFK--EKR-KVTE 562
 WP_044674937 487 SAENFIRMTNYDQYLPDQKVLPKHSLLYKFAVYNELTKVKFIA--EGMRDYqFLDSDGQKKDIVKTLFK--TKR-KVTA 561
 WP_044676715 487 SAENFIRMTNYDQYLPDQKVLPKHSLLYKFAVYNELTKVRYVT--EQKSF-FFDANMKQEIFDGVFK--VYR-KVTK 560
 WP_044680361 487 SAENFIRMTNYDQYLPDQKVLPKHSLLYKFAVYNELTKVRYVT--EQKSF-FFDANMKQEIFDGVFK--VYR-KVTK 560
 WP_044681799 487 SAENFIRMTNYDQYLPDQKVLPKHSLLYKFAVYNELTKVKFIA--EGMRDYqFLDSDGQKKDIVKTLFK--TKR-KVTA 561
 WP_049533112 487 SAEKFI TRMTINDLYLPEEKVLPKHSLLYETFTVYNELTKVKYVN--EQGEAK-FFDANMKQEIFDHVFK--ENR-KVTK 560
 WP_029090905 472 SSEAFIKRMTNKCTYLIHEDVLPKHSFSYAKFEVLNKLKIRLDG-----KP-IDIPLKKRIFEGFLF--EktKVtQ 540
 WP_006506696 499 TAEGFIKRMRSYCTYFPDEEVLPKNSLIVSKYEVYNELNKIRVDD-----kLLEVDVKNDIYNELFM--KNK-TVTE 567
 AIT42264 487 SAQSFIERMTNFDKMLPNEKVLPKHSLLYETFTVYNELTKVKYVT--EGMRKPaFLSGEQKKAIVDILLFK--TNR-KVTV 561
 WP_034440723 494 SAELFIENLTSRDTYLPDPEVLPKRSLLIYQKFTIFNELTKISYID--ERGILQ-NFSSREKIAIFNDLFLK--NksKVTK 567
 AKQ21048 487 SAQSFIERMTNFDKMLPNEKVLPKHSLLYETFTVYNELTKVKYVT--EGMRKPaFLSGEQKKAIVDILLFK--TNR-KVTV 561
 WP_004636532 489 SAEKFIERMTNMDTYLLEEKVLPKRSLLYQTFEVYNELTKVRYTN--EQKTE-KLNRQQAIEITLFLK-qKNR--VRE 562
 WP_002364836 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_016631044 447 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 520
 EMS75795 232 SAMKFIQRMINYDLYLPTEKVLPKHSLLYQKTYIFNELTKVAYKD--ERGIKH-QFSSKEKEKIFKELFQ--QOR-KVTV 305
 WP_002373311 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_002378009 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_002407324 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_002413717 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_010775580 498 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 571
 WP_010818269 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_010824395 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_016622645 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_033624816 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_033625576 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_033789179 496 SATAFIERMTNFDTYLPSEKVLPKHSLLYKFMVFNELTKISYTD--DRGIKA-NFSGKEKEKIFDYLFK--TRR-KVKK 569
 WP_002310644 493 SAVRFIERMINTDMPHNKVLPKNSLLYQKFSIYNELTKVRYQD--ERGQMN-YFSSIEKKEIFHELFE--KNR-KVTK 566
 WP_002312694 494 SAVRFIERMINTDMPHNKVLPKNSLLYQKFSIYNELTKVRYQD--ERGQMN-YFSSIEKKEIFHELFE--KNR-KVTK 567
 WP_002314015 494 SAVRFIERMINTDMPHNKVLPKNSLLYQKFSIYNELTKVRYQD--ERGQMN-YFSSIEKKEIFHELFE--KNR-KVTK 567
 WP_002320716 494 SAVRFIERMINTDMPHNKVLPKNSLLYQKFSIYNELTKVRYQD--ERGQMN-YFSSIEKKEIFHELFE--KNR-KVTK 567

WP_002330729 493 SAVRFIERMINTDMYI PHNKVLPKNSLLYQKFSIYNELTKVRYQD--ERGQMN-YFSSIEKKEIFHELFE--KNR-KVTK 566
 WP_002335161 494 SAVRFIERMINTDMPHNKVLKNSLLYQKFSIYNELTKVRYQD--ERGQMN-YFSSIEKKEIFHELFE--KNR-KVTK 567
 WP_002345439 494 SAVRFIERMINTDMYI PHNKVLPKNSLLYQKFSIYNELTKVRYQD--ERGQMN-YFSSIEKKEIFHELFE--KNR-KVTK 567
 WP_034867970 487 SAMRFIQRMTKQDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--ERGVKQ-YFSGDEKQOIFKQLFQ--KERgKITV 561
 WP_047937432 494 SAVRFIERMINTDMPHNKVLKNSLLYQKFSIYNELTKVRYQD--ERGQMN-YFSSIEKKEIFHELFE--KNR-KVTK 567
 WP_010720994 487 SAMRFIQRMTKQDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--ERGVKQ-YFSGDEKQOIFKQLFQ--KERgKITV 561
 WP_010737004 487 SAMRFIQRMTKQDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--ERGVKQ-YFSGDEKQOIFKQLFQ--KERgKITV 561
 WP_034700478 487 SAMRFIQRMTKQDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--ERGVKQ-YFSGDEKQOIFKQLFQ--KERgKITV 561
 WP_007209003 491 SAVAFIERMTIKDIYL-NENVLPRHSLIYEKFTVFNELTKVRYQD--DRGVFQ-RFSAEEKEDI FEKLFK--SER-KVTK 563
 WP_023519017 486 SAIEFIERMTNQDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--ERGVKQ-YFSGDEKQOIFKQLFQ--KERgKITV 561
 WP_010770040 490 SATKFIERMINTDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--ERGMNQ-RFSGEEKQOIVEELFK--QSR-KVTK 563
 WP_048604708 486 SAELFIERMTNFDLYLSEKVLKNSLLYQKFSIYNELTKVRYQD--EQGKVQ-NFSSEEKERIFIDLFK--QHR-KVTK 559
 WP_010750235 487 SATKFIQRMINDYDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DRGIKH-QFSSEEKERIFIDLFK--QHR-KVTK 560
 AII16583 526 SAQSFIERMINTDKNLPEKVLKNSLLYQKFSIYNELTKVRYQD--EGMRKPaFLSGEQKKAIVDILLFK--TNR-KVTV 600
 WP_029073316 507 TADEFIKRMNFCTYFPDEPVLAKNSLTVSKYEVLEINLKRIND-----hLIIKRDIKDKMLHTLFM--DHK-SISA 575
 WP_031589969 507 TADEFIKRMNFCTYFPDEPVMKNSLTVSKYEVLEINLKRIND-----hLIIKRDMKDKMLHTLFM--DHK-SISA 575
 KDA45870 485 SAEDFIKRMINDLYLPEPVLKNSLLYQKFSIYNELTKVRYQD--ENGEAK-YFDAQTKRSIFE-LFKL--DR-KVSE 557
 WP_039099354 510 SANEFIKRMTTDTYLLAEDVLPKNSLLYQKFSIYNELTKVRYQD--QPITTE-----LKQAITDLMFM--QKtSVTV 578
 AKP02966 484 SSNKFIERMINTDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTS-YFSGQEKQOIFNDLFF--QKR-KVTK 560
 WP_010991369 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTS-YFSGQEKQOIFNDLFF--QKR-KVTK 566
 WP_033838504 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTS-YFSGQEKQOIFNDLFF--QKR-KVTK 566
 EHN60060 496 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTS-YFSGQEKQOIFNDLFF--QKR-KVTK 569
 EFR89594 262 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTS-YFSGQEKQOIFNDLFF--QKR-KVTK 335
 WP_038409211 493 SAIDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTH-HFSGQEKQOIFNGLFK--QQR-KVKK 566
 EFR95520 112 SAIDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTH-HFSGQEKQOIFNGLFK--QQR-KVKK 185
 WP_003723650 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVKK 566
 WP_003727705 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVKK 566
 WP_003730785 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVKK 566
 WP_003733029 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVKK 566
 WP_003739838 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVSK 566
 WP_014601172 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVKK 566
 WP_023548323 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVKK 566
 WP_031665337 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVKK 566
 WP_031669209 493 SAVDFIEKMTNKDTPTEKVLKNSLLYQKFSIYNELTKVRYQD--DQKTN-YFSGREKQOIFNDLFF--QKR-KVKK 566

WP_033920898	493	SAVD	FI	EKMT	NKDT	YLP	PKEN	VLP	PKHS	SLCY	QKYM	VYNEL	TKIR	YID--	DQ	GK	TN--	YF	SG	QEK	QOI	FND	L	FK--	Q	KR--	K	V	K	K	566					
AKI42028	496	SAVD	FI	EKMT	NKDT	YLP	PKEN	VLP	PKHS	SLCY	QKYM	VYNEL	TKVR	YID--	DQ	GK	TN--	YF	SG	QEK	QOI	FND	L	FK--	Q	KR--	K	V	K	K	569					
AKI50529	496	SAVD	FI	EKMT	NKDT	YLP	PKEN	VLP	PKHS	SLCY	QKYM	VYNEL	TKIR	YID--	DQ	GK	TN--	YF	SG	QEK	QOI	FND	L	FK--	Q	KR--	K	V	K	K	569					
EFR83390	1	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	14					
WP_046323366	493	SAID	FI	EKMT	NKDT	YLP	PKEN	VLP	PKHS	MCY	QKYM	VYNEL	TKIR	YTD--	DQ	GK	TH--	YF	SG	QEK	QOI	FND	L	FK--	Q	KR--	K	V	K	K	566					
AKE81011	503	SAQ	S	FI	ERM	TN	FD	KN	LP	PKHS	LLY	EY	FT	VY	NEL	TKV	KY	VT--	EG	MR	KPa	FL	SG	EQ	KKAI	VD	LL	FK--	T	N	R--	K	V	T	V	577
CUO82355	503	TAEG	FI	KRM	RS	YCT	YFP	DEE	VLP	PKNS	LI	VSK	YEV	YNEL	NKIR	VDD	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	571		
WP_033162887	504	TAEG	FI	ERM	KNT	GT	YFP	DE	PV	MA	KNS	LT	VSK	FE	VL	NEL	NKIR	I	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	572		
AGZ01981	520	SAQ	S	FI	ERM	TN	FD	KN	LP	PKHS	LLY	EY	FT	VY	NEL	TKV	KY	VT--	EG	MR	KPa	FL	SG	EQ	KKAI	VD	LL	FK--	T	N	R--	K	V	T	V	594
AKA60242	487	SAQ	S	FI	ERM	TN	FD	KN	LP	PKHS	LLY	EY	FT	VY	NEL	TKV	KY	VT--	EG	MR	KPa	FL	SG	EQ	KKAI	VD	LL	FK--	T	N	R--	K	V	T	V	561
AKS40380	487	SAQ	S	FI	ERM	TN	FD	KN	LP	PKHS	LLY	EY	FT	VY	NEL	TKV	KY	VT--	EG	MR	KPa	FL	SG	EQ	KKAI	VD	LL	FK--	T	N	R--	K	V	T	V	561
4UN5_B	491	SAQ	S	FI	ERM	TN	FD	KN	LP	PKHS	LLY	EY	FT	VY	NEL	TKV	KY	VT--	EG	MR	KPa	FL	SG	EQ	KKAI	VD	LL	FK--	T	N	R--	K	V	T	V	565

WP_010922251	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_039695303	564	EKLLNLYLNKE--FPEYRIKDLIGLDKENkSFNASLGTYHDLKIL-DK	AFLDDKVNEEVIEDIIKTLTLFEDKDMIH	637
WP_045635197	562	KDIIHYLHN--VDGYDGIELKIEKQ--FNASLSTYHDDLKIIKDK	EFMDDAKNEAILENIVHTLTLFEDREMIE	632
5AXW_A	300	KQIAKEIILVNe--EDIKGYRVTSTGKPe---FTNLKVYHDIKIDITARK	-----ENAEALLDQIAKILTIYQSSEDIQ	368
WP_009880683	246	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	317
WP_010922251	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_011054416	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_011284745	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_011285506	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_011527619	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_012560673	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_014407541	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_020905136	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_023080005	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_023610282	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_030125963	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_030126706	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_031488318	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_032460140	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_032461047	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_032462016	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_032462936	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_032464890	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_033888930	387	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	458
WP_038431314	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_038432938	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_038434062	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
BAQ51233	473	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	544
KGE60162		-----	-----	
KGE60856		-----	-----	
WP_002989955	562	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDDLKIIKDK	DFLDNEENEDIILEDIVLTLTLFEDREMIE	633
WP_003030002	561	DKLLNLYLNKE--FEEFRIVNLTGLDKENkAFNSSLGTYHDLRKL-DK	SFLDDKANAKTIEDI IQTLTLFEDREMIR	634
WP_003065552	564	EKLLNLYLNKE--FPEYRIKDLIGLDKENkSFNASLGTYHDLKIL-DK	AFLDDKVNEEVIEDIIKTLTLFEDKDMIH	637
WP_001040076	563	KDIIISFLNK--VDGYEGIAIKIEKQ---FNASLSTYHDDLKIL-GK	DFLDNTDNEILEDIVLTLTLFEDREMIE	632

WP_001040078 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040080 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040081 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040083 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIQTLTLFEDREMİK 635
 WP_001040085 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
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 WP_001040092 562 KQLDFLAKE--FEEFRIVDVTGLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040094 563 KDII SFLNK--VDGYEGIAIKGIEKQ--FNASLSTYHDLKKIL-GK DFLDNTDNEIILEDIVQTLTLFEDREMİR 632
 WP_001040095 563 KDII SFLNK--VDGYEGIAIKGIEKQ--FNASLSTYHDLKKIL-GK DFLDNTDNEIILEDIVQTLTLFEDREMİR 632
 WP_001040096 563 KDII SFLNK--VDGYEGIAIKGIEKQ--FNASLSTYHDLKKIL-GK DFLDNTDNEIILEDIVQTLTLFEDREMİR 632
 WP_001040097 563 KDII SFLNK--VDGYEGIAIKGIEKQ--FNASLSTYHDLKKIL-GK DFLDNTDNEIILEDIVQTLTLFEDREMİR 632
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 WP_001040099 563 KDII SFLNK--VDGYEGIAIKGIEKQ--FNASLSTYHDLKKIL-GK DFLDNTDNEIILEDIVQTLTLFEDREMİR 632
 WP_001040100 563 KDII SFLNK--VDGYEGIAIKGIEKQ--FNASLSTYHDLKKIL-GK DFLDNTDNEIILEDIVQTLTLFEDREMİR 632
 WP_001040104 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040105 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040106 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040107 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040108 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040109 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_001040110 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_015058523 562 KQLDFLAKE--FEEFRIVDVTGLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_017643650 563 KDII SFLNK--VDGYEGIAIKGIEKQ--FNASLSTYHDLKKIL-GK DFLDNTDNEIILEDIVQTLTLFEDREMİR 632
 WP_017647151 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_017648376 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_017649527 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_017771611 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 WP_017771984 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 CFQ25032 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635
 CFV16040 562 KLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGTYHDLKKIL-DK DFLNDPNESILEDIVQTLTLFEDREMİK 635

KLJ37842	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
KLJ72361	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
KL20707	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
KL42645	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
WP_047207273	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
WP_047209694	563	KDII SFLNK--VDGYEGIAKIEKQ--FNASLSTYHDLKIL-GK	DFLDNPDNESILEDIVQTLTLFEDREMIR	632
WP_050198062	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
WP_050201642	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
WP_050204027	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
WP_050881965	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
WP_050886065	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
AHN30376	562	KQLLDFLAKE--FEEFRIVDVTGLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
EAO78426	562	KKLLDFLAKE--YEEFRIVDVI GLDKEnkAFNASLGT YHDLKIL-DK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
CCW42055	562	KQLLDFLAKE--FEEFRIVDVTGLDKEnkAFNASLGT YHDLKIL-GK	DFLDNPDNESILEDIVQTLTLFEDREMIR	635
WP_003041502	561	DKLLNLYLNKE--FEEFRIVNLTGLDKEnkVFNSSLGT YHDLRKIL-NK	SFLDNKANAQIIEIIQTLTLFEDREMIR	634
WP_037593752	562	DKLLNLYLNKE--FEEFRIVNLTGLDKEnkAFNSSLGT YHDLRKIL-DK	SFLDDKANAKTIEIIQTLTLFEDREMIR	635
WP_049516684	562	DKLLNLYLNKE--FEEFRIVNLTGLDKEnkAFNSSLGT YHDLRKIL-DK	SFLDDKANAKTIEIIQTLTLFEDREMIR	635
GAD46167	561	DKLLNLYLNKE--FEEFRIVNLTGLDKEnkAFNSSLGT YHDLRKIL-DK	SFLDDKANAKTIEIIQTLTLFEDREMIR	634
WP_018363470	562	EKLLNYLDKE--FPEYRIQDLVGLDKEnkSFNASLGT YHDLKIL-DK	SFLDDKANEKTIIEIIQTLTLFEDREMIQ	635
WP_003043819	572	KQLKEDYFKK--IECFDSVEIIGVEDR--FNASLGT YHDLKIL-DK	DFLDNEENEDILEDIVLTLTLFEDREMIE	643
WP_006269658	561	DKLLNLYLNKE--FEEFRIVNLTGLDKEnkAFNSSLGT YHDLRKIL-DK	SFLDDKANAKTIEIIQTLTLFEDREMIR	634
WP_048800889	561	DKLLNYLDKE--FDEFRIVDLTGLDKEnkAFNASLGT YHDLRKIL-DK	SFLDDKANAKTIEIIQTLTLFEDREMIR	634
WP_012767106	562	KQLKEDYFKK--IECFDSVEIISGVEDR--FNASLGT YHDLKIL-KIKDK	DFLDNEENEDILEDIVLTLTLFEDKEMIE	633
WP_014612333	562	KQLKEDYFKK--IECFDSVEIISGVEDR--FNASLGT YHDLKIL-KIKDK	DFLDNEENEDILEDIVLTLTLFEDKEMIE	633
WP_015017095	562	KQLKEDYFKK--IECFDSVEIISGVEDR--FNASLGT YHDLKIL-KIKDK	DFLDNEENEDILEDIVLTLTLFEDKEMIE	633
WP_015057649	562	KQLKEDYFKK--IECFDSVEIISGVEDR--FNASLGT YHDLKIL-KIKDK	DFLDNEENEDILEDIVLTLTLFEDKEMIE	633
WP_048327215	562	KQLKEDYFKK--IECFDSVEIISGVEDR--FNASLGT YHDLKIL-KIKDK	DFLDNEENEDILEDIVLTLTLFEDKEMIE	633
WP_049519324	562	KQLKEDYFKK--IECFDSVEIISGVEDR--FNASLGT YHDLKIL-KIKDK	DFLDNEENEDILEDIVLTLTLFEDKEMIE	633
WP_012515931	562	KQLKENYFKK--IECWDSVEITGVEDS--FNASLGT YHDLKIL-KIQDK	DFLDNPDNQKIIIEIILLTLTLFEDKKMIS	633
WP_021320964	562	KQLKENYFKK--IECWDSVEITGVEDS--FNASLGT YHDLKIL-KIQDK	DFLDNPDNQKIIIEIILLTLTLFEDKKMIS	633
WP_037581760	562	KQLKENYFKK--IECWDSVEITGVEDS--FNASLGT YHDLKIL-KIQDK	DFLDNPDNQKIIIEIILLTLTLFEDKKMIS	633
WP_004232481	561	AKLLSYLNNE--FEEFRINDLI GLDKDsKsFNASLGT YHDLKIL-DK	SFLDDKTNEQIIEIIVLTLTLFEDRDMIH	634
WP_009854540	562	EKLLNLYLNKE--FPEYRIKDLI GLDKEnkSFNASLGT YHDLKIL-DK	AFLDDKANEKTIIEIILLTLTLFEDKDMIH	635
WP_012962174	562	DKFLNLYLNKE--FPEYRIQDLI GLDKEnkSFNASLGT YHDLKIL-DK	SFLDDKTNETIIEIILLTLTLFEDRDMIR	635

WP_039695303	564	EKLLNLYLNKE--FPEYRIKDLI GLDKEnkSFNASLGTYHDLCKIL-DK	AFLDDKVNVEEVIEDI IKTLTLFEDKDMIH	637
WP_014334983	561	AKLLSYLNNE--FEFRINDLIGLDDKsKFNASLGTYHDLCKIL-DK	SFLDDKTNGQI IEDIVLTLTLFEDRDMIH	634
WP_003099269	562	KQLKEEYFSK--MKCFHTVTILGVEDR--FNASLGTYHDLCKILFKDK	AFLDDEANQDILEEIVWTLTLFEDQAMIE	633
AHY15608	562	KQLKEEYFSK--MKCFHTVTILGVEDR--FNASLGTYHDLCKILFKDK	AFLDDEANQDILEEIVWTLTLFEDQAMIE	633
AHY17476	562	KQLKEEYFSK--MKCFHTVTILGVEDR--FNASLGTYHDLCKILFKDK	AFLDDEANQDILEEIVWTLTLFEDQAMIE	633
ESR09100		-----	-----	
AGM98575	562	KQLKEEYFSK--MKCFHTVTILGVEDR--FNASLGTYHDLCKILFKDK	AFLDDEANQDILEEIVWTLTLFEDQAMIE	633
ALF27331	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_018372492	574	RKLKDFIEKElgYIDIDNIKGVEEQ--FNASYTTYQDLCKILGDK	EFLDNEENKDLLEEIIYILTVFEDRKMIE	647
WP_045618028	563	KDIIQYLHN--VDSYDGIELKIEKQ--FNASLSTYHDLCKILKIKDK	EFMDDSKNEAILENIVHTLTLIFEDREMIR	633
WP_045635197	562	KDIIHYLHN--VDGYDGIELKIEKQ--FNASLSTYHDLCKILKIKDK	EFMDDAKNEAILENIVHTLTLIFEDREMIR	632
WP_002263549	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002263887	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002264920	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002269043	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002269448	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002271977	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002272766	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002273241	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002275430	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002276448	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002277050	561	KKLRTFLDKN--FDEFRIVDIQGLDKeteTFNASYATYQDLCKILKVIKDK	VFMDNPENAEILENIVLTLTLFEDREMIR	635
WP_002277364	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002279025	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002279859	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002280230	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002281696	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002282247	561	KKLRTFLDKN--FDEFRIVDIQGLDKeteTFNASYATYQDLCKILKVIKDK	VFMDNPENAEILENIVLTLTLFEDREMIR	635
WP_002282906	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002283846	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002287255	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002288990	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002289641	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkVFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634
WP_002290427	561	DKLMDFLEKE--FDEFRIVDLTGLDKEnkAFNASYGTYHDLCKIL-DK	DFLDNSKNEKILEDIVLTLTLFEDREMIR	634

WP_002295753	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkVFNASyGTYHDLCKIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_002296423	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_002304487	571	DKLLNLYLNKE--FEFRIVNLTGLDKEkVFNSSLGTYHDLRkIL-NK	SFLDNkENEQIIEdIQTLTLFEDREMIr	644
WP_002305844	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkVFNASyGTYHDLCKIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_002307203	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_002310390	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_002352408	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_012997688	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_014677909	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_019312892	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_019313659	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkVFNASyGTYHDLCKIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_019314093	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_019315370	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_019803776	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkVFNASyGTYHDLCKIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_019805234	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_024783594	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkVFNASyGTYHDLCKIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_024784288	561	KKLRTFLDKN--FDEFRIVDLQGLDKEteTFNASyATYQDLLKVIKDK	VFMdNPENAEILENIENIVHTTLTLFEDREMIr	635
WP_024784666	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_024784894	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkVFNASyGTYHDLCKIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_024786433	561	KKLRTFLDKN--FDEFRIVDLQGLDKEteTFNASyATYQDLLKVIKDK	VFMdNPENAEILENIENIVHTTLTLFEDREMIr	635
WP_049473442	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkVFNASyGTYHDLCKIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
WP_049474547	561	DKLMDFLEKE--FDEFRIVDLTGLDKEkVFNASyGTYHDLCKIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	634
EMC03581	554	DKLMDFLEKE--FDEFRIVDLTGLDKEkAFNASyGTYHDLRkIL-DK	DFLDnskEKILEdIVLTLTLFEDREMIr	627
WP_000428612	565	KDIIQYLHT--VDGYDGIELKGI EKQ--FNASLSTYHDLKIIKDK	EFMDdPNNEEILENIENIVHTTLTLFEDREMIr	635
WP_000428613	563	KDIIQFLHN--VDGYDGIELKGI EKQ--FNASLSTYHDLKIIKDK	EFMDdSKNEEILENIENIVHTTLTLFEDREMIr	633
WP_049523028	562	KDIIHYLHN--VDGYDGIELKGI EKH--FNSSLSTYHDLKIIKDK	EFMDdPKNEEILENIENIVHTTLTLFEDRVMIr	632
WP_003107102	531	KQLKENYFNK--IRCLDSITISGVEDK--FNASLGTyHDLNIIKNQ	KILdDEQnQDSLEDIVLTLTLFEDEKMIr	602
WP_054279288	564	KQLKEDFFSK--IECFDVTDISGVEDK--FNASLGTyHDLKIIKDK	AFLDNSENIENIEdIILTLTLFEDKEMIA	635
WP_049531101	563	KDLIHYLHN--VDGYDGIELKGI EKQ--FNASLSTYHDLKIIKDK	RFMDPEKQEIENIVHTTLTLFEDREMIr	633
WP_049538452	563	KDIIQYLHN--VDGYDGIELKGI EKQ--FNASLSTYHDLKIIKDK	EFMDdSKNEEILENIENIVHTTLTLFEDREMIr	633
WP_049549711	563	KDIIHYLHT--VDGYDGIELKGI EKQ--FNASLSTYHDLKIIKDK	EFMDdSKNEALENIENIVHTTLTLFEDREMIr	633
WP_007896501	565	KDLKEKYFSQ--IEGLENVdVTGVEGA--FNASLGTyNDLIIKIIKDK	AFLdDEANAEILEEIVLTLTLFQDEKLI E	636
EFR44625	517	KDLKEKYFSQ--IEGLENVdVTGVEGA--FNASLGTyNDLIIKIIKDK	AFLdDEANAEILEEIVLTLTLFQDEKLI E	588
WP_002897477	562	KDIIHYLHN--VDGYDGIELKGI EKQ--FNANLSTYHDLKIIKDK	EFMDdPKNEEILENIENIVHTTLTLFEDREMIr	632

WP_002906454	562	KDIIHYLHN---VDGYDGIELKIEKQ---	FNASLSTYHDDLKIIKDK	EFMDNPKNGEILENIHHTLTI	FEDREMIK	632
WP_009729476	563	KDIIQFLHN---VDGYDGIELKIEKQ---	FNASLSTYHDDLKIIKDK	AFMDDAKNEAILENIVHTLTI	FEDREMIK	633
CQR24647	562	KKILNFDKN--FDEFRITDIQGLDNETg	FNASYGTYHDDLKIIIGDK	EFMSSDNVDVLEDIVLSLTL	FEDREMIK	636
WP_000066813	565	KDIIHYLHN---VDGYDGIELKIEKQ---	FNASLSTYHDDLKIIKDK	AFMDDSKNEEILENIHHTLTI	FEDREMIK	635
WP_009754323	563	KDIIHYLHN---VDGYDGIELKIEKQ---	FNASLSTYHDDLKIIKDK	EFMDHNKQEIENIVHTLTI	FEDREMIK	633
WP_044674937	562	KDIKAYL-EN--SNGYAGVELKLEEQ---	FNASLPTYHDDLKILRDK	AFIDAEENQEILEDIVLTLTL	FEDREMIK	632
WP_044676715	561	EKLMDFLGKE--FDEFRIVDLGLDKDnK	SFNASLGTYHDDLKIIKDK	DLNPNEDILENVLTLTL	FEDREMIK	634
WP_044680361	561	EKLMDFLGKE--FDEFRIVDLGLDKDnK	SFNASLGTYHDDLKIIKDK	DLNPNEDILENVLTLTL	FEDREMIK	634
WP_044681799	562	KDIKAYL-EN--SNGYAGVELKLEEQ---	FNASLPTYHDDLKILRDK	AFIDAEENQEILEDIVLTLTL	FEDREMIK	632
WP_049533112	561	DKLLNYLGKE--FDEFRIVDLGLDKDnK	SFNASLGTYHDDLKIL- DK	SFLDNKENEQIIEDIIQTLTL	FEDREMIK	634
WP_029090905	541	TSLKKWLAEH--EHMTVSVVQGTQKET-	EFATSLQAFHRFVKIF-DR	ETVSNPANEEMFEKIIYWS	VFFEDKKIMR	612
WP_006506696	568	KKLKNWLVANNqCS--KDAEIKGFQKEn-	QFSTSLTPWIDFTNI	FGKI	---	637
AIT42264	562	KQLKEDYFKK--IECFDSVEISGVEDR-	---FNASLGTYHDDLKIIKDK	DFLDNEENEDILEDIVLTLTL	FEDREMIK	633
WP_034440723	568	NQLVKYIENK--EQIIAPEIKGIEDS---	FNSNYSTYIDLSKI	PDMK	---	637
AKQ21048	562	KQLKEDYFKK--IECFDSVEISGVEDR-	---FNASLGTYHDDLKIIKDK	DFLDNEENEDILEDIVLTLTL	FEDREMIK	633
WP_004636532	563	KDIANYLEQ---YGYVDGTDIKGVEDK---	FNASLSTYNDLAKIDGAK	AYLDDPEYADVWEDIIKILTI	FEDKAMRK	633
WP_002364836	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_016631044	521	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	592
EMS75795	306	KKLQFLSAN--YN-IEDAEILGVDKA---	FNSSYATYHDFDLAKPN	ELLEQPEMNAMFEDIVKILTI	FEDREMIK	381
WP_002373311	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_002378009	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_002407324	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_002413717	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_010775580	572	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	643
WP_010818269	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_010824395	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_016622645	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_033624816	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_033625576	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_033789179	570	KDIIQFYRNE--YN-TEIVTLSGLEED---	QFNASFSTYQDLLKCGLTR	AELDHPDNAEKLEDDIIKILTI	FEDRQIR	641
WP_002310644	567	KDLQEFLYLK--YD-IKHAELSGIEKA---	FNASYTYHDFLITMSENK	QWLEDPELASMFEI	IKTLTVFEDREMIK	641
WP_002312694	568	KDLQEFLYLK--YD-IKHAELSGIEKA---	FNASYTYHDFLITMSENK	QWLEDPELASMFEI	IKTLTVFEDREMIK	642
WP_002314015	568	KDLQEFLYLK--YD-IKHAELSGIEKA---	FNASYTYHDFLITMSENK	QWLEDPELASMFEI	IKTLTVFEDREMIK	642
WP_002320716	568	KDLQEFLYLK--YD-IKHAELSGIEKA---	FNASYTYHDFLITMSENK	QWLEDPELASMFEI	IKTLTVFEDREMIK	642

WP_002330729	567	KDLQEFLYLK--YD-IKHAELSGIEKA---FNASYTTYHDFLITMSENK	QWLEDEPELASMFEIEIKTLTVFEDREMİK	641
WP_002335161	568	KDLQEFLYLK--YD-IKHAELSGIEKA---FNASYTTYHDFLITMSENK	QWLEDEPELASMFEIEIKTLTVFEDREMİK	642
WP_002345439	568	KDLQEFLYLK--YD-IKHAELSGIEKA---FNASYTTYHDFLITMSENK	QWLEDEPELASMFEIEIKTLTVFEDREMİK	642
WP_034867970	562	KKLQFLYTH--YH-IENAQIFGIEKA---FNASYTTYHDFLITMSENK	EWLEQPEMEPIFEDIVKILITIFEDRQMIK	637
WP_047937432	568	KDLQEFLYLK--YD-IKHAELSGIEKA---FNASYTTYHDFLITMSENK	QWLEDEPELASMFEIEIKTLTVFEDREMİK	642
WP_010720994	562	KKLQFLYTH--YH-IENAQIFGIEKA---FNASYTTYHDFLITMSENK	EWLEQPEMEPIFEDIVKILITIFEDRQMIK	637
WP_010737004	562	KKLQFLYTH--YH-IENAQIFGIEKA---FNASYTTYHDFLITMSENK	EWLEQPEMEPIFEDIVKILITIFEDRQMIK	637
WP_034700478	562	KKLQFLYTH--YH-IENAQIFGIEKA---FNASYTTYHDFLITMSENK	EWLEQPEMEPIFEDIVKILITIFEDRQMIK	637
WP_007209003	564	KKLENYLRIEL---SISPPVKIEEQ---FNANFGTYLCLKFDELH	PYLDDEKYQDTLEEVIVKILITIFEDRSMIQ	634
WP_023519017	560	KQLRKFLLEIN--EQ-IDSTEIKGIETS---FNASYTTYHDLKLS---	TLDDDPDMTTFEEIIEKILITIFEDREMİR	631
WP_010770040	564	KLLEKFLSNE--FG-LVDVAIKGIE-T--SFNAGYGYHDFLKGITR	EQLDKEENSEITLLEIVKILITIFEDRKMİR	634
WP_048604708	560	KDLSNFLRNE--YN-LDDVLDVVAIKGIE-T--KFNASFNTHDFLKLKIDP	KVLDPPANEPMEFEEIVKILITIFEDRKMİR	630
WP_010750235	561	KKLQHFLSAN--YN-IEDAEILGVDKV---FNSSYATYHDFLELAKPY	ELLEQPEMEEMFEDIVKILITIFEDREMVR	636
AI116583	601	KQLKEDYFKK--IECFDSVEISGVEDR---FNASLGTYHDLKLIKDK	DFLDNEENEDILEDIVLTLTLFEDREMIE	672
WP_029073316	576	NAMKKWLKVNqyFSNTDDIKIEGFQKEN-ACSTSLTPWIDFTKIFGEI	----NNSNYELIEKIIYDVTVFEDKKILR	647
WP_031589969	576	NAMKKWLKVNqyFSNTDDIKIEGFQKEN-ACSTSLTPWIDFTKIFGKI	----NESNYDFIEKIIYDVTVFEDKKILR	647
KDA45870	558	KMVIKHLKV--MPAIRIQALKGLDNGK--FNASYGYKDLVDMGVAP	ELLNDEVNSEKWEDEIIEKTLIFEGRKLİK	630
WP_039099354	579	KNIQDYLVSEK--RYASRPAITGLSDENK-FNSRLSTYHDLKTIIVGDA	--VDDVKQADLEKCIEWSTIFEDGKIYS	650
AKP02966	561	KKLTWLIQAQg--YKNPILIGLSQKd-EFNSTLTTTYLDMKKIFGSS	-FMENKNYNQIEELIEWLITIFEDKQILN	632
WP_010991369	567	KDLELFLRNM--SH-VEPTIEGLE-D--SFNSSYSTYHDLKLVGİKQ	EILDNPVNTMELENIIVKILITVFEDKRMİK	637
WP_033838504	567	KDLELFLRNM--SH-VEPTIEGLE-D--SFNSSYSTYHDLKLVGİKQ	EILDNPVNTMELENIIVKILITVFEDKRMİK	637
EHN60060	570	KDLELFLRNM--SH-VEPTIEGLE-D--SFNSSYSTYHDLKLVGİKQ	EILDNPVNTMELENIIVKILITVFEDKRMİK	640
EFR89594	336	KDLELFLRNM--SH-VEPTIEGLE-D--SFNSSYSTYHDLKLVGİKQ	EILDNPVNTMELENIIVKILITVFEDKRMİK	406
WP_038409211	567	KDLERFLYTI--NH-IESPTIEGLE-D--AFNSSFATYHDLQKGGVTQ	EILDNPLNADMLEEIVKILITVFEDKRMİK	637
EFR95520	186	KDLERFLYTI--NH-IESPTIEGLE-D--AFNSSFATYHDLQKGGVTQ	EILDNPLNADMLEEIVKILITVFEDKRMİK	256
WP_003723650	567	KDLELFLRNI--NH-IESPTIEGLE-D--SFNASYATYHDLKLVGMKQ	EILDNPLNTEMLEDIVKILITVFEDKPMİK	637
WP_003727705	567	KDLELFLRNI--NH-IESPTIEGLE-D--SFNASYATYHDLKLVGMKQ	EILDNPLNTEMLEDIVKILITVFEDKPMİK	637
WP_003730785	567	KDLELFLRNI--NH-IESPTIEGLE-D--SFNASYATYHDLKLVGMKQ	EILDNPLNTEMLEDIVKILITVFEDKPMİK	637
WP_003733029	567	KDLELFLRNI--NQ-IESPTIEGLE-D--SFNASYATYHDLKLVGMKQ	EILDNPLNTEMLEDIVKILITVFEDKPMİK	637
WP_003739838	567	KDLEQFLRNM--SH-IESPTIEGLE-D--SFNSSYATYHDLKLVGİKQ	EVLNPLNTEMLEDIVKILITVFEDKRMİK	637
WP_014601172	567	KDLELFLRNI--NH-IESPTIEGLE-D--SFNSSYATYHDLKLVGMKQ	EILDNPLNTEMLEDIVKILITVFEDKPMİK	637
WP_023548323	567	KDLELFLRNI--NH-VEPTIEGLE-D--SFNASYATYHDLKLVGİKQ	EILDNPLNTEMLEDIVKILITVFEDKRMİK	637
WP_031665337	567	KDLELFLRNI--NQ-IESPTIEGLE-D--SFNASYATYHDLKLVGMKQ	EILDNPLNTEMLEDIVKILITVFEDKRMİK	637
WP_031669209	567	KDLELFLRNI--NQ-IESPTIEGLE-D--SFNASYATYHDLKLVGMKQ	EILDNPLNTEMLEDIVKILITVFEDKRMİK	637

WP_033920898	567	KDLELFLRNI--NH-VESPTIEGLE-D--SFNASYATYHDLMKVGIKQ	EILDNPLNTEMLEDIVKILTVFEDKRMK	637
AKI42028	570	KDLELFLRNI--NH-IESPTIEGLE-D--SFNASYATYHDLKVGKMQ	EILDNPLNTEMLEDIVKILTVFEDKPMK	640
AKI50529	570	KDLELFLRNI--NH-VESPTIEGLE-D--SFNASYATYHDLMKVGIKQ	EILDNPLNTEMLEDIVKILTVFEDKRMK	640
EFR83390	15	KDLELFLRNI--NQ-IESPTIEGLE-D--SFNASYATYHDLKVGKMQ	EILDNPLNTEMLEDIVKILTVFEDKRMK	85
WP_046323366	567	KDLELFLYNM--NH-VESPTVEGVE-D--AFNSSFTTYHDLQKVGVPQ	EILDDPLNTEMLEEIIKILTVFEDKRMN	637
AKE81011	578	KQLKEDYFKK--IECFDSVEISGVEDR--FNASLGTYHDLKIIKDK	DFLDNEENEDILEDIVLTLTLFEDREMIE	649
CUO82355	572	KKLKNWLVNNqCCR--KDAEIKGFQKEn-QFSTSLTPWIDFTNIFGKI	----DQSNFDLIEKIIYDLTVFEDKKIMK	641
WP_033162887	573	KKLKWLVTHqYDINEELKIEGYQKDL-QFSTSLAPWIDFTKIFGEI	----NASYQLIEKIIYDISIFEDKKILK	644
AGZ01981	595	KQLKEDYFKK--IECFDSVEISGVEDR--FNASLGTYHDLKIIKDK	DFLDNEENEDILEDIVLTLTLFEDREMIE	666
AKA60242	562	KQLKEDYFKK--IECFDSVEISGVEDR--FNASLGTYHDLKIIKDK	DFLDNEENEDILEDIVLTLTLFEDREMIE	633
AKS40380	562	KQLKEDYFKK--IECFDSVEISGVEDR--FNASLGTYHDLKIIKDK	DFLDNEENEDILEDIVLTLTLFEDREMIE	633
4UN5_B	566	KQLKEDYFKK--IECFDSVEISGVEDR--FNASLGTYHDLKIIKDK	DFLDNEENEDILEDIVLTLTLFEDREMIE	637

WP_010922251	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNF ^Q QLIHDDSL	702
WP_039695303	638	ERLQKYSDI FTANQLKKLER-RHYTGWGRLSYKLLINGIRNK	ENKKTILDYLI	DDG---	SANRNFQLINDDTL	706
WP_045635197	633	QLAQYDSL FDEKVI KALTR-RHYTGWGKLSAKLLINGICDK	QTGNTILDYLI	DDG---	KINRNFQLINDDGL	701
5AXW A	369	EELTNLSEL TQEIEEQI SNLKGYTGTHNLSLKA INLILDE	-----LW	-----	TNDNQIAI FNRLKL	426
WP_009880683	318	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	386
WP_010922251	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_011054416	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_011284745	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_011285506	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_011527619	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_012560673	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_014407541	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_020905136	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_023080005	634	ERLKYANL FDDKVMKQLKR-RHYTGWGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLINDDSL	702
WP_023610282	634	ERLKYANL FDDKVMKQLKR-RHYTGWGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLINDDSL	702
WP_030125963	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_030126706	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_031488318	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_032460140	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_032461047	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_032462016	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_032462936	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_032464890	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_033888930	459	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	527
WP_038431314	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_038432938	634	ERLKYANL FDDKVMKQLKR-RHYTGWGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLINDDSL	702
WP_038434062	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
BAQ51233	545	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	613
KGE60162		-----	-----	-----	-----	
KGE60856		-----	-----	-----	-----	
WP_002989955	634	ERLKYAHLFDDKVMKQLKR-RRYTGWRGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf---	ANRNFQLIHDDSL	702
WP_003030002	635	QLQKYSDI FTKAQLKKLER-RHYTGWGRLSYKLLINGIRNK	ENKKTILDYLI	DDG---	YANRNFQLINDDAL	703
WP_003065552	638	ERLQKYSDI FTADQLKKLER-RHYTGWGRLSYKLLINGIRNK	ENKKTILDYLI	DDG---	SANRNFQLINDDTL	706
WP_001040076	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLLINGIRNK	ENQKTILDYLI	DDG---	SANRNFQLIKDAGL	701

WP_001040078	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040080	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040081	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040083	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040085	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040087	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040088	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040089	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040090	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040091	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040092	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDR	ESQKTILDYLI	SDG---	RANRFMQLINDDGL	704
WP_001040094	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLINGIRNK	ENQKTILDYLI	DDG---	SANRFMQLIKDAGL	701
WP_001040095	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLINGIRNK	ENQKTILDYLI	DDG---	SANRFMQLIKDAGL	701
WP_001040096	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLINGIRNK	ENQKTILDYLI	DDG---	SANRFMQLIKDAGL	701
WP_001040097	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLINGIRNK	ENQKTILDYLI	DDG---	SANRFMQLIKDAGL	701
WP_001040098	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLINGIRNK	ENQKTILDYLI	DDG---	SANRFMQLIKDAGL	701
WP_001040099	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLINGIRNK	ENQKTILDYLI	DDG---	SANRFMQLIKDAGL	701
WP_001040100	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLINGIRNK	ENQKTILDYLI	DDG---	SANRFMQLIKDAGL	701
WP_001040104	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040105	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_001040106	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	SDG---	RANRFMQLIHDDGL	704
WP_001040107	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	SDG---	RANRFMQLIHDDGL	704
WP_001040108	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	SDG---	RANRFMQLIHDDGL	704
WP_001040109	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	SDG---	RANRFMQLIHDDGL	704
WP_001040110	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	SDG---	RANRFMQLIHDDGL	704
WP_015058523	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDR	ESQKTILDYLI	SDG---	RANRFMQLINDDGL	704
WP_017643650	633	KRLDIYKDFFTESQLKKLYR-RHYTGWGRLSAKLINGIRNK	ENQKTILDYLI	DDG---	SANRFMQLIKDAGL	701
WP_017647151	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	KSNRFMQLIHDDGL	704
WP_017648376	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	KSNRFMQLIHDDGL	704
WP_017649527	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
WP_017771611	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	SDG---	RANRFMQLIHDDGL	704
WP_017771984	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
CFQ25032	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704
CFV16040	636	KRLENYKDLFTESQLKKLYR-RHYTGWGRLSAKLINGIRDK	ESQKTILDYLI	DDG---	RSNRFMQLINDDGL	704

KLJ37842	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	704
KLJ72361	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	704
KL20707	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	718
KL242645	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	SDG---	RANRFMQLIHDDGL	704
WP_047207273	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	704
WP_047209694	633	KRLDIYKDFFTESQLKLLYR-RHYTGWGRLSAKLINGIRNK	ENQKTIIDYLI	DDG---	SANRFMQLIKDAGL	701
WP_050198062	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	704
WP_050201642	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	704
WP_050204027	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	SDG---	RANRFMQLIHDDGL	704
WP_050881965	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	704
WP_050886065	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	704
AHN30376	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDR	ESQKIIDYLI	SDG---	RANRFMQLINDDGL	704
EA078426	636	KRLNFKDLFTESQLKLLYR-RHYTGWGRLSAKLINGIRDK	ESQKTIIDYLI	DDG---	RSNRFMQLINDDGL	704
CCW42055	636	KRLDIYKDFFTESQLKLLYR-RHYTGWGRLSAKLINGIRNK	ENQKTIIDYLI	DDG---	SANRFMQLIKDAGL	704
WP_003041502	635	QRLQKYSDI FTKAQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	703
WP_0037593752	636	QRLQKYSDI FTKAQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	704
WP_049516684	636	QRLQKYSDI FTKAQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	704
GAD46167	635	QRLQKYSDI FTKAQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	704
WP_018363470	636	QRLQKYSDI FTKAQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	704
WP_003043819	644	ERLKYAHLFDDKVMKQKLR-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	703
WP_006269658	635	QRLQKYSDI FTKAQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	704
WP_048800889	635	QRLQKYSDI FTKAQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	703
WP_012767106	634	ERLKYANLFDKVMKQKLR-RHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	703
WP_014612333	634	ERLKYANLFDKVMKQKLR-RHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	702
WP_015017095	634	ERLKYAHLFDDKVMKQKLR-RHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	702
WP_015057649	634	ERLKYANLFDKVMKQKLR-RHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	702
WP_048327215	634	ERLKYAHLFDDKVMKQKLR-RHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	702
WP_049519324	634	ERLKYAHLFDDKVMKQKLR-RHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	702
WP_012515931	634	KRLDQYAHLFDKVVLNKLER-HHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	702
WP_021320964	634	KRLDQYAHLFDKVVLNKLER-HHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	702
WP_037581760	634	KRLDQYAHLFDKVVLNKLER-HHYTGWGRLSYKLLINGIRDK	ENKKTIIIDYLI	DDG---	YANRFMQLINDDGL	702
WP_004232481	635	ERLQKYSDI FTSQQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	DANRFMQLINDDGL	703
WP_009854540	636	ERLQKYSDI FTANQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	SANRFMQLINDDGL	704
WP_012962174	636	QRLQKYSDI FTTPQQLKLLER-RHYTGWGRLSYKLLINGIRNK	ENKKTIIIDYLI	DDG---	YANRFMQLISDDTL	704

WP_039695303	638	ERLQKYSDI FTANQLKKLER-RHYTGWGRLSYKLLINGIRNK	ENKKTILDYLI	DDG---	SANRFMQLINDDTL	706
WP_014334983	635	ERLQKYSDFFTSQQLKKLER-RHYTGWGRLSYKLLINGIRNK	ENKKTILDYLI	DDG---	HANRFMQLINDESL	703
WP_003099269	634	RRLVKYADVFEKSVLKKKK-RHYTGWGRLSQKLLINGIKDK	QTGKTILGFLK	-DGV--	ANRFMQLINDSSL	702
AHY15608	634	RRLVKYADVFEKSVLKKKK-RHYTGWGRLSQKLLINGIKDK	QTGKTILGFLK	-DGV--	ANRFMQLINDSSL	702
AHY17476	634	RRLVKYADVFEKSVLKKKK-RHYTGWGRLSQKLLINGIKDK	QTGKTILGFLK	-DGV--	ANRFMQLINDSSL	702
ESR09100		-----	-----	-----	-----	
AGM98575	634	RRLVKYADVFEKSVLKKKK-RHYTGWGRLSQKLLINGIKDK	QTGKTILGFLK	-DGV--	ANRFMQLINDSSL	702
ALF27331	635	KRLNYSDDLTTKEQVKNLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_018372492	648	KRLSELNIPFENKIKKAR-KKYGWGNLSRKLIDGIRNR	ETNRTILGHLI	DDGf--	SNRNLMLINDDGL	716
WP_045618028	634	QLAHYASIFDEKVIKALTR-RHYTGWGKLSAKLINGIYDK	QSKKTILDYLI	DDG---	EINRFMQLINDDGL	702
WP_045635197	633	QLAQYDSLDFEKVVKALTR-RHYTGWGKLSAKLINGICDK	QTGNTILDYLI	DDG---	KINRFMQLINDDGL	701
WP_002263549	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002263887	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002264920	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002269043	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002269448	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002271977	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002272766	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002273241	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002275430	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002276448	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002277050	636	QLAKYADVFDKVIDQLAR-RHYTGWGRLSAKLLINGIRDK	QSKKTIMDYLI	DDA---	QSNRNLMLITDDNL	704
WP_002277364	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002279025	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002279859	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002280230	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002281696	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002282247	636	QLAKYADVFDKVIDQLAR-RHYTGWGRLSAKLLINGIRDK	QSKKTIMDYLI	DDA---	QSNRNLMLITDDNL	704
WP_002282906	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002283846	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002287255	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002288990	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002289641	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002290427	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703

WP_002295753	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002296423	635	KRLKNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002304487	645	QRLQKYSDIFTKAQKKLER-RHYTGWGRLSYKLLINGIRDK	QSNKTILGYLI	DDG---	YSNRNFMQLINDDAL	713
WP_002305844	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002307203	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTLLDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002310390	635	KRLKNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_002352408	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTLLDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_012997688	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_014677909	635	KRLKNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_019312892	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_019313659	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_019314093	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTLLDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_019315370	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTLLDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_019803776	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_019805234	635	KRLKNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_024783594	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_024784288	636	QRLAKYADVFKKVIDQLAR-RHYTGWGRLSAKLLNGIRDK	QSCKTIMDYLI	DDA---	QSNRNLMLITDDNL	704
WP_024784666	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_024784894	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_024786433	636	QRLAKYADVFKKVIDQLAR-RHYTGWGRLSAKLLNGIRDK	QSCKTIMDYLI	DDA---	QSNRNLMLITDDNL	704
WP_049473442	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
WP_049474547	635	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	703
EMC03581	628	KRLNYSDDLTTKEQVKKLER-RHYTGWGRLSAELIHGIRNK	ESRKTILDYLI	DDG---	NSNRNFMQLINDDAL	696
WP_000428612	636	QRLAQYDSLFDKVIKALTR-RHYTGWGKLSKLLINGIRDK	QTKTILDYLM	DDG---	YNNRNFQMLINDDEL	704
WP_000428613	634	QRLAQYDSLFDKVIKALIR-RHYTGWGKLSAKLLIDGICDK	QTKTILDYLI	DDG---	KNNRNFQMLINDDGL	702
WP_049523028	633	QRLNQYDSIFDEKVIKALTR-RHYTGWGKLSAKLLINGIRDK	KTSKTILDYLI	DDG---	YSNRNFMQLINDDGL	701
WP_003107102	603	KRLSKYESIFDPSILKLLKK-RHYTGWGRLSQKLLINGIRDK	QTKTILDFLI	-DGq---	ANRNFQMLINDPFL	671
WP_054279288	636	NRLAVYEDLFDQNLVKQLKR-RHYTGWGRLSKQLINGMRDK	HTGKTILDFLK	-Dgf---	INRNFQMLINDDNL	704
WP_049531101	634	QRLAQYASIFDEKVIKALTR-RHYTGWGKLSAKLLINCIRDR	KTKTILDYLI	DDG---	YNNRNFQMLINDDGL	702
WP_049538452	634	QRLAQYDSIFDEKVIKALTR-RHYTGWGKLSAKLLINGIRDK	QTKTILDYLI	DDG---	YSNRNFMQLINDDGL	702
WP_049549711	634	QRLAQYDSLFDKVIKALTR-RHYTGWGKLSAKLLINGICDK	QTKTILDYLI	DDG---	EINRNFQMLINDDGL	702
WP_007896501	637	KRLAKYANLFEKSVLKKLRK-RHYRGGWGRLSRQLIDGMKDK	ASGKTILDFLK	-DDf---	ANRNFQMLINDSSL	705
EFR44625	589	KRLAKYANLFEKSVLKKLRK-RHYRGGWGRLSRQLIDGMKDK	ASGKTILDFLK	-DDf---	ANRNFQMLINDSSL	657
WP_002897477	633	QRLAQYDTLFDKVIKALTR-RHYTGWGKLSAKLLINGIRDK	QSGKTILDYLI	DDD---	KINRNFQMLINDDGL	701

WP_002906454	633	QRLAQYDTLFDKVIKALTR-RHYTGWGKLSAKLINGIRDK	QTKGTILEYLI	DDG---	DCNRNFMQLINDDGL	701
WP_009729476	634	QRLAQYDSLFDKVIKALTR-RHYTGWGKLSAKLINGISDK	QTKGTILDYLI	DDG---	EINRNFQMLINDDGL	702
QOR24647	637	QRLKYEYEDIFSFKVIANLTR-RHYTGWGRLSACLINGIKDK	HSRKTILDYLI	DDG---	HSNRNFQMLINDDNL	705
WP_000066813	636	QRLAQYDSLFDKVIKALTR-RHYTGWGKLSAKLINGIRDK	KSGKTILDYLI	DDG---	EINRNFQMLIHDDGL	704
WP_009754323	634	QRLAQYDSIFDEKVIKALTR-RHYTGWGKLSAKLINGICDK	KTGKTILDYLI	DDG---	YNNRNFQMLINDDGL	702
WP_044674937	633	KRLEKYKDILTEEQKLLER-RHYTGWGRLSACLINGILDK	VTRKTILGYLI	DDG---	TSNRNFQMLINDDTL	701
WP_044676715	635	KRLEKYKDVLTEEQKLLER-RHYTGWGRLSACLINGIRDK	VTRKTILDYLI	DDG---	TSNRNFQMLINDDTL	703
WP_044680361	635	KRLEKYKDVLTEEQKLLER-RHYTGWGRLSACLINGIRDK	VTRKTILDYLI	DDG---	TSNRNFQMLINDDTL	703
WP_044681799	633	KRLEKYKDILTEEQKLLER-RHYTGWGRLSACLINGILDK	VTRKTILGYLI	DDG---	TSNRNFQMLINDDTL	701
WP_049533112	635	QRLQKYSDIFFKAQLKLLER-CHYTGWGRLSYKLINGIRNK	ENKKTILDYLI	DDG---	YANRNFQMLINDDAL	703
WP_029090905	613	RKLSEYPLTEQQVQLAQV--RFRGWGRLSQRLINRIKTP	EDHKLSINEIL	-----	QTENFMQIIRNKDY	682
WP_006506696	638	RRLKKYALPDDKVQIKL--KYKDWRLSKLLDGIADVAD	SV--TVLDVLE	-----	SRLNLMELIINDKDL	705
AIT42264	634	ERLKYAHLFDDKVMKQLKR-RRYTGWGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf--	ANRNFQMLIHDDSL	702
WP_034440723	638	RQLMKFKDKLSEKAINQLSK-KHYTGWGQLSEKLLINGIRDE	QSNKTILDYLI	DNGcpk	NMNRNFQMLINDDTL	710
AKQ21048	634	ERLKYAHLFDDKVMKQLKR-RRYTGWGRLSRKLLINGIRDK	QSGKTILDFLK	-Dgf--	ANRNFQMLIHDDSL	702
WP_004636532	634	KQLQTYSDTLSPEILKLLER-KHYTGWGRFSKLLINGIRDE	GSNKTILDYLK	DEGsgp	TNRNFQMLIRDNTL	706
WP_002364836	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLI	DDGvs	kHYNRNFQMLINDSQL	714
WP_016631044	593	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLV	DDGvs	kHYNRNFQMLINDSQL	665
EMS75795	382	TQLKKYQSVLGDGFVKLVK-KHYTGWGRLSERLLINGIRDK	KTNKTILDYLI	DDDFpy	NRNRNFQMLINDSQL	454
WP_002373311	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLV	DDGvs	kHYNRNFQMLINDSQL	714
WP_002378009	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLI	DDGvs	kHYNRNFQMLINDSQL	714
WP_002407324	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLV	DDGvs	kHYNRNFQMLINDSQL	714
WP_002413717	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLV	DDGvs	kHYNRNFQMLINDSQL	714
WP_010775580	644	TQLSTFKGQFSEEVKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLI	DDGvs	kHYNRNFQMLINDSQL	716
WP_010818269	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILGYLI	DDGvs	kHYNRNFQMLINDSQL	714
WP_010824395	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLI	DDGvs	kHYNRNFQMLINDSQL	714
WP_016622645	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLV	DDGvs	kHYNRNFQMLINDSQL	714
WP_033624816	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLI	DDGvs	kHYNRNFQMLINDSQL	714
WP_033625576	642	TQLSTFKGQFSAEVLKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLI	DDGvs	kHYNRNFQMLINDSQL	714
WP_033789179	642	TQLSTFKGQFSEEVKLLER-KHYTGWGRLSKLLINGIYDK	ESGKTILDYLI	DDGvs	kHYNRNFQMLINDSQL	714
WP_002310644	642	TRLSHHEATLGKHIKKLTK-KHYTGWGRLSKELIQGIRDK	QSNKTILDYLI	DDDfph	HRNRNFQMLINDDNL	714
WP_002312694	643	TRLSHHEATLGKHIKKLTK-KHYTGWGRLSKELIQGIRDK	QSNKTILDYLI	DDDfph	HRNRNFQMLINDDNL	715
WP_002314015	643	TRLSHHEATLGKHIKKLTK-KHYTGWGRLSKELIQGIRDK	QSNKTILDYLI	DDDfph	HRNRNFQMLINDDNL	715
WP_002320716	643	TRLSHHEATLGKHIKKLTK-KHYTGWGRLSKELIQGIRDK	QSNKTILDYLI	DDDfph	HRNRNFQMLINDDNL	715

WP_002330729	642	TRLSHHEATLGKHI IKKLTK-KHYTGWGRLSKELIQGIRDK	QSNKTILDYLI	QSNKTILDYLI	DDDFpHRRNRFMQLINDDSL	714
WP_002335161	643	TRLSHHEATLGKHI IKKLTK-KHYTGWGRLSKELIQGIRDK	QSNKTILDYLI	QSNKTILDYLI	DDDFpHRRNRFMQLINDDSL	715
WP_002345439	643	TRLSHHEATLGKHI IKKLTK-KHYTGWGRLSKELIQGIRDK	QSNKTILDYLI	QSNKTILDYLI	DDDFpHRRNRFMQLINDDSL	715
WP_034867970	638	HQLSKYQEVFGEKLLKEFAR-KHYTGWGRFSAKLIHGIRDR	KTNKTILDYLI	KTNKTILDYLI	DDDDvpaNRNRFMQLINDEHL	710
WP_047937432	643	TRLSHHEATLGKHI IKKLTK-KHYTGWGRLSKELIQGIRDK	QSNKTILDYLI	QSNKTILDYLI	DDDFpHRRNRFMQLINDDSL	715
WP_010720994	638	HQLSKYQEVFGEKLLKEFAR-KHYTGWGRFSAKLIHGIRDR	KTNKTILDYLI	KTNKTILDYLI	DDDDvpaNRNRFMQLINDEHL	710
WP_010737004	638	HQLSKYQEVFGEKLLKEFAR-KHYTGWGRFSAKLIHGIRDR	KTNKTILDYLI	KTNKTILDYLI	DDDDvpaNRNRFMQLINDEHL	710
WP_034700478	638	HQLSKYQEVFGEKLLKEFAR-KHYTGWGRFSAKLIHGIRDR	KTNKTILDYLI	KTNKTILDYLI	DDDDvpaNRNRFMQLINDEHL	710
WP_007209003	635	NQLEQLPINALSTKTIKALSR-RKYTGWGRLSARLIDGIHDK	NSGKTILDYLI	NSGKTILDYLI	DESdsyIVNRNFMQLINDDHL	707
WP_023519017	632	EQLKPYETVGLPAIKKAK-KHYTGWGRSEKMIQGMREK	QSRKTILDYLI	QSRKTILDYLI	DDDFpcNRNRFMQLINDDHL	704
WP_010770040	635	EQLKKYTYLFDDEEVLLKLER-RHYTGWGRLSAKLLIGIKEK	RTHKTILDYLI	RTHKTILDYLI	DDGGkqPINRNLMLQINDSDL	707
WP_048604708	631	EQLSKFSDRLSEKTIKDLER-RHYTGWGRLSAKLINGIHDK	QSNKTILDYLI	QSNKTILDYLI	DDApkKNIINRNFMLINDNRL	703
WP_010750235	637	TQLKKYQRIILGEEIFKKLVK-KKYTGWGRLSKRLINGIRDQ	KTNKTILDYLI	KTNKTILDYLI	DDDFpyNRNRFMQLINDDHL	709
AI116583	673	ERLKTIAHLFDDKVMKQLKR-RRYTGWGRLSRKLINGIRDK	QSGKTILDFLK	QSGKTILDFLK	-DGF---ANRNFMLIHDDSL	741
WP_029073316	648	RLKKEYDLDEEKIKKILKL--KYSGWSRLSKLLSGIKTK	RTPETVLEVME	RTPETVLEVME	-----TNMNLMLQVINDEKL	717
WP_031589969	648	RLKKEYDLDEEKIKKILKL--KYSGWSRLSKLLSGIKTK	RTPETVLEVME	RTPETVLEVME	-----TNMNLMLQVINDEKL	717
KDA45870	631	RLLENYRDFLGDGVLKLER-RHYTGWGRLSAKLLDGIYDK	KTHKTILDCLM	KTHKTILDCLM	EDYs-----QNFMQLINDDYY	698
WP_039099354	651	AKLNEIDWLTQQRVQAAK--RYRKGWGRLSAKLLTQIVN-	ANGQRIMDLLW	ANGQRIMDLLW	-----TTDNFMRIHVHSE--	712
AKP02966	633	EKLHSSNYSYSDQIKKISN-MRYKKGWGRLSKLLTCTITTE	TNTPKSLQLSN	TNTPKSLQLSN	-DLnt-wTTNNNFISIIISNDKY	706
WP_010991369	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLMGIRDK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
WP_033838504	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLMGIRDK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
EHN60060	641	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLMGIRDK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	709
EFR89594	407	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLMGIRDK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	475
WP_038409211	638	EQLQSFSDVLDGTLKLLER-RHYTGWGRLSAKLLTGIRDK	HSHLTILDYLM	HSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
EFR95520	257	EQLQSFSDVLDGTLKLLER-RHYTGWGRLSAKLLTGIRDK	HSHLTILDYLM	HSHLTILDYLM	DDG---LNRNLMQLINDSNL	325
WP_003723650	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
WP_003727705	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
WP_003730785	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
WP_003733029	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
WP_003739838	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
WP_014601172	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
WP_023548323	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706
WP_031665337	638	EQLQQFSDVLDGVLKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILEYLM	QSHLTILEYLM	DDG---LNRNLMQLINDSNL	706
WP_031669209	638	EQLQQFSDVLDGTVLKKLER-RHYTGWGRLSAKLLVGIKREK	QSHLTILDYLM	QSHLTILDYLM	DDG---LNRNLMQLINDSNL	706

WP_033920898	638	EQLQQFSDVLDGTVLKKLER-RHYTGWGRLSAKLLVGI	QSHLTILDYLM	DDG----	LNRNLMQLINDSNL	706
AKI42028	641	EQLQQFSDVLDGGVLLKKLER-RHYTGWGRLSAKLLVGI	QSHLTILDYLM	DDG----	LNRNLMQLINDSNL	709
AKI50529	641	EQLQQFSDVLDGTVLKKLER-RHYTGWGRLSAKLLVGI	QSHLTILDYLM	DDG----	LNRNLMQLINDSNL	709
EFR83390	86	EQLQQFSDVLDGTVLKKLER-RHYTGWGRLSAKLLVGI	QSHLTILEYLM	DDG----	LNRNLMQLINDSNL	154
WP_046323366	638	ERLQEFNSVLEAVLKKLER-RHYTGWGRLSAKLLVGI	ESHLTILDYLM	DDK----	HNRNLMQLINDSNL	706
AKE81011	650	ERLKTIAHLFDDKVMKQ_LKR-RRYTGWGRLSRKLINGI	QSGKTIIDFLK	-Dgf----	ANRNFMLIHDDSL	718
CUO82355	642	RRLKKYALPDDKIQLKL--KYKDW_SRLSKLLDGI	SV--TVLVDVLE	-----	SRLNMEIINDKEL	709
WP_033162887	645	RRLKKVQLDLDLVDKIQLKL--NYTGW_SRLSEKLLT	KA--TVLTVLE	-----	SNKNLMEIINDEKL	712
AGZ01981	667	ERLKTIAHLFDDKVMKQ_LKR-RRYTGWGRLSRKLINGI	QSGKTIIDFLK	-Dgf----	ANRNFMLIHDDSL	735
AKA60242	634	ERLKTIAHLFDDKVMKQ_LKR-RRYTGWGRLSRKLINGI	QSGKTIIDFLK	-Dgf----	ANRNFMLIHDDSL	702
AKS40380	634	ERLKTIAHLFDDKVMKQ_LKR-RRYTGWGRLSRKLINGI	QSGKTIIDFLK	-Dgf----	ANRNFMLIHDDSL	702
4UN5_B	638	ERLKTIAHLFDDKVMKQ_LKR-RRYTGWGRLSRKLINGI	QSGKTIIDFLK	-Dgf----	ANRNFMLIHDDSL	706

WP_010922251	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_039695303	707	PFKQIIQKSQVVG-DVDD-IEAVVHDLPGSPAIIKKGILQSVKIVDELVKVMG-GNPDNIVIVEMARENQ	TTNRRRSQS	780
WP_045635197	702	SFKEIIQKAQVIG-KTDD-VKQVQELSGSPAIIKKGILQSIKIVDELVKVMG-HAPESIVIVEMARENQ	TTARGKNS	775
5AXW A	427	VPKKVDLSQQKEI---PT---TLVDDFILSPVVKRSFIQSIKVINAIKKYG--LPNDIIIELAREKN	-----S	487
WP_009880683	387	TFKEDLQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	461
WP_010922251	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_011054416	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_011284745	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_011285506	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_011527619	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_012560673	703	TFKEDLQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_014407541	703	TFKEDIQKAQVSG-QGHS-LHEQIANLAGSPAIIKKGILQTVKIVDELVKVMG-HKPENIVIVEMARENQ	TTQKGQKNS	776
WP_020905136	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_023080005	703	TFKEAIQKAQVSG-QGHS-LHEQIANLAGSPAIIKKGILQTVKIVDELVKVMG-HKPENIVIVEMARENQ	TTQKGQKNS	776
WP_023610282	703	TFKEAIQKAQVSG-QGHS-LHEQIANLAGSPAIIKKGILQTVKIVDELVKVMG-HKPENIVIVEMARENQ	TTQKGQKNS	776
WP_030125963	703	TFKEDLQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_030126706	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_031488318	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_032460140	703	TFKEDLQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_032461047	703	TFKEDLQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_032462016	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_032462936	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_032464890	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_033888930	528	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	602
WP_038431314	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_038432938	703	TFKEAIQKAQVSG-QGHS-LHEQIANLAGSPAIIKKGILQTVKIVDELVKVMG-HKPENIVIVEMARENQ	TTQKGQKNS	776
WP_038434062	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKIVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
BAQ51233	614	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	688
KGE60162		-----	-----	
KGE60856		-----	-----	
WP_002989955	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKNS	777
WP_003030002	704	SFKEIARAQIIG-DVDD-IANVHDLPGSPAIIKKGILQSVKIVDELVKVMG-HNPNANIIEMARENQ	MTDKRRNS	777
WP_003065552	707	PFKQIIQKSQVVG-DVDD-IEAVVHDLPGSPAIIKKGILQSVKIVDELVKVMG-DNPDNIVIVEMARENQ	TTNRRRSQS	780
WP_001040076	702	SFKPIIDKARTGS-HSDN-LKEVIGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRS	775

WP_001040078	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040080	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040081	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040083	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040085	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040087	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040088	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040089	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040090	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040091	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040092	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040094	702	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRS	775
WP_001040095	702	SFKPIIDKARTGS-HLDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRS	775
WP_001040096	702	SFKPIIDKARTGS-HLDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRS	775
WP_001040097	702	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRL	775
WP_001040098	702	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRS	775
WP_001040099	702	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRS	775
WP_001040100	702	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRS	775
WP_001040104	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040105	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_001040106	705	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNT	778
WP_001040107	705	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNT	778
WP_001040108	705	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNT	778
WP_001040109	705	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNT	778
WP_001040110	705	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNT	778
WP_015058523	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_017643650	702	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRL	775
WP_017647151	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_017648376	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_017649527	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_017771611	705	SFKPIIDKARTGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNT	778
WP_017771984	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
CFQ25032	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
CFV16040	705	SFKSII SKAQAGS-HSDN-LKEVVGELAGSPAIIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778

KLJ37842	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
KLJ72361	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
KL20707	719	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	792
KL242645	705	SFKPIIDKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNKRRNT	778
WP_047207273	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_047209694	702	SFKPIIDKARTGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTAKGLSRS	775
WP_050198062	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_050201642	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_050204027	705	SFKPIIDKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNT	778
WP_050881965	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_050886065	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
AHN30376	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
EAO78426	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
CCW42055	705	SFKSII SKAQAGS-HSDN-LKEVWVGEIAGSPAIKKGILQSLKIVDELVKVMG-YEPEQIVVEMARENQ	TTNQRRNS	778
WP_003041502	704	SFKEEIAKAQIIG-DVDD-IANVHDLPGSPAIKKGILQSLKIVDELVKVMG-HNPANIIEMARENQ	TTDRGRNS	777
WP_037593752	705	SFKEEIAARAQIIG-DVDD-IANVHDLPGSPAIKKGILQSLKIVDELVKVMG-HNPANIIEMARENQ	TTDKGRNS	778
WP_049516684	705	SFKEEIAARAQIIG-DVDD-IANVHDLPGSPAIKKGILQSLKIVDELVKVMG-HNPANIIEMARENQ	TTDKGRNS	778
GAD46167	704	SFKEEIAARAQIIG-DVDD-IANVHDLPGSPAIKKGILQSLKIVDELVKVMG-HNPANIIEMARENQ	TTDKGRNS	777
WP_018363470	705	SFKQIIQEAQVVG-DVDD-IETVHDLPGSPAIKKGILQSLKIVDELVKVMG-DNPDNIVEMARENQ	TTNRRSQS	778
WP_003043819	713	TFKEEIEKAQVVG-DVDD-IETVHDLPGSPAIKKGILQSLKIVDELVKVMG-HKPNIVIVEMARENQ	TTTKGLQOS	786
WP_006269658	704	SFKEEIAARAQIID-DVDD-IANVHDLPGSPAIKKGILQSLKIVDELVKVMG-HNPANIIEMARENQ	TTDKGRNS	777
WP_048800889	704	PFKQIIKDAQAID-DVDD-IELIVHDLPGSPAIKKGILQSLKIVDELVKVMG-YNPDNIVIVEMARENQ	TTTKGRNS	777
WP_012767106	703	TFKEAIQKAQVVG-QGHS-LHEQIANLAGSPAIKKGILQSLKIVDELVKVMG-HKPNIVIVEMARENQ	TTQKGQKNS	776
WP_014612333	703	TFKEAIQKAQVVG-QGHS-LHEQIANLAGSPAIKKGILQSLKIVDELVKVMG-HKPNIVIVEMARENQ	TTQKGQKNS	776
WP_015017095	703	TFKEAIQKAQVVG-QGHS-LHEQIANLAGSPAIKKGILQSLKIVDELVKVMG-HKPNIVIVEMARENQ	TTQKGQKNS	776
WP_015057649	703	TFKEAIQKAQVVG-QGHS-LHEQIANLAGSPAIKKGILQSLKIVDELVKVMG-HKPNIVIVEMARENQ	TTQKGQKNS	776
WP_048327215	703	TFKEAIQKAQVVG-QGHS-LHEQIANLAGSPAIKKGILQSLKIVDELVKVMG-HKPNIVIVEMARENQ	TTQKGQKNS	776
WP_049519324	703	TFKEAIQKAQVVG-QGHS-LHEQIANLAGSPAIKKGILQSLKIVDELVKVMG-HKPNIVIVEMARENQ	TTQKGQKNS	776
WP_012515931	703	SFIDEIAKAQVIG-KTEY-SKDLVGNLAGSPAIKKGISQTIKIVDELVKIMG-YLPQQIVIVEMARENQ	TTAQGIKNA	776
WP_021320964	703	SFIDEIAKAQVIG-KTEY-SKDLVGNLAGSPAIKKGISQTIKIVDELVKIMG-YLPQQIVIVEMARENQ	TTAQGIKNA	776
WP_037581760	703	SFIDEIAKAQVIG-KTEY-SKDLVGNLAGSPAIKKGISQTIKIVDELVKIMG-YLPQQIVIVEMARENQ	TTAQGIKNA	776
WP_004232481	704	SFKTTIQEAQVVG-DVDD-IEAVHDLPGSPAIKKGILQSLKIVDELVKVMG-HNPQNIVIVEMARENQ	IITGYGRNS	777
WP_009854540	705	PFKQIIQKSQVVG-DVDD-IEAVHDLPGSPAIKKGILQSLKIVDELVKVMG-DNPDNIVIVEMARENQ	TTNRRSQS	778
WP_012962174	705	PFKQIIKDAQIIG-DIDD-VTSVRELPGSPAIKKGILQSLKIVDELVKVMG-HNPDNIVIVEMARENQ	TTNRRGRNS	778

WP_039695303	707	PFKQIIQKSQVVG-DVDD-IEAVVHDLPGSPAIAKKGILQSVKIVDELVKVMG-GNPDNIVIVEMARENQ	TTNRRSQS	780
WP_014334983	704	SFKTIIQEAQVVG-DVDD-IEAVVHDLPGSPAIAKKGILQSVKIVDELVKVMG-DNPDNIVIVEMARENQ	TTGYGRNKS	777
WP_003099269	703	DFAKIIKNEQEKTIKNES-LEETIANLAGSPAIAKKGILQSIKIVDEIVKIMG-QNPDNIVIVEMARENQ	STMQGIKNS	777
AHY15608	703	DFAKIIKNEQEKTIKNES-LEETIANLAGSPAIAKKGILQSIKIVDEIVKIMG-QNPDNIVIVEMARENQ	STMQGIKNS	777
AHY17476	703	DFAKIIKNEQEKTIKNES-LEETIANLAGSPAIAKKGILQSIKIVDEIVKIMG-QNPDNIVIVEMARENQ	STMQGIKNS	777
ESR09100		-----	-----	
AGM98575	703	DFAKIIKNEQEKTIKNES-LEETIANLAGSPAIAKKGILQSIKIVDEIVKIMG-QNPDNIVIVEMARENQ	STMQGIKNS	777
ALF27331	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_018372492	717	DFKEIIRKAQTIE-NIDT-NQALVSSLPSPAIAKKGILQSLNIVDEIIAIMG-YAPTIVIVEMARENQ	TTQKGRDNS	790
WP_045618028	703	SFKEIIQKAQVVG-KTND-VKQVWQELPGSPAIAKKGILQSIKIVDELVKVMG-HAPESIVIVEMARENQ	TTARGKKNs	776
WP_045635197	702	SFKEIIQKAQVIG-KTDD-VKQVWQELSGSPAIAKKGILQSIKIVDELVKVMG-HAPESIVIVEMARENQ	TTARGKKNs	775
WP_002263549	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002263887	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002264920	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002269043	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002269448	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002271977	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002272766	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002273241	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTKQGRNRS	777
WP_002275430	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002276448	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTKQGRNRS	777
WP_002277050	705	TFKDDIVKAQYVD-NSDD-LHQVWQSLAGSPAIAKKGILQSLKIVDELVKIMG-KEPEQIVIVEMARENQ	TTAKGRNRS	778
WP_002277364	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002279025	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002279859	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002280230	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTKQGRNRS	777
WP_002281696	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002282247	705	TFKDDIVKAQYVD-NSDD-LHQVWQSLAGSPAIAKKGILQSLKIVDELVKIMG-KEPEQIVIVEMARENQ	TTAKGRNRS	778
WP_002282906	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002283846	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002287255	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002288990	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002289641	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777
WP_002290427	704	SFKEEIAKAQVIG-ETDN-LNQVSDIAGSPAIAKKGILQSLKIVDELVKIMG-HQPENIVIVEMARENQ	FTNQGRNRS	777

WP_002295753	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTKQGRNS	777
WP_002296423	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_002304487	714	SFKEEIAKAQVIG-EMDG-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HNPANIVVEMARENQ	TTAKGRSS	787
WP_002305844	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTKQGRNS	777
WP_002307203	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_002310390	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSLAIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_002352408	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_012997688	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_014677909	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_019312892	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_019313659	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_019314093	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_019315370	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_019803776	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_019805234	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSLAIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_024783594	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_024784288	705	TFKDDIVKAQVVD-NSDD-LHQQVQSLAGSPAIIKGGILQSLKIVDELVKIMG-KEPEQIVVEMARENQ	TTAKGRNS	778
WP_024784666	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_024784894	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_024786433	705	TFKDDIVKAQVVD-NSDD-LHQQVQSLAGSPAIIKGGILQSLKIVDELVKIMG-KEPEQIVVEMARENQ	TTAKGRNS	778
WP_049473442	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
WP_049474547	704	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	777
EMC03581	697	SFKEEIAKAQVIG-ETDN-LNQVWSDIAGSPAIIKGGILQSLKIVDELVKIMG-HQPENIVVEMARENQ	FTNQGRNS	770
WP_000428612	705	SFKEEIAKAQVIG-ETDN-LNQVWQELPGSPAIIKGGILQSLKIVDELVKIMG-HEPESIVVEMARENQ	TTARGKNS	778
WP_000428613	703	SFKEEIAKAQVIG-ETDN-LNQVWQELPGSPAIIKGGILQSLKIVDELVKIMG-HTPESIVVEMARENQ	TTARGKNS	776
WP_049523028	702	SFKEEIAKAQVIG-ETDN-LNQVWQELPGSPAIIKGGILQSLKIVDELVKIMG-HAPESVVIEMARENQ	TTNKGSKS	775
WP_003107102	672	DFASIIKEAQEKTIKSEK-LEETIANLAGSPAIIKGGILQSLKIVDELVKIMG-YEPSNIVVEMARENQ	STQRGINNS	746
WP_054279288	705	SFKEEIAKAQVIG-ETDN-LNQVWQELPGSPAIIKGGILQSLKIVDELVKIMG-KAPQHIIVVEMARDVQ	KTDIGVKQS	778
WP_049531101	703	SFKEEIAKAQVIG-ETDN-LNQVWQELPGSPAIIKGGILQSLKIVDELVKIMG-HDPESIVVEMARENQ	TTARGKNS	776
WP_049538452	703	SFKEEIAKAQVIG-ETDN-LNQVWQELPGSPAIIKGGILQSLKIVDELVKIMG-HEPESIVVEMARENQ	TTTRGKNS	776
WP_049549711	703	SFKEEIAKAQVIG-ETDN-LNQVWQELPGSPAIIKGGILQSLKIVDELVKIMG-HAPESIVVEMARENQ	TTARGKNS	776
WP_007896501	706	DFEKLIDDAQKKAiKRES-LTEAVANLAGSPAIIKGGILQSLKIVDELVKIMG-HNPDNIVVEMARENQ	TTAQGLKNA	780
EFR44625	658	DFEKLIDDAQKKAiKRES-LTEAVANLAGSPAIIKGGILQSLKIVDELVKIMG-HNPDNIVVEMARENQ	TTAQGLKNA	732
WP_002897477	702	SFKEEIAKAQVIG-ETDN-LNQVWQELPGSPAIIKGGILQSLKIVDELVKIMG-YALESIVVEMARENQ	TTARGKNS	775

WP_002906454	702	SFKEIIQKAQVVG-KTDD-VKQVQVEIPGSPAIKKGILQSIKIVDELVKVMG-HNPESIVIVEMARENQ	TTAKGKKN	775
WP_009729476	703	SFKEIIQKAQVVG-KTND-VKQVQVELPGSPAIKKGILQSIKIVDELVKVMG-HAPESIVIVEMARENQ	TTARGKKN	776
CQR24647	706	SFKDEIANSQVIG-DGDD-LHQVQVELAGSPAIKKGILQSLKIVDELVKVMG-YNPEQIVVEMARENQ	TTARGRNN	779
WP_000066813	705	SFKEIIQKAQVVG-KTND-VKQVQVELPGSPAIKKGILQSIKIVDELVKVMG-HAPESIVIVEMARENQ	TTARGKKN	778
WP_009754323	703	SFKEIIQKAQVVG-KTDD-LTQVRELSPAIKKGILQSIKIVDELVKVMG-YAPESIVIVEMARENQ	TTAKGKKN	776
WP_044674937	702	SFVDEIRLAQSG-EAED-YRAEVQNLGSPAIKKGILQSLKIVDELIVMGM-YDPEHIVVEMARENQ	FTNQRRNS	775
WP_044676715	704	SFVDEIRLAQSG-EAED-YRAEVQNLGSPAIKKGILQSLKIVDELIVMGM-YDPEHIVVEMARENQ	FTNQRRNS	777
WP_044680361	704	SFVDEIRLAQSG-EAED-YRAEVQNLGSPAIKKGILQSLKIVDELIVMGM-YDPEHIVVEMARENQ	FTNQRRNS	777
WP_044681799	702	SFVDEIRLAQSG-EAED-YRAEVQNLGSPAIKKGILQSLKIVDELIVMGM-YDPEHIVVEMARENQ	FTNQRRNS	775
WP_049533112	704	SFKEEIIKAQVIG-ETDD-LNQVSDIAGSPAIKKGILQSLKIVDELVKVMG-YNPANIVIVEMARENQ	TTDKGRNS	777
WP_029090905	683	LFKKIEEQFENETALLN--KQRIDELAASPANKKIWIQAIKIVKELEKVLQ-QPAENIFIEFARSD	ES---	752
WP_006506696	706	GYAQMIEEATSCPEdGKF-TYEEVERLAGSPALKRGIWQSLQIVEEITKVMK-CRPKIYIIEFERSEE	----	776
AIT42264	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKN	777
WP_034440723	711	SFKEKIRKAQDIN-QVND-IKEIVKDLPGSPAIKKGIIYQSIRIVDEIIRKMK-DRPKNIVIVEMARENQ	TTQEGKKN	784
AKQ21048	703	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIKKGILQTVKVVDELVKVMGrHKPENIVIVEMARENQ	TTQKGQKN	777
WP_004636532	707	SFKKKIEDAQTIIE-DTTH-IYDTVAELPGSPAIKKGIRQALKIVVEEIIIDIIIG-YEPENIVVEMARESQ	TTKKGKDL	780
WP_002364836	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_016631044	666	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	739
EMS75795	455	SFKEELANELALA-GNQS-LLEVVEALLGSPAIKKGIIWQTLKIVVEELIEIIG-YNPKNIVVEMARENQ	RT---	524
WP_002373311	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_002378009	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_002407324	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_002413717	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_010775580	717	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	790
WP_010818269	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_010824395	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_016622645	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_033624816	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_033625576	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_033789179	715	SFKNAIQKAQSE-HEET-LSETVNELAGSPAIKKGIIYQSLKIVDELVAIMG-YAPKRIVVEMARENQ	TTSTGRRS	788
WP_002310644	715	SFKKEIKKAQMIT-DTEN-LEEIVKELTGPSAIKKGILQSLKIVDEIVGIMG-YEPANIVVEMARENQ	TTGRGLKSS	788
WP_002312694	716	SFKKEIKKAQMIT-DTEN-LEEIVKELTGPSAIKKGILQSLKIVDEIVGIMG-YEPANIVVEMARENQ	TTGRGLKSS	789
WP_002314015	716	SFKKEIKKAQMIT-DTEN-LEEIVKELTGPSAIKKGILQSLKIVDEIVGIMG-YEPANIVVEMARENQ	TTGRGLKSS	789
WP_002320716	716	SFKKEIKKAQMIT-DTEN-LEEIVKELTGPSAIKKGILQSLKIVDEIVGIMG-YEPANIVVEMARENQ	TTGRGLKSS	789

WP_002330729	715	SFKKEIKKAQMIT-DTEN-LEEVKELTGSPAIKKGILQSLKIVDEIVGIMG-YEPANIVVEMARENQ	TTGRGLKSS	788
WP_002335161	716	SFKKEIKKAQMIT-DTEN-LEEVKELTGSPAIKKGILQSLKIVDEIVGIMG-YEPANIVVEMARENQ	TTGRGLKSS	789
WP_002345439	716	SFKKEIKKAQMIT-DTEN-LEEVKELTGSPAIKKGILQSLKIVDEIVGIMG-YEPANIVVEMARENQ	TTGRGLKSS	789
WP_034867970	711	SFKEEIAKATVFS-KHKS-LVDVIQDLPSPAIKKGILQSLKIVDEIVGIMG-YKPKNIVVEMARENQ	KT----HRT	780
WP_047937432	716	SFKKEIKKAQMIT-DTEN-LEEVKELTGSPAIKKGILQSLKIVDEIVGIMG-YEPANIVVEMARENQ	TTGRGLKSS	789
WP_010720994	711	SFKEEIAKATVFS-KHKS-LVDVIQDLPSPAIKKGILQSLKIVDEIVGIMG-YKPKNIVVEMARENQ	KT----HRT	780
WP_010737004	711	SFKEEIAKATVFS-KHKS-LVDVIQDLPSPAIKKGILQSLKIVDEIVGIMG-YKPKNIVVEMARENQ	KT----HRT	780
WP_034700478	711	SFKEEIAKATVFS-KHKS-LVDVIQDLPSPAIKKGILQSLKIVDEIVGIMG-YKPKNIVVEMARENQ	KT----HRT	780
WP_007209003	708	SFKKIEDSQPK-EQQS-AEEIVSELSPAIKKGILQSLKIVDELVAIMG-YKPKNIVVEMARENQ	TTGRGKQNS	781
WP_023519017	705	SFKETIANELIMS-DSNV-LLDQVKAI PGSPAVKGIWQSIKIVEEIIIGIIG-KAPKNIVVEMARENQ	RTSR----S	774
WP_010770040	708	SFKSEIAEAQSDM-NTED-LHEVVQNLAGSPAIKKGILQSLKIVDELVDIMG-SLPKNIVVEMARENQ	TTSRGRTNS	781
WP_048604708	704	TFKEEIEKEQLKA-NSEESLIEIVQNLAGSPAIKKGIFQSLKIVDELVEIMG-YAPTNI VVEMARENQ	TTANGRRNS	778
WP_010750235	710	SFKEEIAKELTLS-DKQS-LLEVVEAIPGSPAIKKGILQSLKIVDEIVGIMG-YKPKNIVVEMARENQ	TTTTGGKNS	783
AI116583	742	TFKEDIQKAQVSG-QGDS-LHEHIANLAGSPAIKKGILQSLKIVDELVAIMG-YKPKNIVVEMARENQ	TTQKQKNS	816
WP_029073316	718	GFKKTIDDANSTSVSGKF-SYAEVQELAGSPAIKKGILQSLKIVDEIVGIMG-HEPAHVYIEFARNED	-----KERK	788
WP_031589969	718	GFKKTIDDANSTSVSGKF-SYAEVQELAGSPAIKKGILQSLKIVDEIVGIMG-HEPAHVYIEFARNED	-----KERK	788
KDA45870	699	SFKETIKNAQVIE-KEET-LAKTVQELPGSPAIKKGILQSLKIVDEIVGIMG-YKPKSIVVEMARETQ	---THGTRKR	771
WP_039099354	713	DFDKLITEANQMM-LAENdVQVINDLTPSPQNKALRQILLVNDIQKAMKQAPERILIEFAREDE	VNPRLSVQR	788
AKP02966	707	DFKYYIENHNLKnedQn-ISNLVNDIHVSPALKRGITQSIKIVQEI VKFVG-HAPKYIFIEVTRETK	TTSRGKRIQ	785
WP_010991369	707	SFKSIEKEQVTT-ADKD-IQSI VADLAGSPAIKKGILQSLKIVDELVSVMG-YPPQTI VVEMARENQ	TTGKGKNS	780
WP_033838504	707	SFKSIEKEQVTT-ADKD-IQSI VADLAGSPAIKKGILQSLKIVDELVSVMG-YPPQTI VVEMARENQ	TTGKGKNS	780
EHN60060	710	SFKSIEKEQVTT-ADKD-IQSI VADLAGSPAIKKGILQSLKIVDELVSVMG-YPPQTI VVEMARENQ	TTGKGKNS	783
EFR89594	476	SFKSIEKEQVTT-ADKD-IQSI VADLAGSPAIKKGILQSLKIVDELVSVMG-YPPQTI VVEMARENQ	TTGKGKNS	549
WP_038409211	707	SFKSIEKEQVST-ADKG-IQSI VAEELAGSPAIKKGILQSLKIVDELVGIMG-YPPQTI VVEMARENQ	TTGKGKNS	780
EFR95520	326	SFKSIEKEQVST-ADKG-IQSI VAEELAGSPAIKKGILQSLKIVDELVGIMG-YPPQTI VVEMARENQ	TTGKGKNS	399
WP_003723650	707	SFKSIEKEQVST-TDKD-LQSI VAEELAGSPAIKKGILQSLKIVDELVSIMG-YPPQTI VVEMARENQ	TTGKGKNS	780
WP_003727705	707	SFKSIEKEQVST-TDKD-LQSI VADLAGSPAIKKGILQSLKIVDELVSIMG-YPPQTI VVEMARENQ	TTGKGKNS	780
WP_003730785	707	SFKSIEKEQVST-TDKD-LQSI VADLAGSPAIKKGILQSLKIVDELVSIMG-YPPQTI VVEMARENQ	TTGKGKNS	780
WP_003733029	707	SFKSIEKEQVST-TDKD-LQSI VAEELAGSPAIKKGILQSLKIVDELVSIMG-YPPQTI VVEMARENQ	TTNKGGKNS	780
WP_003739838	707	SFKSIEKEQVST-TDKD-LQSI VADLAGSPAIKKGILQSLKIVDELVSIMG-YPPQTI VVEMARENQ	TTVKGGKNS	780
WP_014601172	707	SFKSIEKEQVST-TDKD-LQSI VADLAGSPAIKKGILQSLKIVDELVSIMG-YPPQTI VVEMARENQ	TTGKGKNS	780
WP_023548323	707	SFKSIEKEQVST-ADKD-LQSI VADLAGSPAIKKGILQSLKIVDELVSVMG-YPPQTI VVEMARENQ	TTNKGGKNS	780
WP_031665337	707	SFKSIEKEQVST-TDKD-LQSI VAEELAGSPAIKKGILQSLKIVDELVSIMG-YPPQTI VVEMARENQ	TTGKGKNS	780
WP_031669209	707	SFKSIEKEQVST-ADKD-LQSI VADLAGSPAIKKGILQSLKIVDELVSVMG-YPPQTI VVEMARENQ	TTNKGGKNS	780

WP_033920898	707	SFKSII	IEKEQVST-ADKD-LQSI	VADLAGSPA	IKKGI	LQSLK	VEELV	SVMG-YP	PQTIV	VEMARE	NQ	780	TTNKGKNN
AKI42028	710	SFKSII	IEKEQVST-TDKD-LQSI	VADLAGSPA	IKKGI	LQSLK	IVDEL	VSIMG-YP	PQTIV	VEMARE	NQ	783	TTGKGKNN
AKI50529	710	SFKSII	IEKEQVST-ADKD-LQSI	VADLAGSPA	IKKGI	LQSLK	VEELV	SVMG-YP	PQTIV	VEMARE	NQ	783	TTNKGKNN
EFR83390	155	SFKSII	IEKEQVST-TDKD-LQSI	VADLAGSPA	IKKGI	LQSLK	IVDEL	VSIMG-YP	PQTIV	VEMARE	NQ	228	TTVKGKNN
WP_046323366	707	SFKSII	IEKEQVST-ADKD-IQSI	VADLAGSPA	IKKGI	LQSLK	IVDEL	VGIMG-YP	PQTIV	VEMARE	NQ	780	TTGKGKNN
AKE81011	719	TFKEDI	QKAQVSG-QGDS-LHEH	IANLAGSPA	IKKGI	LQTVK	VDEL	VKMGr	HKPENI	VIEMARE	NQ	793	TTQKGQKN
CUO82355	710	GYAQMI	EEASSCPkDGKF-TYEE	VAKLAGSPA	IKRGI	WQSLQ	IVVEE	ITKVMK-CR	PKYI	YIEFER	SEE	780	-----KERT
WP_033162887	713	GYKQII	EESNMQDiEGPF-KYDE	VKKLAGSPA	IKRGI	WQALLV	REIT	KFMK-HEP	SHI	YIEFA	REEQ	783	-----KVRK
AGZ01981	736	TFKEDI	QKAQVSG-QGDS-LHEH	IANLAGSPA	IKKGI	LQTVK	VDEL	VKMGr	HKPENI	VIEMARE	NQ	810	TTQKGQKN
AKA60242	703	TFKEDI	QKAQVSG-QGDS-LHEH	IANLAGSPA	IKKGI	LQTVK	VDEL	VKMGr	HKPENI	VIEMARE	NQ	777	TTQKGQKN
AKS40380	703	TFKEDI	QKAQVSG-QGDS-LHEH	IANLAGSPA	IKKGI	LQTVK	VDEL	VKMGr	HKPENI	VIEMARE	NQ	777	TTQKGQKN
4UN5_B	707	TFKEDI	QKAQVSG-QGDS-LHEH	IANLAGSPA	IKKGI	LQTVK	VDEL	VKMGr	HKPENI	VIEMARE	NQ	781	TTQKGQKN

WP	ERMKRIEEGIK	ELGS	QILKEHP--VE--NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_010922251	ERMKRIEEGIK	PSYI	E---DK--VE---NSHLQNDQLFLYYIQNGKDMYTGDEL--D--IDHLSYDIDHI	841
WP_039695303	QRLKQLQNSLK	PSYI	NILKENP--TD---NNQLQNDRLFLYYIQNGKDMYTGEAL--D--INQLSSYDIDHI	851
WP_045635197	QRYKRIEDSLK	ILAS	EIIRTTGK--E---NAKYLIEKIKLHDMQEGKCLYSLEAIpleLNNPFYEVVDHI	843
5AXW_A	KDAQKMINEMQK	QTNE	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	561
WP_009880683	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	525
WP_010922251	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_011054416	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_011284745	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_011285506	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_011527619	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_012560673	ERMKRIEEGIK	ELGS	QILKEHP--VE---TTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_014407541	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	840
WP_020905136	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_023080005	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	840
WP_023610282	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	840
WP_030125963	ERMKRIEEGIK	ELGS	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_030126706	ERMKRIEEGIK	ELGS	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_031488318	ERMKRIEEGIK	ELGS	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_032460140	ERMKRIEEGIK	ELGS	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_032461047	ERMKRIEEGIK	ELGS	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_032462016	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_032462936	ERMKRIEEGIK	ELGS	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_032464890	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_033888930	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	666
WP_038431314	ERMKRIEEGIK	ELGS	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_038432938	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	840
WP_038434062	ERMKRIEEGIK	ELGS	DILKEYP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
BAQ51233	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	752
KGE60162	-----	----	-----QEL--D--INRLSGYDVDHI	16
KGE60856	-----	----	-----	16
WP_002989955	ERMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
WP_003030002	QRLKLLQDSLK	PVNI	K---N--VE---NQQLQNDRLFLYYIQNGKDMYTGETL--D--INNLSQYDIDHI	840
WP_003065552	QRLKQLQNSLK	PSYI	E---DK--VE---NSHLQNDQLFLYYIQNGKDMYTGDEL--D--IDHLSYDIDHI	851
WP_001040076	RQLTTLRESLA	NLKS	EKKPKYV--KdqveNHHLSDDRFLYYLQNGKDMYTDEL--D--IDNLSQYDIDHI	846

WP_001040078	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040080	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040081	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040083	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040085	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040087	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040088	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040089	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040090	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040091	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040092	779	RQRYKLLDDGVK	NLAS	DILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040094	776	RQRLTTLRESLA	NLKS	EKKPKYV--KDQVHHLSDDRLFLYYLQNGKDMYTDEAL--D--IDNLSQYDIDHI	846
WP_001040095	776	RQRLTTLRESLA	NLKS	EKKPKYV--KDQVHHLSDDRLFLYYLQNGKDMYTDEAL--D--IDNLSQYDIDHI	846
WP_001040096	776	RQRLTTLRESLA	NLKS	EKKPKYV--KDQVHHLSDDRLFLYYLQNGKDMYTDEAL--D--IDNLSQYDIDHI	846
WP_001040097	776	RQRLTTLRESLA	NLKS	EKKPKYV--KDQVHHLSDDRLFLYYLQNGKDMYTDEAL--D--IDNLSQYDIDHI	846
WP_001040098	776	RQRLTTLRESLA	NLKS	EKKPKYV--KDQVHHLSDDRLFLYYLQNGKDMYTDEAL--D--IDNLSQYDIDHI	846
WP_001040099	776	RQRLTTLRESLA	NLKS	EKKPKYV--KDQVHHLSDDRLFLYYLQNGKDMYTDEAL--D--IDNLSQYDIDHI	846
WP_001040100	776	RQRLTTLRESLA	NLKS	EKKPKYV--KDQVHHLSDDRLFLYYLQNGKDMYTDEAL--D--IDNLSQYDIDHI	846
WP_001040104	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040105	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040106	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040107	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040108	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040109	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_001040110	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_015058523	779	RQRYKLLDDGVK	NLAS	DILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_017643650	776	RQRLTTLRESLA	NLKS	EKKPKYV--KDQVHHLSDDRLFLYYLQNGKDMYTDEAL--D--IDNLSQYDIDHI	846
WP_017647151	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_017648376	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_017649527	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_017771611	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_017771984	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
CFQ25032	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
CFV16040	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD--NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846

KLJ37842	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
KLJ72361	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
KL20707	793	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	860
KL42645	779	RQRYKLLDEGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_047207273	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_047209694	776	RQRLTTLRESLA	NLKS	EKKPKYV--KdqvEHHLSDDRFLYYLQNGKDMYTDEL--D--IDNLSQYDIDHI	846
WP_050198062	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_050201642	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_050204027	779	RQRYKLLDEGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_050881965	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
WP_050886065	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
AHN30376	779	RQRYKLLDDGVK	NLAS	DILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
EA078426	779	RQRYKLLDDGVK	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTGEAL--D--IDNLSQYDIDHI	846
CCW42055	779	RQRYKLLDDGVR	NLAS	NILKEYP--TD---NQALQNERLFLYYLQNGRDMYTEKAL--D--IDNLSQYDIDHI	846
WP_003041502	778	QQRLLKLLQDSLK	PVNI	K-----N-VE---NQQLQNDRLFLYYIQNGKDMYTGETL--D--INNLSQYDIDHI	840
WP_037593752	779	QQRLLKLLQDSLK	PVNI	K-----N-VE---NQQLQNDRLFLYYIQNGKDMYTGETL--D--INNLSQYDIDHI	841
WP_049516684	779	QQRLLKLLQDSLK	PVNI	K-----N-VE---NQQLQNDRLFLYYIQNGKDMYTGETL--D--INNLSQYDIDHI	841
GAD46167	778	QQRLLKLLQDSLK	PVNI	K-----N-VE---NQQLQNDRLFLYYIQNGKDMYTGETL--D--INNLSQYDIDHI	841
WP_018363470	779	QQRLLKLLQDSLK	PSYI	E-----DK-VE---NSHLQNDQLFLYYIQNGKDMYTGETL--D--IDHLSQYDIDHI	849
WP_003043819	787	RERKKRIEEGK	ELES	QILKENP--VE---NTQLQNEKLYLQNGRDMYDQEL--D--INRLSDYDIDHI	850
WP_006269658	778	QQRLLKLLQDSLK	PVNI	K-----N-VE---NQQLQNDRLFLYYIQNGKDMYTGETL--D--INNLSQYDIDHI	840
WP_048800889	778	QQRLLKLLQDSLK	PVSI	K-----N-VE---NQQLQNDRLFLYYIQNGKDMYTGETL--D--INNLSQYDIDHI	840
WP_012767106	777	RERMKRIEEGK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYDQEL--D--INRLSDYDIDHI	840
WP_014612333	777	RERMKRIEEGK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYDQEL--D--INRLSDYDIDHI	840
WP_015017095	777	RERMKRIEEGK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYDQEL--D--INRLSDYDIDHI	840
WP_015057649	777	RERMKRIEEGK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYDQEL--D--INRLSDYDIDHI	840
WP_048327215	777	RERMKRIEEGK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYDQEL--D--INRLSDYDIDHI	840
WP_049519324	777	RERMKRIEEGK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYDQEL--D--INRLSDYDIDHI	840
WP_012515931	777	RQMRKLEETAK	KLGS	NILKEHP--VD---NSQLQNDKRYLYLQNGKDMYTGDDL--D--IDYLSYDIDHI	840
WP_021320964	777	RQMRKLEETAK	KLGS	NILKEHP--VD---NSQLQNDKRYLYLQNGKDMYTGDDL--D--IDYLSYDIDHI	840
WP_037581760	777	RQMRKLEETAK	KLGS	NILKEHP--VD---NSQLQNDKRYLYLQNGKDMYTGDDL--D--IDYLSYDIDHI	840
WP_004232481	778	NQRLKRLQDSLK	PSYV	D-----SK-VE---NSHLQNDRLFLYYIQNGKDMYTGEEL--D--IDHLSQYDIDHI	848
WP_009854540	779	QQRLLKLLQDSLK	PSYI	E-----DK-VE---NSHLQNDQLFLYYIQNGKDMYTGEAL--D--IDHLSQYDIDHI	849
WP_012962174	779	QQRLLKLLQDSLK	PSYI	E-----GK-VE---NNHLQDDRLFLYYIQNGKDMYTGDEL--D--IDHLSQYDIDHI	849

WP_039695303	781	QQRLLKQLNSLK	PSYI	E----	DK--VE---	NSHLQNDQLFLYYIQNGKDMYTGDEL--D--IDHLSQYDIDHI	851
WP_014334983	778	NQRLKRLQDSLK	PSYV	D----	SK--VE---	NSHLQNDRLFLYYIQNGKDMYTGDEL--D--IDRLSDYDIDHI	848
WP_003099269	778	RQRLKLEEVHK	NTGS	KILKEYN--VS---	NTQLQSDRLYLLQDGKDMYTGKEL--D--YDNLQYDIDHI	841	
AHY15608	778	RQRLKLEEVHK	NTGS	KILKEYN--VS---	NTQLQSDRLYLLQDGKDMYTGKEL--D--YDNLQYDIDHI	841	
AHY17476	778	RQRLKLEEVHK	NTGS	KILKEYN--VS---	NTQLQSDRLYLLQDGKDMYTGKEL--D--YDNLQYDIDHI	841	
ESR09100							
AGM98575	778	RQRLKLEEVHK	NTGS	KILKEYN--VS---	NTQLQSDRLYLLQDGKDMYTGKEL--D--YDNLQYDIDHI	841	
ALF27331	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGDEL--D--IDYLSQYDIDHI	841	
WP_018372492	791	AQRLKKIEDGK	-LGS	DLKQNP--IQd--	NKDLQKEKFLYYMQNGIDLYTGQPlncD--PDSLAFYDIDHI	857	
WP_045618028	777	QQRKRIEDALK	NLAH	NILKEHP--TD--	NIQLQNDRLFLYYLQNGKDMYTGKSL--D--INQLSSCDIDHI	844	
WP_045635197	776	QQRKRIEDSLK	ILAS	NILKENP--TD--	NNQLQNDRLFLYYLQNGKDMYTGEL--D--INQLSSYDIDHI	843	
WP_002263549	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002263887	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002264920	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002269043	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002269448	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002271977	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002272766	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002273241	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002275430	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002276448	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002277050	779	QQRKRLKEAIK	DLNH	KILKEHP--TD--	NQALQNNRLFLYYLQNGRDMYTGESL--D--INRLSDYDIDHV	846	
WP_002277364	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	HSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002279025	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002279859	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002280230	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002281696	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002282247	779	QQRKRLKEAIK	DLNH	KILKEHP--TD--	NQALQNNRLFLYYLQNGRDMYTGESL--D--INRLSDYDIDHV	846	
WP_002282906	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002283846	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002287255	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	HSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002288990	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002289641	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	
WP_002290427	778	QQRLLKGLTDSIK	EFGS	QILKEHP--VE---	NSQLQNDRLFLYYLQNGRDMYTGEL--D--IDYLSQYDIDHI	841	

WP_002295753	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_002296423	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_002304487	788	QKRYKRLKEAIAK	DLNH	KILKEHP--TD---NQALQNDRLFLYYLQNGRDMYTGEEL--D--INRLSDYDIDHI	855
WP_002305844	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_002307203	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_002310390	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_002352408	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_012997688	778	QQRKGLTDSIK	EFGS	QILKEHP--VK---HSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_014677909	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_019312892	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_019313659	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_019314093	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_019315370	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_019803776	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_019805234	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_024783594	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_024784288	779	QQRKGLTDSIK	DLNH	KILKEHP--TD---NQALQNDRLFLYYLQNGRDMYTGEEL--D--INRLSDYDIDHI	846
WP_024784666	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---HSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_024784894	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_024786433	779	QQRKGLTDSIK	DLNH	KILKEHP--TD---NQALQNDRLFLYYLQNGRDMYTGEEL--D--INRLSDYDIDHI	846
WP_049473442	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
WP_049474547	778	QQRKGLTDSIK	EFGS	QILKEHP--VE---NSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	841
EMC03581	771	QQRKGLTDSIK	EFGS	QILKEHP--VE---HSQLQNDRLFLYYLQNGRDMYTGEEL--D--IDYLSQYDIDHI	834
WP_000428612	779	QQRKRIEDSLK	ILAS	KILKEHP--TD---NIQLQNDRLFLYYLQNGRDMYTGEEL--D--INQLSSYDIDHI	846
WP_000428613	777	QQRKRIEDALK	NLAS	NILKEHP--TN---NIQLQNDRLFLYYLQNGRDMYTGEEL--D--INQLSSYDIDHI	844
WP_049523028	776	QQRKTLSDAIS	ELG-	NILKEHP--TD---NIQLQNDRLFLYYLQNGRDMYTGEEL--D--INQLSNYDIDHI	839
WP_003107102	747	RERLKRLEEVHK	NIGS	KILKEHE--IS---NAQLQSDRVLYLQNGRDMYTGEEL--D--FDRLSQYDIDHI	810
WP_054279288	779	RERMKRVQEVLK	KLGS	QLLKEHP--VE---NFQLQNERLYLQNGRDMYTGEEL--S--ISNLSHYDIDHI	842
WP_049531101	777	QQRKRIEDSLK	ILAS	NILKEHP--TD---NIQLQNDRLFLYYLQNGRDMYTGEEL--D--INHLSSYDIDHI	844
WP_049538452	777	QQRKRIENSJK	ILAS	KILKEHP--TD---NNQLQNDRLFLYYLQNGRDMYTGEEL--D--INQLSSCDIDHI	844
WP_049549711	777	QQRKRIEDSLK	ILAS	NILKENP--TD---NNQLQNDRLFLYYLQNGRDMYTGEEL--D--INQLSSYDIDHI	844
WP_007896501	781	RQRLKKIKEVHK	KTGS	RILEDNserIT---NLTLQDNRLYLLQNGRDMYTGEEL--D--INNLSQYDIDHI	846
EFR44625	733	RQRLKKIKEVHK	KTGS	RILEDNserIT---NLTLQDNRLYLLQNGRDMYTGEEL--D--INNLSQYDIDHI	798
WP_002897477	776	QQRKRIEDALK	NLAP	NILKENP--TD---NIQLKNDRLFLYYLQNGRDMYTGEEL--D--INQLSSYDIDHI	843

WP_002906454	776	QQRKRIEDALK	NLAP	NILKENP--TD---NIQLQNDRLFLYYLQNGKDMYTGKAI--D--INQLSNYDIDHI	843
WP_009729476	777	QQRKRIEDSLK	I LAS	KILKEHP--TD---NIQLQNDRLFLYYLQNGKDMYTGKAI--D--INQLSSCDIDHI	844
CQR24647	780	QQRGLSITKAIQ	DFGS	DILKRYP--VE---NNQLQNDQLYYLQNGKDMYTGDTL--D--IHNLQSDYDIDHI	843
WP_000066813	779	QQRKRIEDSLK	NLAS	NILKENP--TD---NIQLQNDRLFLYYLQNGRDMYTGKPL--E--INQLSNYDIDHI	846
WP_009754323	777	QQRKRIEDALK	NLAP	TISKENP--TD---NIQLQNDRLFLYYLQNGKDMYTGKAI--D--INQLSSYDIDHI	844
WP_044674937	776	QQRKKIENAIK	NLNS	KILKEYP--TN---NQALQNDRLFLYYLQNGKDMYDDEEL--D--IDQLSQYDIDHI	843
WP_044676715	778	QQRKKIENAIK	NLNS	KILKEYP--TN---NQALQNDRLFLYYLQNGKDMYDDEEL--D--IDQLSQYDIDHI	845
WP_044680361	778	QQRKKIENAIK	NLNS	KILKEYP--TN---NQALQNDRLFLYYLQNGKDMYDDEEL--D--IDQLSQYDIDHI	845
WP_044681799	776	QQRKKIENAIK	NLNS	KILKEYP--TN---NQALQNDRLFLYYLQNGKDMYDDEEL--D--IDQLSQYDIDHI	843
WP_049533112	778	QQRKLLQDSLK	PVNI	K-----N-VE---NQQLQNDRLFLYYIQLNGKDMYTGTEL--D--INNLSQYDIDHI	840
WP_029090905	753	TPRDKFIEKAYA	ETDT	EHLKELK--Qr---SKQLSSQRLFLYFIQNGKCMYSGEHL--D--IERLDSYEVVDHI	823
WP_006506696	777	ESKIKKLENVYK	DEQT	SVLEELKg-FDn---TKKISSDSLFLYFTQLGKCMYSGKKL--D--IDSLDKYQIDHI	849
AIT42264	778	REMKRIEIEGK	ELGS	QILKEHP--VE---NTQLQNEKLYLYLQNGRDMYVDQEL--D--INRLSDYDIDHI	841
WP_034440723	785	KARLKKIQEGLE	NLDS	HVEKQAL--D---EEMLKSPKYLYLQNGKDIYTGKDL--D--IGQLQTYDIDHI	848
AKQ21048	778	REMKRIEIEGK	ELGS	QILKEHP--VE---NTQLQNEKLYLYLQNGRDMYVDQEL--D--INRLSDYDIDHI	841
WP_004636532	781	KERLEKITEAIK	EFDG	--VKVD--LK---NENLRNDRLYLYLQNGRDMYDNEPL--D--INNLSKYDIDHI	845
WP_002364836	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_016631044	740	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	803
EMS75795	525	KPRLKALEEALK	SFDS	PLLKEQP--VD---NQALQKDRLYLYLQNGKDMYTGDEL--D--IDRLSEYDIDHI	588
WP_002373311	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_002378009	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_002407324	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_002413717	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_010775580	791	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	854
WP_010818269	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_010824395	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_016622645	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_033624816	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_033625576	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_033789179	789	IQRKIVEKAMA	EIGS	NLLKEQP--TT---NEQLRDTRLFLYMQNGKDMYTGDEL--S--LHRLSHYDIDHI	852
WP_002310644	789	RPRLKALEEALK	DFGS	QLLKEYP--TD---NSSLQKDRLYLYLQNGRDMYTGAPL--D--IHRLSDYDIDHI	852
WP_002312694	790	RPRLKALEEALK	DFGS	QLLKEYP--TD---NSSLQKDRLYLYLQNGRDMYTGAPL--D--IHRLSDYDIDHI	853
WP_002314015	790	RPRLKALEEALK	DFGS	QLLKEYP--TD---NSSLQKDRLYLYLQNGRDMYTGAPL--D--IHRLSDYDIDHI	853
WP_002320716	790	RPRLKALEEALK	DFGS	QLLKEYP--TD---NSSLQKDRLYLYLQNGRDMYTGAPL--D--IHRLSDYDIDHI	853

WP_002330729	789	RPRLKALEESLK	DFGS	QLLKEYP--TD---NSSLQKDRLLYLLQNGRDMYTGAPL--D--IHRLSYDIDDI	852
WP_002335161	790	RPRLKALEESLK	DFGS	QLLKEYP--TD---NSSLQKDRLLYLLQNGRDMYTGAPL--D--IHRLSYDIDDI	853
WP_002345439	790	RPRLKALEESLK	DFGS	QLLKEYP--TD---NSSLQKDRLLYLLQNGRDMYTGAPL--D--IHRLSYDIDDI	853
WP_034867970	781	SPRLKALEENGLK	QIGS	TLLKEQP--TD---NKALQKERLLYLLQNGRDMYTGAPL--E--IENLHQYEVVDHI	844
WP_047937432	790	RPRLKALEESLK	DFGS	QLLKEYP--TD---NSSLQKDRLLYLLQNGRDMYTGAPL--D--IHRLSYDIDDI	853
WP_010720994	781	KPRLKALEENGLK	QIGS	TLLKEQP--TD---NKALQKERLLYLLQNGRDMYTGAPL--E--IENLHQYEVVDHI	844
WP_010737004	781	SPRLKALEENGLK	QIGS	TLLKEQP--TD---NKALQKERLLYLLQNGRDMYTGAPL--E--IENLHQYEVVDHI	844
WP_034700478	781	KPRLKALEENGLK	QIGS	TLLKEQP--TD---NKALQKERLLYLLQNGRDMYTGAPL--E--IENLHQYEVVDHI	844
WP_007209003	782	KPRLKGIENGLK	EFSD	SVLKGSS--ID---NKQLQNDRLYLLQNGKDMYTGHEL--D--IDHLSYDIDDI	845
WP_023519017	775	RPRLKALEEALK	NIDS	PLLKDYP--TD---NQALQKDRLLYLLQNGKDMYTGAPL--E--IHRLEYDIDDI	838
WP_010770040	782	NPRMKALEEAMR	NLRS	NLLKEYP--TD---NQALQNDRLYLLQNGKDMYTGAPL--S--LHNLSSYDIDDI	845
WP_048604708	779	RPRLKNLEKAI	DLDS	EILKKHP--VD---NKALQKDRLLYLLQNGKDMYTGAPL--D--IHKLSYDIDDI	842
WP_010750235	784	KPRLKSLEEALK	NFDS	QLLKERP--VD---NQSLQKDRLLYLLQNGKDMYTGAPL--D--IDRLSEYDIDDI	847
AII16583	817	REMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLLYLLQNGRDMYVDQEL--D--INRLSDYDIDDI	880
WP_029073316	789	DSFVNQMLKLYK	DFED	EANKHLKg--Eda--KSKIRSERLKLYYTQMGKCMYTGKSL--D--IDRLDYQYVDHI	860
WP_031589969	789	DSFVNQMLKLYK	DFED	EANKHLKg--Eda--KSKIRSERLKLYYTQMGKCMYTGKSL--D--IDRLDYQYVDHI	860
KDA45870	772	EDRVQIVKNLK	ELPK	---P---S---NAELSDERKLYLQNGRDMYTGAPL--D--YDHLQFYDIDDI	833
WP_039099354	789	KRQVEQVYQNIS	EL--	EIRNELK--Dl--sNALSNTFLYFMQGGRRDMYTGDSL--N--IDRLSYDIDDI	856
AKP02966	786	RLQSKLLKANG	-LVP	EELKHKn--D---LSSERIMLYFLQNGKSLYSEESL--N--INKLSDYQYVDHI	858
WP_010991369	781	RPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELRNRLYLLQNGKDMYTGQDL--D--IHNLSDYDIDDI	844
WP_033838504	781	RPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELRNRLYLLQNGKDMYTGQDL--D--IHNLSDYDIDDI	844
EHN60060	784	RPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELRNRLYLLQNGKDMYTGQDL--D--IHNLSDYDIDDI	847
EFR89594	550	RPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLLQNGKDMYTGQDL--D--IHNLSDYDIDDI	613
WP_038409211	781	KPRFISLEKAIK	EFGS	QILKEHP--TD---NQCLKNNRLYLLQNGKDMYTGKEL--D--IHNLSDYDIDDI	844
EFR95520	400	KPRFISLEKAIK	EFGS	QILKEHP--TD---NQCLKNNRLYLLQNGKDMYTGKEL--D--IHNLSDYDIDDI	463
WP_003723650	781	KPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLLQNGKDMYTGQEL--D--IHNLSDYDIDDI	844
WP_003727705	781	KPRYKSLEKAIK	DFGS	QILKEHP--TD---NQELKNNRLYLLQNGKDIYTGQEL--D--IHNLSDYDIDDI	844
WP_003730785	781	KPRYKSLEKAIK	DFGS	QILKEHP--TD---NQELKNNRLYLLQNGKDIYTGQEL--D--IHNLSDYDIDDI	844
WP_003733029	781	KPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLLQNGKDIYTGQEL--D--IHNLSDYDIDDI	844
WP_003739838	781	RPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELRNRLYLLQNGKDMYTGQEL--D--IHNLSDYDIDDI	844
WP_014601172	781	KPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELRNRLYLLQNGKDMYTGQEL--D--IHNLSDYDIDDI	844
WP_023548323	781	KPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLLQNGKDMYTGQEL--D--IHNLSDYDIDDI	844
WP_031665337	781	KPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLLQNGKDMYTGQEL--D--IHNLSDYDIDDI	844
WP_031669209	781	KPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLLQNGKDMYTGQEL--D--IHNLSDYDIDDI	844

WP_033920898	781	KPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLQNGKDMYTGQEL--D--IHNLSDYDIDHI	844
AKI42028	784	KPRYKSLEKAIK	EFGS	KILKEHP--TD---NQELKNNRLYLQNGKDMYTGQEL--D--IHNLSDYDIDHI	847
AKI50529	784	KPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLQNGKDMYTGQEL--D--IHNLSDYDIDHI	847
EFR83390	229	RPRYKSLEKAIK	EFGS	QILKEHP--TD---NQELKNNRLYLQNGKDIYTGQEL--D--IHNLSDYDIDHI	292
WP_046323366	781	KPRFTSLEKAIK	ELGS	QILKEHP--TD---NQGLKNDRLYLQNGKDMYTGQEL--D--IHNLSDYDIDHV	844
AKE81011	794	REMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	857
CUO82355	781	ESKIKKLENVYK	DEQT	SVLEELKg-FDn--TKKISSDSLFLYFTQLGKCMYSGKKL--D--IDSLDKYQIDHI	853
WP_033162887	784	ESKIAKLQIYE	NLQT	QVYESLkK-Eda--KKRMETDALYLQNGKCMYSGKPL--D--IDKLSYQIDHI	855
AGZ01981	811	REMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	874
AKA60242	778	REMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDAI	841
AKS40380	778	REMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDHI	841
4UN5_B	782	REMKRIEEGIK	ELGS	QILKEHP--VE---NTQLQNEKLYLQNGRDMYVDQEL--D--INRLSDYDVDAI	845

WP_010922251	842	VPQSF	FLKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_039695303	852	IPQAFI	KDDSIDNRVLTSSAKNRG-KSDD--VP	S--LDIVRARKA-EWVRLYKSGLI	SKRKFNDLTKA--ERGG	LITE	920
WP_045635197	844	IPQAFI	KDDSIDNRVLTSSKDNRG-KSDN--VP	S--IEVQKRKA-FWQQLLDSKLI	SERKFNNLTKA--ERGG	LDE	912
5AXW A	562	IPRSV	FDNSFNNKVLVQEEASK-KGNR--TP	Fqy-LSSSDSKI-SYETF	FKHILNLA	KGGRISKTk-KEYLLEE	632
WP_009880683	526	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWKQLLNAKLI	TQRKFDNLTKA--ERGG	594
WP_010922251	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_011054416	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_011284745	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_011285506	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_011527619	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_012560673	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWKQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_014407541	841	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	909
WP_020905136	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_023080005	841	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	909
WP_023610282	841	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	909
WP_030125963	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_030126706	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_031488318	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWKQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_032460140	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWKQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_032461047	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWKQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_032462016	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_032462936	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_032464890	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWKQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_033888930	667	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	735
WP_038431314	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_038432938	841	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	909
WP_038434062	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWKQLLNAKLI	TQRKFDNLTKA--ERGG	910
BAQ51233	753	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	821
KGE60162	17	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	85
KGE60856							
WP_002989955	842	VPQSF	LKDDSIDNKVLT	TRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLI	TQRKFDNLTKA--ERGG	910
WP_003030002	841	IPQAFI	KDNLNDRVLT	TRSDKNRG-KSDD--VP	S--IEVHEMKS-FWSKLLSVKLI	TQRKFDNLTKA--ERGG	909
WP_003065552	852	IPQAFI	KDDSIDNRVLTSSAKNRG-KSDD--VP	S--LDIVRARKA-EWVRLYKSGLI	SKRKFNDLTKA--ERGG	LITE	920
WP_001040076	847	IPQAFI	KDDSIDNRVLTSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLLD	AKLMSQRKYDNL	TKA--ERGG	915

WP_001040078	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040080	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040081	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040083	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040085	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040087	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040088	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040089	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040090	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040091	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040092	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--IDIVKARKA-FWKLLDACLMSQRKYDNLTKA--ERGG LTP	915
WP_001040094	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--VEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040095	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040096	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040097	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--VEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040098	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--VEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040099	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--VEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040100	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--VEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040104	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040105	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040106	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDN--VP	S--IDIVKARKA-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040107	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDN--VP	S--IDIVKARKA-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040108	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDN--VP	S--IDIVKARKA-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040109	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDN--VP	S--IDIVKARKA-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_001040110	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDN--VP	S--IDIVKARKA-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_015058523	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--IDIVKARKA-FWKLLDACLMSQRKYDNLTKA--ERGG LTP	915
WP_017643650	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--VEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_017647151	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_017648376	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_017649527	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_017771611	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDN--VP	S--IDIVKARKA-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
WP_017771984	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
CFQ25032	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915
CFV16040	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDACLMSQRKYDNLTKA--ERGG LTS	915

KLJ37842	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
KLJ72361	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
KL20707	861	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	929
KL42645	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDN--VP	S--IDIVKARKA-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
WP_047207273	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
WP_047209694	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--VEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
WP_050198062	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
WP_050201642	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
WP_050204027	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDN--VP	S--IDIVKARKA-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
WP_050881965	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
WP_050886065	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
AHN30376	847	VPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--IDIVKARKA-FWKLLDAKLMSQRKYDNLTKA--ERGGTTP	915
EA078426	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
CCW42055	847	IPQAFIKDDSIDNRVLVSSAKNRG-KSDD--VP	S--LEIVKDKV-FWKLLDAKLMSQRKYDNLTKA--ERGGTTS	915
WP_003041502	841	IPQAYIKDDSFDRNRLVTSSENRRG-KSDN--VP	S--IEVVCARKA-DWMRLRKAAGLISQRKFDNLTKA--ERGGLTE	909
WP_037593752	842	IPQAFIKDNLDRNRLTRSDKNRG-KSDD--VP	S--IEVHEMKS-FWSKLLSVKLIQTKRFDNLTKA--ERGGLTE	910
WP_049516684	842	IPQAFIKDNLDRNRLTRSDKNRG-KSDD--VP	S--IEVHEMKS-FWSKLLSVKLIQTKRFDNLTKA--ERGGLTE	910
GAD46167	841	IPQAFIKDNLDRNRLTRSDKNRG-KSDD--VP	S--IEVHEMKS-FWSKLLSVKLIQTKRFDNLTKA--ERGGLTE	909
WP_018363470	850	IPQAFIKDDSIDNRVLTSSAKNRG-KSDD--VP	S--LGIVRARKA-EWRLYKSGLISKRKFDNLTKA--ERGGLTE	918
WP_003043819	851	VPQSFIKDDSIDNKVLTSSAKNRG-KSDD--VP	S--EEVVKMKN-YWRQLLNAKLITQTKRFDNLTKA--ERGGTSE	919
WP_006269658	841	IPQAFIKDNLDRNRLTRSDKNRG-KSDD--VP	S--IEVHEMKS-FWSKLLSVKLIQTKRFDNLTKA--ERGGTSE	909
WP_048800889	841	IPQAFIKDDSIDNRVLTSSAKNRG-KSDD--VP	N--LEVVCDRKA-DWIRLREAGLISQRKFDNLTKA--ERGGTSE	909
WP_012767106	841	VPQSFIKDDSIDNKILTRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLITQTKRFDNLTKA--ERGGTSE	909
WP_014612333	841	VPQSFIKDDSIDNKVLTSSAKNRG-KSDD--VP	S--EEVVKMKN-YWRQLLNAKLITQTKRFDNLTKA--ERGGTSE	909
WP_015017095	841	VPQSFIKDDSIDNKVLTSSAKNRG-KSDD--VP	S--EEVVKMKN-YWRQLLNAKLITQTKRFDNLTKA--ERGGTSE	909
WP_015057649	841	VPQSFIKDDSIDNKVLTSSAKNRG-KSDD--VP	S--EEVVKMKN-YWRQLLNAKLITQTKRFDNLTKA--ERGGTSE	909
WP_048327215	841	VPQSFIKDDSIDNKVLTSSAKNRG-KSDD--VP	S--EEVVKMKN-YWRQLLNAKLITQTKRFDNLTKA--ERGGTSE	909
WP_049519324	841	VPQSFIKDDSIDNKVLTSSAKNRG-KSDD--VP	S--EEVVKMKN-YWRQLLNAKLITQTKRFDNLTKA--ERGGTSE	909
WP_012515931	841	IPQSFIKNNSIDNKVLTSSAKNRG-KSDN--VP	S--EAIVRKMG-YWQSLLRAGLISQRKFDNLTKA--ERGGLTQ	909
WP_021320964	841	IPQSFIKNNSIDNKVLTSSAKNRG-KSDN--VP	S--EAIVRKMG-YWQSLLRAGLISQRKFDNLTKA--ERGGLTQ	909
WP_037581760	841	IPQSFIKNNSIDNKVLTSSAKNRG-KSDN--VP	S--EAIVRKMG-YWQSLLRAGLISQRKFDNLTKA--ERGGLTQ	909
WP_004232481	849	IPQAFIKDNLDRNRLVTSSENRRG-KSDD--VP	S--IEIVRNRKS-YWYKLYKSGLISKRKFDNLTKA--ERGGLTE	917
WP_009854540	850	IPQAFIKDDSIDNRVLTSSAKNRG-KSDD--VP	S--LDIVRARKA-EWVRLYKSGLISKRKFDNLTKA--ERGGTSE	918
WP_012962174	850	IPQAFIKDDSIDNRVLTSSAKNRG-KSDD--VP	S--LDIVHDRKA-DWIRLKYKSGLISKRKFDNLTKA--ERGGTSE	918

WP_039695303	852	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--LDIVRARKA-EMVRLYKSGLSIKRKFDFNLTKA--ERGGGLTE	920
WP_014334983	849	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--IEIVNRNRS-YWYKLYKSGLSIKRKFDFNLTKA--ERGGGLTE	917
WP_003099269	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDN--VP	N--IETVNMKS-FWYKQLKSGAISQRKFDFNLTKA--ERGALS	910
AHY15608	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDN--VP	N--IETVNMKS-FWYKQLKSGAISQRKFDFNLTKA--ERGALS	910
AHY17476	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDN--VP	N--IETVNMKS-FWYKQLKSGAISQRKFDFNLTKA--ERGALS	910
ESR09100				
AGM98575	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDN--VP	N--IETVNMKS-FWYKQLKSGAISQRKFDFNLTKA--ERGALS	910
ALF27331	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_018372492	858	VPRSYKNDSDFNKVLTSKGNRK-KLDD--VP	A--KEVVEKEMEN-TWRRLHAAGLISDIKLSYLMKGe-----LITE	923
WP_045618028	845	IPQAFIKDSDSDNRVLTSSAKNRG-KSDN--VP	S--LEIVQKRKA-FWQQLLSDSKLISERKFDFNLTKA--ERGGGLD	913
WP_045635197	844	IPQAFIKDSDSDNRVLTSSAKNRG-KSDN--VP	S--IEVQKRKA-FWQQLLSDSKLISERKFDFNLTKA--ERGGGLD	912
WP_002263549	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002263887	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002264920	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002269043	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002269448	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--EDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002271977	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002272766	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002273241	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002275430	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002276448	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002277050	847	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--EDVVRMRP-FWNKLLSSGLISQRKYNNLTKK--E--LTP	912
WP_002277364	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002279025	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002279859	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002280230	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002281696	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002282247	847	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--EDVVRMRP-FWNKLLSSGLISQRKYNNLTKK--E--LTL	912
WP_002282906	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002283846	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002287255	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002288990	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002289641	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910
WP_002290427	842	IPQAFIKDSDSDNRVLTSSAKNRG-KSDD--VP	S--KDVVRKMS-YWSKLLSAKLITQRKFDFNLTKA--ERGGGLD	910

WP_002295753	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_002296423	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KNVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_002304487	856	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--EEVVRKMKP-FWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	924
WP_002305844	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_002307203	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_002310390	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_002352408	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_012997688	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KNVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_014677909	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_019312892	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KNVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_019313659	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_019314093	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_019315370	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMKP-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_019803776	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_019805234	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_024783594	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_024784288	847	IPQAFIKDNSIDNRVLTSSKANRG-KSDD--VP	S--EDVVRMRP-FWNKLLSSGLISQRKYNNLTKK--E---LTL	912
WP_024784666	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KNVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_024784894	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_024786433	847	IPQAFIKDNSIDNRVLTSSKANRG-KSDD--VP	S--EDVVRMRP-FWNKLLSSGLISQRKYNNLTKK--E---LTL	912
WP_049473442	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMKP-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
WP_049474547	842	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KDVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	910
EMC03581	835	IPQAFIKDNSIDNRVLTSSKENRG-KSDD--VP	S--KNVVRKMK-SYWSKLLSAKLITQRKFDNLTKA--ERGGGLTD	903
WP_000428612	847	VPQAFIKDDSLDNRVLTSLKDNRG-KSDN--VP	S--LEVEKMKT-FWQQLLDSKLSYRKFNLTKA--ERGGGLDE	915
WP_000428613	845	VPQAFIKDDSLDNRVLTSLKDNRG-KSDN--VP	S--IEVVRKKA-FWQQLLDSKLSERKFNLTKA--ERGGGLDE	913
WP_049523028	840	IPQAFIKDDSLDNRVLTSSKDNRG-KSDN--VP	S--LEIVEKMG-FWQQLLDSKLSERKFNLTKA--ERGGGLDE	908
WP_003107102	811	IPQSFIKDNSIDNIVLTQESNRG-KSDN--VP	Y--IAIVNRKMK-SYWSKLLSAGAIQRKFDNLTKA--ERGGGLSE	879
WP_054279288	843	IPRSFIKDDSIDNKVLTTRSEHNRG-KTDN--VP	S--IEVVRKMKP-YWQKLLDTKVIQRKFDNLTKA--ERGGLOE	911
WP_049531101	845	IPQAFIKDDSLDNRVLTSSKDNRG-KSDN--VP	S--LEVVRKKA-FWQQLLDSKLSERKFNLTKA--ERGGGLNE	913
WP_049538452	845	IPQAFIKDDSLDNRVLTSSKENRG-KSDN--VP	C--LEVVRKMKV-FWQQLLDFKLSYRKFNLTKA--ERGGGLDE	913
WP_049549711	845	IPQAFIKDDSLDNRVLTSSKDNRG-KSDN--VP	S--LEVVRKKA-FWQQLLDSKLSERKFNLTKAerERDGLNE	915
WP_007896501	847	IPQSFIKDNSIDNLVLTQKANRG-KSDN--VP	S--IEVVRMKDrVWRRQLANGAISQRKFDHLTKA--ERGGGLAD	916
EFR44625	799	IPQSFIKDNSIDNLVLTQKANRG-KSDN--VP	S--IEVVRMKDrVWRRQLANGAISQRKFDHLTKA--ERGGGLAD	868
WP_002897477	844	IPQAFIKDDSIDNRVLTSSKDNRG-KSDN--VP	S--LEVVRKKA-FWQQLLDSKLSERKFNLTKA--ERGGGLDE	912

WP_002906454	844	IPQAFIKDDSLDNRVLTSSKDNRG-KSDN--VP	S--IEVQKRKA-FWQQLDLSKLI SERKFNNLTKA--KRGGLDE	912
WP_009729476	845	IPQAFIKDDSLDNRVLTSSKDNRG-KSDN--VP	S--LEVVDKMKV-FWQQLDLSKLI SYRKFNNTKA--ERGGLE	913
CQR24647	844	IPQSFIKDNLNDRVLTNSKSNRG-KSDN--VP	S--NEVVKRMKG-FWLKQLDAKLI SQRKFDNLTKA--ERGGLSA	912
WP_000066813	847	IPQAFIKDDSLDNRVLTSSKDNRG-KSDN--VP	S--LEVEKMKKA-FWQQLDLSKLI SERKFNNLTKAerERGGLE	917
WP_009754323	845	IPQAFIKDDSLDNRVLTSSKDNRG-KSDN--VP	S--LEVVKRKA-FWQQLDLSKLI SERKFNNLTKA--ERGGLE	913
WP_044674937	844	IPQAFIKDDSLDNKVLTKSAKNG-KSDD--VP	S--LEIVHKKKN-FWKQLDLSQLI SQRKFDNLTKA--ERGGLTN	912
WP_044676715	846	IPQAFIKDDSLDNKVLTKSAKNG-KSDD--VP	S--LEIVHKKKN-FWKQLDLSQLI SQRKFDNLTKA--ERGGLTN	914
WP_044680361	846	IPQAFIKDDSLDNKVLTKSAKNG-KSDD--VP	S--LEIVHKKKN-FWKQLDLSQLI SQRKFDNLTKA--ERGGLTN	914
WP_044681799	844	IPQAFIKDDSLDNKVLTKSAKNG-KSDD--VP	S--LEIVHKKKN-FWKQLDLSQLI SQRKFDNLTKA--ERGGLTN	912
WP_049533112	841	IPQAFIKDDSFDRVLTSSSENRG-KSDN--VP	S--IEVVRARKA-DWMRLRKAGLI SQRKFDNLTKA--ERGGLTE	909
WP_029090905	824	LPQSYIKDNSIENLALVKVENQR-KKDSL1LN	S---SIINQNS-RWEQLKNAGLIGEKKFRNLTTRk-----ITD	890
WP_006506696	850	VPQSLVKDDSFDRVLTSSSENQR-KLDD1VVP	---FDIRDKMYR-FWKLLFDHELISPKKFYSLIKTe-----YTE	916
AIT42264	842	VPQSFVKDDSIDNKVLTTRSDKNRG-KSDN--VP	S--EEVVKMKKN-YWRQLLNAKLITQRKFDNLTKA--ERGGLE	910
WP_034440723	849	IPRSFITDNSFDNLVLTSSTVNRG-KLDN--VP	Sp--DIVRQKKG-FWKQLLRAGLMSQRKFNNTKGGk-----LTD	914
AKQ21048	842	VPQSFVKDDSIDNKVLTTRSDKNRG-KSDN--VP	S--EEVVKMKKN-YWRQLLNAKLITQRKFDNLTKA--ERGGLE	910
WP_004636532	846	IPQSFITDNSIDNKVLTSTRTKNQnKSD--VP	S--INIVHKMKP-FWRQLHKAAGLISDRKFNLTKA--EHGGLTE	915
WP_002364836	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_016631044	804	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	872
EMS75795	589	IPRSFIVDNSIDNKVIVSSKENRL-KMDD--VP	D--QKVIRMR-WEKLLRANLISERKFAYLTkLe-----LTP	654
WP_002373311	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KVVVKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_002378009	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_002407324	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_002413717	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_010775580	855	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	923
WP_010818269	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_010824395	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_016622645	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_033624816	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_033625576	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_033789179	853	IPQSFMKDDSLDNLVVGSTENRG-KSDD--VP	S--KEVKDKMKA-YWEKLYAAGLISQRKFQRLTKG--EQGGTLT	921
WP_002310644	853	IPRSFTTDNSIDNKVIVSSKENRL-KKDD--VP	S--EKVVKMRS-FWYDLYSSKLI SKRKL DNLTKik-----LTe	918
WP_002312694	854	IPRSFTTDNSIDNKVIVSSKENRL-KKDD--VP	S--EKVVKMRS-FWYDLYSSKLI SKRKL DNLTKik-----LTe	919
WP_002314015	854	IPRSFTTDNSIDNKVIVSSKENRL-KKDD--VP	S--EKVVKMRS-FWYDLYSSKLI SKRKL DNLTKik-----LTe	919
WP_002320716	854	IPRSFTTDNSIDNKVIVSSKENRL-KKDD--VP	S--EKVVKMRS-FWYDLYSSKLI SKRKL DNLTKik-----LTe	919

WP_002330729	853	IPRSFTTDSIDNKVLSVSSKENRL-KKDD--VP	S--EKVVKMRS-FWYDLYSSKLI SKRKL DNLTKIK-----LTE	918
WP_002335161	854	IPRSFTTDSIDNKVLSVSSKENRL-KKDD--VP	S--EKVVKMRS-FWYDLYSSKLI SKRKL DNLTKIK-----LTE	919
WP_002345439	854	IPRSFTTDSIDNKVLSVSSKENRL-KKDD--VP	S--EKVVKMRS-FWYDLYSSKLI SKRKL DNLTKIK-----LTE	919
WP_034867970	845	IPRSFIVDSIDDKVAVASKQNK-KRDD--VP	K--KQIVNEQRI-FWNQLKEAKLIS TKKAYALTKE-----LTP	910
WP_047937432	854	IPRSFTTDSIDNKVLSVSSKENRL-KKDD--VP	S--EKVVKMRS-FWYDLYSSKLI SKRKL DNLTKIK-----LTE	919
WP_010720994	845	IPRSFIVDSIDNKVAVASKQNK-KRDD--VP	K--KQIVNEQRI-FWNQLKEAKLIS PKKYAYLTKE-----LTP	910
WP_010737004	845	IPRSFIVDSIDNKVAVASKQNK-KRDD--VP	K--KQIVNEQRI-FWNQLKEAKLIS PKKYAYLTKE-----LTP	910
WP_034700478	845	IPRSFIVDSIDNKVAVASKQNK-KRDD--VP	N--KQIVNEQRI-FWNQLKEAKLIS PKKYAYLTKE-----LTP	910
WP_007209003	846	IPQSFLTDSIDNRVLTTSKSNRG-KSDN--VP	S--EEVVRKMDR-FWRKLLNAKLI SERKYTNLTKE-----LTE	911
WP_023519017	839	IPRSFIVDSIDNKVLSVSKVNRG-KLDN--AP	D--PLVVKRMS-HWEKLGQA KLI SDKLANLTKQN-----LTE	904
WP_010770040	846	VPQSFTTDSIDNRVLSVSSKENRG-KKDD--VP	S--KEVWQKNIT-LWETLKN SNLIS QKKYDNLTKG--LRGGLTE	914
WP_048604708	843	IPQSFI VDSIDNRVLSVSSKENRG-KLDD--VP	S--KEVWVKMRA-FWESLYRSGLIS KKKFDNLVKA--ESGGLSE	911
WP_010750235	848	IPRSFIVDHSIDNKVLSVSSKENRL-KKDD--VP	D--SKVVKRMA-YWEKLLRANLIS ERKFSYLTKE-----LTD	913
AII16583	881	VPQSFLKDDSIDNKVLT RSDKNRG-KSDN--VP	S--EEVVKMKMKN-YWRQLLNAKLI TQRKFDNLTKA--ERGGGLSE	949
WP_029073316	861	VPQSLLKDDSIDNKVLSVSSSENQR-KLDDVIP	---EMIRNKMGF-FWNKLYENKII SPKKFYSLIKSE-----YSD	927
WP_031589969	861	VPQSLLKDDSIDNKVLSVSSSENQR-KLDDVIP	---SSIRNKMYG-FWEKLFNNKII SPKKFYSLIKTE-----FNE	927
KDA45870	834	IPQSFLKDDSIENKVLTIKKENVR-KTNG--LP	S--EAVIQMGS-FWKLLDAGAMTNKKYDNLRRN1--HGGLNE	902
WP_039099354	857	LPQSFIKDDSIDNRVLSVQRMNR-KADQ--VP	S--VELGQKMQI-QWEQMLRAGLITKKKYDNLTLNp-----	923
AKP02966	859	LPRTYIPDDSENKALVLAKENQR-KADD1LN	S--NVIDKNLE-RWTYMLNNMMGLKKFKNLTTRV-----ITD	925
WP_010991369	845	VPQSFI TDSIDNLVLTSSAGNRE-KGDD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
WP_033838504	845	VPQSFI TDSIDNLVLTSSAGNRE-KGDD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
EHN60060	848	VPQSFI TDSIDNLVLTSSAGNRE-KGDD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	916
EFR89594	614	VPQSFI TDSIDNLVLTSSAGNRE-KGND--VP	P--LEIVQKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	682
WP_038409211	845	IPQSFI TDSIDNRVLSVSTANRE-KGDN--VP	L--LEIVRKRKA-FWEKLYQAKLMSKRKFDYLTKA--ERGGGLTE	913
EFR95520	464	IPQSFI TDSIDNRVLSVSTANRE-KGDN--VP	L--LEIVRKRKA-FWEKLYQAKLMSKRKFDYLTKA--ERGGGLTE	532
WP_003723650	845	VPQSFI TDSIDNLVLTSSAGNRE-KGGD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
WP_003727705	845	VPQSFI TDSIDNLVLTSSAGNRE-KGGD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
WP_003730785	845	VPQSFI TDSIDNLVLTSSAGNRE-KGGD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
WP_003733029	845	VPQSFI TDSV DNLVLTSSAGNRE-KGDN--VP	P--LEIVQKRKI-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
WP_003739838	845	VPQSFI TDSIDNLVLTSSAGNRE-KGDD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
WP_014601172	845	VPQSFI TDSIDNLVLTSSAGNRE-KGGD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTD	913
WP_023548323	845	VPQSFI TDSIDNLVLTSSAGNRE-KGDN--VP	P--LEIVQKRKI-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
WP_031665337	845	VPQSFI TDSIDNLVLTSSAGNRE-KGGD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913
WP_031669209	845	VPQSFI TDSV DNLVLTSSAGNRE-KGDN--VP	P--LEIVQKRKI-FWEKLYQGNLMSKRKFDYLTKA--ERGGGLTE	913

WP_033920898	845	VPQSFITDNSIDNVLVTSSAGNRE-KGDN--VP	P--LEIVQKRKI-FWEKLYQGNLMSKRKFDYLTKA--ERGGLTE	913
AKI42028	848	VPQSFITDNSIDNVLVTSSAGNRE-KGGD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGLTID	916
AKI50529	848	VPQSFITDNSIDNVLVTSSAGNRE-KGDN--VP	P--LEIVQKRKI-FWEKLYQGNLMSKRKFDYLTKA--ERGGLTE	916
EFR83390	293	VPQSFITDNSIDNVLVTSSAGNRE-KGDD--VP	P--LEIVRKRKV-FWEKLYQGNLMSKRKFDYLTKA--ERGGLTE	361
WP_046323366	845	VPQSFITDNSIDNRVLASSAANRE-KGDN--VP	S--LEVVRKRKV-YWEKLYQAKLMSKRKFDYLTKA--ERGGLTE	913
AKE81011	858	VPQSFVKDDSIDNKVLTTRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLITQRKFDNLTKA--ERGGTSE	926
CUO82355	854	VPQSLVKDDSFDRVLPSENOQ-KLDDLVVP	---FDIRDKMYR-FWKLLFDHELISPKKFYSLIKTE-----YTE	920
WP_033162887	856	LPQSLIKDDSFDRVLPPEENQW-KLDSetVP	---FEIRNKMIG-FWQMLHENGMSNKKFFSLIRTd-----FSD	922
AGZ01981	875	VPQSFVKDDSIDNKVLTTRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLITQRKFDNLTKA--ERGGTSE	943
AKA60242	842	VPQSFVKDDSIDNKVLTTRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLITQRKFDNLTKA--ERGGTSE	910
AKS40380	842	VPQSFVKDDSIDNKVLTTRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLITQRKFDNLTKA--ERGGTSE	910
4UN5_B	846	VPQSFVKDDSIDNKVLTTRSDKNRG-KSDN--VP	S--EEVVKMKN-YWRQLLNAKLITQRKFDNLTKA--ERGGTSE	914

WP_010922251	911	Ld	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_039695303	921	AD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	EYKVRREINDY	991
WP_045635197	913	RD	KVGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--TVKIIITLKSNLVSNFRKEF	RLYKVRREINDY	983
5AXW A	633	RD	QKDFINRNLDVTRYATRGLMNLRSYFR-----VnLDVKVKSINGGFTSFLRRKW	KFKKERNGYK	702
WP_009880683	595	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVRVITLKSKLVSDFRCKDF	QFYKVRREINNY	665
WP_010922251	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_011054416	911	LD	KVGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVRVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_011284745	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_011285506	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_011527619	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_012560673	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVRVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_014407541	910	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	980
WP_020905136	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_023080005	910	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	980
WP_023610282	910	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	980
WP_030125963	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_030126706	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_031488318	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_032460140	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVRVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_032461047	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVRVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_032462016	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVRVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_032462936	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVRVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_032464890	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_033888930	736	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	806
WP_038431314	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_038432938	910	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	980
WP_038434062	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
BAQ51233	822	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	892
KGE60162	86	LD	KVGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVRVITLKSKLVSDFRCKDF	QFYKVRREINNY	156
KGE60856		--	-----	-----	
WP_002989955	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	QFYKVRREINNY	981
WP_003030002	910	ED	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--NVKIIITLKSNLVSNFRKEF	ELYKVRREINDY	980
WP_003065552	921	AD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSKLVSDFRCKDF	EYKVRREINDY	991
WP_001040076	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRIR--KVKIVITLKSNLVSNFRKEF	VFYKIREVANNY	986

WP_001040078	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040080	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040081	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040083	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040085	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040087	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040088	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040089	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040090	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040091	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040092	916	DD	KAGFIQRQLVETRQITKHVARILDERFNNKVDNNKPIR--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040094	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040095	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040096	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040097	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040098	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040099	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040100	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040104	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040105	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040106	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040107	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040108	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040109	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_001040110	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_015058523	916	DD	KAGFIQRQLVETRQITKHVARILDERFNNKVDNNKPIR--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_017643650	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_017647151	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_017648376	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_017649527	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_017771611	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
WP_017771984	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
CFQ25032	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986
CFV16040	916	DD	KARFIQRQLVEIRQITKHVARILDERFNNELDSKGRRI--KVIVTLKSNLVSFRKEF	GFYKIREVNDY	986

KLJ37842	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
KLJ72361	916	DD	KARFIQRQLVETRQITKHVARILDELFFNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
KL20707	930	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	1000
KL42645	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNDY	986
WP_047207273	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
WP_047209694	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
WP_050198062	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
WP_050201642	916	DD	KARFIQRQLVETRQITKHVASILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
WP_050204027	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNDY	986
WP_050881965	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
WP_050886065	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
AHN30376	916	DD	KAGFIQRQLVETRQITKHVARILDERFNNKVDNPKPIR--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
EA078426	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNNY	986
CCW42055	916	DD	KARFIQRQLVETRQITKHVARILDERFNNELDSKGRRI--KVKIIVTKSNLVSFRKEF	GFYKIREVNDY	986
WP_003041502	910	ND	KAGFIKRQLVETRQITKHVAQVLDARFNKHDENKVVIR--DVKIITLKSNIQSFRKDF	KFYKIREVNDY	980
WP_037593752	911	ED	KAGFIKRQLVETRQITKHVAQILDERFTEFDGAQRRIR--NVKIITLKSNIQSFRKEF	ELYKIREVNDY	981
WP_049516684	911	ED	KAGFIKRQLVETRQITKHVAQILDERFTEFDGAQRRIR--NVKIITLKSNIQSFRKEF	ELYKIREVNDY	981
GAD46167	910	ED	KAGFIKRQLVETRQITKHVAQILDERFTEFDGAQRRIR--NVKIITLKSNIQSFRKEF	ELYKIREVNDY	980
WP_018363470	919	AD	KAGFIKRQLVETRQITKHVAQILDSRMTEKRDNDKPIR--EVKVIITLKSNIQSFRKDF	KFYKIREVNDY	989
WP_003043819	920	AD	KAGFIKRQLVETRQITKHVAQILDERFTEFDGNKRRIR--NVKIITLKSNIQSFRKDF	QLYKVRDINNY	990
WP_006269658	910	ED	KAGFIKRQLVETRQITKHVAQILDERFTEFDGNKRRIR--NVKIITLKSNIQSFRKDF	ELYKIREVNDY	980
WP_048800889	910	ND	KAGFIHRQLVETRQITKHVAQILDSRMTEKRDNDKPIR--EVKVIITLKSNIQSFRKDF	KLYKVRDINNY	980
WP_012767106	910	LD	KAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIR--EVKVIITLKSNIQSFRKDF	QFYKVRDINNY	980
WP_014612333	910	LD	KAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIR--EVKVIITLKSNIQSFRKDF	QFYKVRDINNY	980
WP_015017095	910	LD	KAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIR--EVKVIITLKSNIQSFRKDF	QFYKVRDINNY	980
WP_015057649	910	LD	KAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIR--EVKVIITLKSNIQSFRKDF	QFYKVRDINNY	980
WP_048327215	910	LD	KAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIR--EVKVIITLKSNIQSFRKDF	QFYKVRDINNY	980
WP_049519324	910	LD	KAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIR--EVKVIITLKSNIQSFRKDF	QFYKVRDINNY	980
WP_012515931	910	VD	KAGFIQRQLVETRQITKHVAQILDSRFTEFDHDKRRIR--KVHIITLKSNIQSFRKDF	GLYKIRDINHY	980
WP_021320964	910	VD	KAGFIQRQLVETRQITKHVAQILDSRFTEFDHDKRRIR--KVHIITLKSNIQSFRKDF	GLYKIRDINHY	980
WP_037581760	910	VD	KAGFIQRQLVETRQITKHVAQILDSRFTEFDHDKRRIR--KVHIITLKSNIQSFRKDF	GLYKIRDINHY	980
WP_004232481	918	TD	KAGFIKRQLVETRQITKHVAQILDSRFTEFDHDKRRIR--DVKVIITLKSNIQSFRKDF	KFYKVRDINNY	988
WP_009854540	919	AD	KAGFIKRQLVETRQITKHVAQILDSRFTEHDENDKVVIR--DVKVIITLKSNIQSFRKDF	EFYKVRDINNY	989
WP_012962174	919	ND	KAGFIKRQLVETRQITKHVAQILDSRFTEHDENDKVVIR--NVKVIITLKSNIQSFRKDF	KFYKVRDINNY	989

WP_039695303	921	AD	KAGFIKRQLVETRQITKHVAQIILDARFNTEHDENDKVR--DVKVIITLKSNIIVSQFRKDF	EFYKVRREINDY	991
WP_014334983	918	AD	KAGFIKRQLVETRQITKHVAQIILDARFNTRKRENDKVR--DVKVIITLKSNIIVSQFRKEF	KFYKVRREINDY	988
WP_003099269	911	FD	KAGFIKRQLVETRQITKHVAQIILDSRFNSNLTEDESKNR--NVKIIITLKSNIIVSDFRKF	GFYKLRREINDY	981
AHY15608	911	FD	KAGFIKRQLVETRQITKHVAQIILDSRFNSNLTEDESKNR--NVKIIITLKSNIIVSDFRKF	GFYKLRREINDY	981
AHY17476	911	FD	KAGFIKRQLVETRQITKHVAQIILDSRFNSNLTEDESKNR--NVKIIITLKSNIIVSDFRKF	GFYKLRREINDY	981
ESR09100		--	-----	-----	
AGM98575	911	FD	KAGFIKRQLVETRQITKHVAQIILDSRFNSNLTEDESKNR--NVKIIITLKSNIIVSDFRKF	GFYKLRREINDY	981
ALF27331	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_018372492	924	ED	KAGFIKRQLVETRQITKHVARILDEKLNKKNENGEKLR--TTKIIITLKSNIIVSDFRFRANF	DLYKLRREINDY	994
WP_045618028	914	RD	KVGFIKRQLVETRQITKHVAQIILDARFNTEVTEKDKKDR--SVKIIITLKSNIIVSNFRKEF	RLYKVRREINDY	984
WP_045635197	913	RD	KVGFIKRQLVETRQITKHVAQIILDARFNTEVNEKDKKDR--TVKIIITLKSNIIVSNFRKEF	RLYKVRREINDY	983
WP_002263549	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002263887	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002264920	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002269043	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002269448	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002271977	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002272766	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002273241	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002275430	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002276448	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002277050	913	DD	KAGFIKRQLVETRQITKHVARMLDERFNKEFDDNKKIR--RVKIVITLKSNIIVSFRKEF	ELYKVRREINDY	983
WP_002277364	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002279025	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002279859	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002280230	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002281696	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002282247	913	DD	KAGFIKRQLVETRQITKHVARMLDERFNKEFDDNKKIR--RVKIVITLKSNIIVSFRKEF	ELYKVRREINDY	983
WP_002282906	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002283846	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002287255	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002288990	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002289641	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981
WP_002290427	911	DD	KAGFIKRQLVETRQITKHVARILDERFNTETDENKKIR--QVKIVITLKSNIIVSNFRKEF	ELYKVRREINDY	981

WP_002295753	911	DD	KAGFIKRQLVETRQITKHVARILDERFYTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_002296423	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_002304487	925	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	995
WP_002305844	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_002307203	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_002310390	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_002352408	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_012997688	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_014677909	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_019312892	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_019313659	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_019314093	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_019315370	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_019803776	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_019805234	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_024783594	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_024784288	913	DD	KAGFIKRQLVETRQITKHVARMLDERFHTETDENKKIR--RVKIVTLKSNLVSFRKEF	ELYKVRREINDY	983
WP_024784666	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_024784894	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_024786433	913	DD	KAGFIKRQLVETRQITKHVARMLDERFHTETDENKKIR--RVKIVTLKSNLVSFRKEF	ELYKVRREINDY	983
WP_049473442	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
WP_049474547	911	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	981
EMC03581	904	DD	KAGFIKRQLVETRQITKHVARILDERFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	974
WP_000428612	916	RD	KVGFIKRQLVETRQITKHVAQILDARFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	986
WP_000428613	914	RD	KVGFIKRQLVETRQITKHVAQILDARFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	ELYKVRREINDY	984
WP_049523028	909	RD	KVGFIKRQLVETRQITKHVAQILDDRFAEVEKKNR--TVKIIITLKSNLVSFRKEF	ELYKVRREINDY	979
WP_003107102	880	YD	KAGFIKRQLVETRQITKHVAQILNRRFNNVDDSSKNR--SVKIIITLKSNLVSFRKEF	GLYKVRREINDY	950
WP_054279288	912	SD	KANFIQRQLVETRQITKHVAQILDSRFNTERDEKDRPIR--RVKIVTLKSKFVSDFRQDF	GFYKIREVNDY	982
WP_049531101	914	RD	KVGFIKRQLVETRQITKHVAQILDSRFNTERDEKDRPIR--RVKIVTLKSKFVSDFRQDF	GFYKIREVNDY	982
WP_049538452	914	RD	KVGFIRRQLVETRQITKHVAQILDSRFNTERDEKDKNR--NVKIIITLKSNLVSFRKEF	ELYKVRREINDY	984
WP_049549711	916	LD	KVGFIKRQLVETRQITKHVAQILDSRFNTERDEKDKNR--NVKIIITLKSNLVSFRKEF	ELYKVRREINDY	984
WP_007896501	917	SD	KARFLRRQLVETRQITKHVAQILLDSRFNSKNQKKLAR--NVKIIITLKSKIVSDFRKF	GLYKLRVANNY	987
EFR44625	869	SD	KARFLRRQLVETRQITKHVAQILLDSRFNSKNQKKLAR--NVKIIITLKSKIVSDFRKF	GLYKLRVANNY	939
WP_002897477	913	RD	KVGFIRRQLVETRQITKNVAQILDARFHTETDENKKIR--QVKIVTLKSNLVSFRKEF	GLYKVRREINNY	983

WP_002906454	913	RD	KVGFIKRQLVETRQITKHVAQVLLDTRFNTEVNEENQKIR--TVKIIITLKSNLVSNFRKEF	GLYKVVREINDY	983
WP_009729476	914	LD	KVGFIKRQLVETRQITKHVAQVLLDARFNKEVTEKDKKNR--TVKIIITLKSNLVSNFRKEF	ELYKVVREINDY	984
CQR24647	913	ED	KAGFIKRQLVETRQITKHVARILDERFNDRDFDKDKRIR--NVKIVITLKSNLVSNFRKEF	GFYKVVREINNF	983
WP_000066813	918	LD	KVGFIKRQLVETRQITKHVAQVLLDARFNKEVTEKDKKNR--NVKIIITLKSNLVSNFRKEF	GLYKVVREINDY	988
WP_009754323	914	RD	KVGFIKRQLVETRQITKHVARILDARFNTEVSEKNQKIR--SVKIIITLKSNLVSNFRKEF	KLYKVVREINDY	984
WP_044674937	913	ED	KARFIQRQLVETRQITKHVARILDTRFNTKLDEAGNRIRdpKVNIITLKSNLVSNFRKDY	QLYKVVREINNY	985
WP_044676715	915	ED	KARFIQRQLVETRQITKHVARILDTRFNTKLDEAGNRIRdpKVNIITLKSNLVSNFRKDY	QLYKVVREINNY	987
WP_044680361	915	ED	KARFIQRQLVETRQITKHVARILDTRFNTKLDEAGNRIRdpKVNIITLKSNLVSNFRKDY	QLYKVVREINNY	987
WP_044681799	913	ED	KARFIQRQLVETRQITKHVARILDTRFNTKLDEAGNRIRdpKVNIITLKSNLVSNFRKDY	QLYKVVREINNY	985
WP_049533112	910	ND	KAGFIKRQLVETRQITKHVAQVLLDARFNAKHDKKVVIR--DVKIIITLKSNLVSNFRKDF	KFYKVVREINDY	980
WP_029090905	891	RD	KEGFIARQLVETRQITKHVTQLLQOEY-----K-dTTKVFAIKATLVSLGLRRKF	EFIKNRNVNDY	951
WP_006506696	917	RD	EERFINRQLVETRQITKNVTQIIEHDHYSY-----TKVAAIRANLSHEFRVKN	HIYKNRDINDY	976
AIT42264	911	LD	KAGFIKRQLVETRQITKHVAQVLLDSRMNTKYDENDKLIR--EVKVIITLKSCLVSDFRKDF	QFYKVVREINNY	981
WP_034440723	915	RD	RQOFINRQLVETRQITKHVANLLSHHLNEK---KEVG--EINIVLLKSALTSQFRKKF	DFYKVVREINDY	980
AKQ21048	911	LD	KAGFIKRQLVETRQITKHVAQVLLDSRMNTKYDENDKLIR--EVKVIITLKSCLVSDFRKDF	QFYKVVREINNY	981
WP_004636532	916	AD	RAHFLNRQLVETRQITKHVANLLDSQYNTAEQ-----R--INIVLLKSSMTRSFRKEF	KLYKVVREINDY	980
WP_002364836	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_016631044	873	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	938
EMS75795	655	ED	KARFIQRQLVETRQITKHVAAILDQYFN-QPEE-SK-NK--GIRIITLKSLLVSNFRKTF	GINKVVREINNH	722
WP_002373311	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_002378009	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_002407324	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_002413717	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_010775580	924	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	989
WP_010818269	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_010824395	922	ED	KAHFIQRQLVETRQITKNVAGILDQLYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_016622645	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_033624816	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_033625576	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_033789179	922	ED	KAHFIQRQLVETRQITKNVAGILDQRYNANSKE-----K--KVQIITLKSALTSQFRSIF	GLYKVVREINDY	987
WP_002310644	919	ED	KAGFIKRQLVETRQITKHVAGILHHRFN-KAEDTNEPIR--KVRITLKSALVSNFRNRF	GIYKVVREINEY	988
WP_002312694	920	ED	KAGFIKRQLVETRQITKHVAGILHHRFN-KAEDTNDPIR--KVRITLKSALVSNFRNRF	GIYKVVREINEY	989
WP_002314015	920	ED	KAGFIKRQLVETRQITKHVAGILHHRFN-KAEDTNEPIR--KVRITLKSALVSNFRNRF	GIYKVVREINEY	989
WP_002320716	920	ED	KAGFIKRQLVETRQITKHVAGILHHRFN-KAEDTNEPIR--KVRITLKSALVSNFRNRF	GIYKVVREINEY	989

WP_002330729	919	ED	KAGFIKRQLVETRQITKHVAGILHHRFN-KAEDTNEPIR--KVRITLTKSALVSQFRNRF	GIYKVRINEY	988
WP_002335161	920	ED	KAGFIKRQLVETRQITKHVAGILHHRFN-KAEDTNEPIR--KVRITLTKSALVSQFRNRF	GIYKVRINEY	989
WP_002345439	920	ED	KAGFIKRQLVETRQITKHVAGILHHRFN-KAEDTNEPIR--KVRITLTKSALVSQFRNRF	GIYKVRINEY	989
WP_034867970	911	ED	KARFIQRQLVETRQITKHVANIHQSFN-QEEEGTD-CD--GVQIITLTKATLTSQFRQTF	GLYKVRINEPH	979
WP_047937432	920	ED	KAGFIKRQLVETRQITKHVAGILHHRFN-KAEDTNEPIR--KVRITLTKSALVSQFRNRF	GIYKVRINEY	989
WP_010720994	911	ED	KARFIQRQLVETRQITKHVANIHQSFN-QEEEGTD-CD--GVQIITLTKATLTSQFRQTF	GLYKVRINEPH	979
WP_010737004	911	ED	KARFIQRQLVETRQITKHVANIHQSFN-QEEEGTD-CD--GVQIITLTKATLTSQFRQTF	GLYKVRINEPH	979
WP_034700478	911	ED	KARFIQRQLVETRQITKHVANIHQSFN-QEEEGTD-CD--GVQIITLTKATLTSQFRQTF	GLYKVRINEPH	979
WP_007209003	912	SD	KAGFLKRQLVETRQITKHVATILDSKFNE--DSNDRDVQ-----IITLKSALVSEFRKTF	NLYKVRINEIDL	977
WP_023519017	905	AD	KARFIQRQLVETRQITKHVANIHQSFN-LPEEVSA-TE--KTSIITLKSTLTSQFRQMF	DIYKVRINEHH	973
WP_010770040	915	DD	RAHFIKRQLVETRQITKHVARILDQRFNSQKDEEGKTIR--AVRVVTLKSSLTSQFRKQF	AHKVREINDY	985
WP_048604708	912	DD	KAGFIHRQLVETRQITKNVARIHQRFNSEKDEEGNLIR--KVRITLTKSALTSQFRKNY	GIYKIREINDY	982
WP_010750235	914	DD	KARFIQRQLVETRQITKHVAAILHQYFN-QTQELEK-EK--DIRIITLKSLLVSQFRQVF	GIHKVREINHH	982
AI116583	950	LD	KAGFIKRQLVETRQITKHVAQILDSRMTKYDENDKLIR--EVKVITLKSCLVSDFRKDF	QFYKVRINEINNY	1020
WP_029073316	928	KD	KERFINRQIVETRQITKHVAQIISNHYET-----TKVTVRADLISHAFRERY	HIYKNRDINDF	987
WP_031589969	928	KD	QERFINRQIVETRQITKHVAQIISNHYEN-----TKVTVRADLISHQFRERY	HIYKNRDINDF	987
KDA45870	903	KL	KERFIERQLVETRQITKYVAQLLDQRLN--YDNGVVELD-eKIAIVTLKAQLASQFRSEF	KLKVRALNLL	972
WP_039099354	924	-D	MKGFINRQLVETRQIKLATNLMEQYGED-----NIELITVKSGLTHQMRTEF	DFPKNRNLNH	990
AKP02966	926	KD	KLGFHRQLVQTSQMVKGVANILNSMYK--NQGTTCIQ-----ARANLSTAFRKAL	ELVKNRNINDF	999
WP_010991369	914	AD	KARFIHRQLVETRQITKNVANIHQRFNYEKDDHGNTMK--QVRIVTLKSALVSQFRKQF	QLYKVRDVNDY	984
WP_033838504	914	AD	KARFIHRQLVETRQITKNVANIHQRFNYEKDDHGNTMK--QVRIVTLKSALVSQFRKQF	QLYKVRDVNDY	984
EHN60060	917	AD	KARFIHRQLVETRQITKNVANIHQRFNYEKDDHGNTMK--QVRIVTLKSALVSQFRKQF	QLYKVRDVNDY	987
EFR89594	683	AD	KARFIHRQLVETRQITKNVANIHQRFNYEKDDHGNTMK--QVRIVTLKSALVSQFRKQF	QLYKVRGVNDY	753
WP_038409211	914	AD	KANFIQRQLVETRQITKNVANIYQRFNCKQDENGNEVE--QVRIVTLKSTLVSQFRKQF	QLYKVRVNDY	984
EFR95520	533	AD	KANFIQRQLVETRQITKNVANIYQRFNCKQDENGNEVE--QVRIVTLKSTLVSQFRKQF	QLYKVRVNDY	984
WP_003723650	914	AD	KARFIHRQLVETRQITKNVANIYQRFNCKQDENGNEVE--QVRIVTLKSTLVSQFRKQF	QLYKVRVNGY	984
WP_003727705	914	AD	KARFIHRQLVETRQITKNVANIYQRFNCKQDENGNEVE--QVRIVTLKSTLVSQFRKQF	QLYKVRVNDY	984
WP_003730785	914	AD	KARFIHRQLVETRQITKNVANIYQRFNCKQDENGNEVE--QVRIVTLKSTLVSQFRKQF	QLYKVRVNDY	984
WP_003733029	914	AD	KARFIHRQLVETRQITKNVANIYQRFNCKQDENGNEVE--QVRIVTLKSTLVSQFRKQF	QFYKVRVNDY	984
WP_003739838	914	AD	KATFIHRQLVETRQITKNVANIHQRFNNETDNHGNTME--QVRIVTLKSALVSQFRKQF	QLYKVRVNDY	984
WP_014601172	914	AD	KARFIHRQLVETRQITKNVANIHQRFNNETDNHGNTME--QVRIVTLKSALVSQFRKQF	QLYKVRVNDY	984
WP_023548323	914	AD	KARFIHRQLVETRQITKNVANIHQRFNKTDDNEDTME--PVRIVTLKSALVSQFRKQF	QLYKVRVNDY	984
WP_031665337	914	AD	KARFIHRQLVETRQITKNVANIHQRFNCKQDENGNEVE--QVRIVTLKSALVSQFRKQF	QLYKVRVNDY	984
WP_031669209	914	AD	KARFIHRQLVETRQITKNVANIHQRFNCKQDENGNEVE--QVRIVTLKSALVSQFRKQF	QFYKVRVNDY	984

WP_033920898	914	AD	KARFIHRQLVETRQITTKNVANILHQRFNFKTDDNEDTME--PVRIVTLKSALVSQFRKQF	QLYKVVREVN	984
AKI42028	917	AD	KARFIHRQLVETRQITTKNVANILHQRFNNETDNHGNTME--QVRIVTLKSALVSQFRKQF	QLYKVVREVN	987
AKI50529	917	AD	KARFIHRQLVETRQITTKNVANILHQRFNFKTDDNEDTME--PVRIVTLKSALVSQFRKQF	QLYKVVREVN	987
EFR83390	362	AD	KARFIHRQLVETRQITTKNVANILHQRFNNETDNHGNTME--QVRIVTLKSALVSQFRKQF	QLYKVVREVN	432
WP_046323366	914	AD	KARFIHRQLVETRQITTKNVANILHQRFNCKKDESGNVIE--QVRIVTLKAALVSQFRKQF	QLYKVVREVN	984
AKE81011	927	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSCLVSDFRKDF	QFYKVVREINN	997
CUO82355	921	RD	EERFINRQLVETRQITKNVTQIIEHDHYS-----TKVAAIRANLSHEFRVKN	HIYKNRDIND	980
WP_033162887	923	KD	KERFINRQLVETRQIKNVAVIINDHYTN-----TNIVTVRAELSHQFRERY	KIYKNRDIND	982
AGZ01981	944	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSCLVSDFRKDF	QFYKVVREINN	1014
AKA60242	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSCLVSDFRKDF	QFYKVVREINN	981
AKS40380	911	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSCLVSDFRKDF	QFYKVVREINN	981
4UN5_B	915	LD	KAGFIKRQLVETRQITKHVAQIILDSRMNTKYDENDKLIR--EVKVITLKSCLVSDFRKDF	QFYKVVREINN	985

WP_010922251	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_039695303	992	HHADAYLNAVVGTTALIKKYPKL-ASEFVYGEYKYYDI	S---SD-----	KATAK--YfFYSNLM-NFFKTKVK	1058
WP_045635197	984	HHADAYLNAVWAKAILKKYPKL-EPEFVYGEYQYDL	SkdpKEV---EK	ATEKY--F-FYSNLL-NFFKKEVH	1055
5AXW A	703	HHAEADALI-----IaNAFDIFKEWKKLDK	Nq-mFE-----EK	ETEQEYkEiFITPHQI.KHIKDFKD	771
WP_009880683	666	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDI	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	735
WP_010922251	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_011054416	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_011284745	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_011285506	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_011527619	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_012560673	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDI	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_014407541	981	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1050
WP_020905136	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_023080005	981	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1050
WP_023610282	981	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1050
WP_030125963	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_030126706	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_031488318	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_032460140	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDI	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_032461047	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDI	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_032462016	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_032462936	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_032464890	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_033888930	807	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	876
WP_038431314	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_038432938	981	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1050
WP_038434062	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
BAQ51233	893	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	962
KGE60162	157	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	226
KGE60856		-----	-----	-----	
WP_002989955	982	HHADAYLNAVVGTTALIKKYPKL-ESEFVYGDYKVVYDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_003030002	981	HHADAYLNAVGNALLKYPQL-EPEFVYGEYPKYN-	S---YR---sRK	SATEK--FlFYSNIL-RFFKKE--	1041
WP_003065552	992	HHADAYLNAVVGTTALIKKYPKL-ASEFVYGEYKYYDI	S---SD-----	KATAK--YfFYSNLM-NFFKRVIR	1058
WP_001040076	987	HHADAYLNAVWAKAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049

WP_001040078	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGLYRKK-	L---SKI---VR	ATRKM--F-FYSNLM-NMFKRVVR	1057
WP_001040080	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040081	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040083	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040085	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040087	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040088	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040089	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040090	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040091	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040092	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040094	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040095	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040096	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040097	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040098	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040099	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040100	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040104	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040105	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040106	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040107	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040108	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040109	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_001040110	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_015058523	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_017643650	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_017647151	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_017648376	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_017649527	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_017771611	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_017771984	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
CFQ25032	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
CFV16040	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049

KLJ37842	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
KLJ72361	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
KL20707	1001	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1063
KL242645	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_047207273	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_047209694	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_050198062	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_050201642	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_050204027	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_050881965	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_050886065	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
AHN30376	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
EA078426	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
CCW42055	987	HHADAYLNAVWAKAAILTKYPQL-EPEFVYGDYPKYN-	S---YKT---RK	ATEKL--F-FYSNIM-NFFKTKVT	1049
WP_003041502	981	HHADAYLNAVIGTALLKKYPKL-ASEFVYGEFKKYDV	S---DK---eIG	KATAK--YfFYSNLM-NFFKKEVK	1050
WP_037593752	982	HHADAYLNAVIGNALLKKYPQL-EPEFVYGEYPKYN-	S---YR---sRK	SATEK--FlFYSNIL-RFFKKE--	1042
WP_049516684	982	HHADAYLNAVIGNALLKKYPQL-EPEFVYGEYPKYN-	S---YR---sRK	SATEK--FlFYSNIL-RFFKKE--	1042
GAD46167	981	HHADAYLNAVIGNALLKKYPQL-EPEFVYGEYPKYN-	S---YR---sRK	SATEK--FlFYSNIL-RFFKKE--	1041
WP_018363470	990	HHADAYLNAVIGTALLKKYPKL-APEFVYGEYKKYDV	S---SDDhseMG	KATAK--YfFYSNLM-NFFKRVIR	1062
WP_003043819	991	HHADAYLNAVIGTALLKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKR--F-FYSNIM-NFFKTEVK	1060
WP_006269658	981	HHADAYLNAVIGNALLKKYPQL-EPEFVYGEYPKYN-	S---YR---sRK	SATEK--FlFYSNIL-RFFKKE--	1041
WP_048800889	981	HHADAYLNAVIGTALLKKYPKL-TSEFVYGEYKKYDV	S---DND---eIG	KATAK--YfFYSNLM-NFFKTEVK	1051
WP_012767106	981	HHADAYLNAVIGTALLKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKR--F-FYSNIM-NFFKTEIT	1050
WP_014612333	981	HHADAYLNAVIGTALLKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKR--F-FYSNIM-NFFKTEIT	1050
WP_015017095	981	HHADAYLNAVIGTALLKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKR--F-FYSNIM-NFFKTEIT	1050
WP_015057649	981	HHADAYLNAVIGTALLKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKR--F-FYSNIM-NFFKTEIT	1050
WP_048327215	981	HHADAYLNAVIGTALLKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKR--F-FYSNIM-NFFKTEIT	1050
WP_049519324	981	HHADAYLNAVIGTALLKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKR--F-FYSNIM-NFFKTEIT	1050
WP_012515931	981	HHADAYLNAVWAKAAILGKYPQL-APEFVYGDYPKYN-	S---FKEr---QK	ATQKM--L-FYSNIL-KFFKDQES	1043
WP_021320964	981	HHADAYLNAVWAKAAILGKYPQL-APEFVYGDYPKYN-	S---FKEr---QK	ATQKT--L-FYSNIL-KFFKDQES	1043
WP_037581760	981	HHADAYLNAVWAKAAILGKYPQL-APEFVYGDYPKYN-	S---FKEr---QK	ATQKT--L-FYSNIL-KFFKDQES	1043
WP_004232481	989	HHADAYLNAVIGTALLKKYPKL-APEFVYGEYKKYDV	S---SDNhselG	KATAK--YfFYSNLM-NFFKTEVK	1061
WP_009854540	990	HHADAYLNAVIGTALLKKYPKL-APEFVYGEYKKYDI	S---SD-----	KATAK--YfFYSNLM-NFFKTKVK	1056
WP_012962174	990	HHADAYLNAVIGTALLKKYPKL-APEFVYGEYKKYDI	S---GD-----	KATAK--YfFYSNLM-NFFKRVIR	1056

WP_039695303	992	HHAHDAYLNAVVGTTALLKKYPKL-ASEFVYGEYKYYDI	S---SD-----	KATAK--YfFYSNLM-NFFKTKVK	1058
WP_014334983	989	HHAHDAYLNAVVGTTALLKKYPKL-TPEFVYGEYKYYDV	S---SDDysemG	KATAK--YfFYSNLM-NFFKTEVK	1061
WP_003099269	982	HHAQDAYLNAVVGTTALLKKYPKL-EAEFVYGDYKHYDL	P---DSS1--GK	ATTRM--F-FYSNLM-NFFKKEIK	1051
AHY15608	982	HHAQDAYLNAVVGTTALLKKYPKL-EAEFVYGDYKHYDL	P---DSS1--GK	ATTRM--F-FYSNLM-NFFKKEIK	1051
AHY17476	982	HHAQDAYLNAVVGTTALLKKYPKL-EAEFVYGDYKHYDL	P---DSS1--GK	ATTRM--F-FYSNLM-NFFKKEIK	1051
ESR09100					
AGM98575	982	HHAQDAYLNAVVGTTALLKKYPKL-EAEFVYGDYKHYDL	P---DSS1--GK	ATTRM--F-FYSNLM-NFFKKEIK	1051
ALF27331	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_018372492	995	HHAHDAYLNAVVAQALLKVYKF-ERELVYGSYVKESI	---FS---RK	ATERM---rMYNNIL-KFISKD--	1055
WP_045618028	985	HHAHDAYLNAVVAKAILKKYPKL-EPEFVYGDYQKYDL	TkdpKEV---EK	ATEKY--F-FYSNLL-NFFKKEVH	1056
WP_045635197	984	HHAHDAYLNAVVAKAILKKYPKL-EPEFVYGEYQKYDL	SkdpKEV---EK	ATEKY--F-FYSNLL-NFFKKEVH	1055
WP_002263549	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002263887	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002264920	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002269043	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002269448	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002271977	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002272766	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HE---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002273241	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002275430	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HE---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002276448	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002277050	984	HHAHDAYLNAVVKALLVKYPKL-EPEFVYGEYPKYN-	S---YR---eRK	ATQKM--F-FYSNIM-NMFKSKVK	1046
WP_002277364	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002279025	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HE---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002279859	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002280230	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002281696	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002282247	984	HHAHDAYLNAVVKALLVKYPKL-EPEFVYGEYPKYN-	S---YR---eRK	ATQKM--F-FYSNIM-NMFKSKVK	1046
WP_002282906	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002283846	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002287255	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002288990	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002289641	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002290427	982	HHAHDAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041

WP_002295753	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002296423	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002304487	996	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKG--	1055
WP_002305844	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002307203	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002310390	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_002352408	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_012997688	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_014677909	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_019312892	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_019313659	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_019314093	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_019315370	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HE---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_019803776	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_019805234	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_024783594	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_024784288	984	HHADAYLNAVVKALLVKYPKL-EPEFVYGEYLKYN-	S---YR---eRK	ATQKM--F-FYSNIM-NMFKSKVK	1046
WP_024784666	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_024784894	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_024786433	984	HHADAYLNAVVKALLVKYPKL-EPEFVYGEYPKYN-	S---YR---eRK	ATQKM--F-FYSNIM-NMFKSKVK	1046
WP_049473442	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HE---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
WP_049474547	982	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1041
EMC03581	975	HHADAYLNAVIGKALLGVYPQL-EPEFVYGDYPHFH-	G---HK---eNK	ATAKK--F-FYSNIM-NFFKKD--	1034
WP_000428612	987	HHADAYLNAVWAKAILKKYPKL-EPEFVYDQKYDL	SkdpKEI---EK	ATEKY--F-FYSNLL-NFFKEEVH	1058
WP_000428613	985	HHADAYLNAVWAKAILKKYPKL-EPEFVYDQKYDL	SrnpKEV---EK	ATEKY--F-FYSNLL-NFFKEEVH	1056
WP_049523028	980	HHADAYLNAVWAKAILKKYPKL-EPEFVYDQKYDL	TkdpKEI---EK	ATEKY--F-FYSNLL-NFFKDKVY	1051
WP_003107102	951	HHADAYLNAVGTALLKKYPKL-EAEFVYDQKYDL	S---DTS1---GK	ATAKM--F-FYSNIM-NFFKKEVR	1020
WP_054279288	983	HHADAYLNAVGTALLKMPKL-ASEFVYDQKYDL	S---GKAs---GH	ATAKY--F-FYSNLM-NFFKSEVK	1052
WP_049531101	985	HHADAYLNAVWAKAILKKYPKL-EPEFVYDQKYDL	SrnpKEI---EK	ATEKY--F-FYSNLL-NFFKEEVH	1056
WP_049538452	985	HHADAYLNAVWAKAILKKYPKL-EPEFVYDQKYDL	SkdpKDI---EK	ATEKY--F-FYSNLL-NFFKEEVH	1056
WP_049549711	987	HHADAYLNAVWAKAILKKYPKL-EPEFVYDQKNDL	SkdpKDI---EK	ATEKY--F-FYSNLL-NFFKEEVH	1058
WP_007896501	988	HHADAYLNAVGTALLKKYPKL-EAEFVYDQKHFDL	S---DPS1---GK	ATAKV--F-FYSNIM-NFFKEELS	1057
EFR44625	940	HHADAYLNAVGTALLKKYPKL-EAEFVYDQKHFDL	S---DPS1---GK	ATAKV--F-FYSNIM-NFFKEELS	1009
WP_002897477	984	HHADAYLNAVWAKAILKKYPKL-EPEFVYDQKYDL	FkpsKEI---EK	ATEKY--F-FYSNLL-NFFKEEVL	1055

WP_002906454	984	HHADAYLNAVAKAILKKYPKL-EPEFVYGDYQKYDL	SkasNTI---DK	ATEKY--F-FYSNLL-NFFKKEKVR	1055
WP_009729476	985	HHADAYLNAVAKAILKKYPKL-EPEFVYGDYQKYDL	skdpKEI---EK	ATEKY--F-FYSNLL-NFFKKEEVH	1056
CQR24647	984	HHADAYLNAVAKALLIRYPKL-EPEFVYGEYPKYN-	S---YRE---RK	ATEKM--F-FYSNIM-NMFKTTIK	1046
WP_000066813	989	HHADAYLNAVAKAILKKYPKL-EPEFVYGDYQKYDL	SrepKEV---EK	ATQKY--F-FYSNLL-NFFKKEEVH	1060
WP_009754323	985	HHADAYLNAVAKAILKKYPKL-EPEFVYGDYQKYDL	skdpKEV---EK	ATEKY--F-FYSNLL-NFFKKEEVH	1056
WP_044674937	986	HHADAYLNAVATALLKKYPQL-APEFVYGDYPKYN-	S---YKS---RK	ATEKV--L-FYSNIM-NFFRRVLV	1048
WP_044676715	988	HHADAYLNAVATALLKKYPQL-APEFVYGDYPKYN-	S---YKS---RK	ATEKV--L-FYSNIM-NFFRRVLV	1050
WP_044680361	988	HHADAYLNAVATALLKKYPQL-APEFVYGDYPKYN-	S---YKS---RK	ATEKV--L-FYSNIM-NFFRRVLV	1050
WP_044681799	986	HHADAYLNAVATALLKKYPQL-APEFVYGDYPKYN-	S---YKS---RK	ATEKV--L-FYSNIM-NFFRRVLV	1048
WP_049533112	981	HHADAYLNAVIGTALLKKYPKL-ASEFVYGEFKKYDV	S---DK---eIG	KATAK--YFFYSNLM-NFFKKEVK	1050
WP_029090905	952	HHQDAFLVAFLGTNITSNPKI-EMEYLFKGQHYLN	-----Ev-GK	AAKPKftF-IVENLS-----	1007
WP_006506696	977	HHADAYIVALIGFMRDRYPNMhDSKAIVSEYMKMFR	-----NKNd--QK	-----g---FVINSM-NYPY-EV-	1038
AIT42264	982	HHADAYLNAVGTALIKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_034440723	981	HHADAYLNGVIALKLELYPYM-AKDLIYGKYSYHRK	G-----DK	ATQAK--Y-KMSNII-ERFSQDL-	1041
AKQ21048	982	HHADAYLNAVGTALIKKYPKL-ESEFVYGDYKVDV	S---EQEi---GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
WP_004636532	981	HHGDAYLNAVATIMKVYPNL-ESEFVYGYKKTSM	-----FKE---EK	ATARK--H-FYSNIT-KFFKKEKV	1042
WP_002364836	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQT	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	1047
WP_016631044	939	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQT	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	998
EMS75795	723	HHADAYLNGVAVIALKKYPKL-EPEFVYGNVTKFNL	-----AT---eNK	ATAKK--E-FYSNIL-RFFEKE--	782
WP_002373311	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQA	-----FKE---NK	ATAKT--I-IYTNLM-RFFTED--	1047
WP_002378009	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQT	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	1047
WP_002407324	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQT	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	1047
WP_002413717	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQT	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	1047
WP_010775580	990	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQA	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	1049
WP_010818269	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQT	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	1047
WP_010824395	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQT	-----FKE---NK	ATAKT--I-IYTNLM-RFFTED--	1047
WP_016622645	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQT	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	1047
WP_033624816	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQA	-----FKE---NK	AMAKA--I-IYTNLL-RFFTED--	1047
WP_033625576	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQA	-----FKE---NK	ATAKA--I-IYTNLM-RFFTEV--	1047
WP_033789179	988	HHGDAYLNCVATLTKVYPNL-APEFVYGEYPKFQA	-----FKE---NK	ATAKA--I-IYTNLL-RFFTED--	1047
WP_002310644	989	HHADAYLNGVAVIALKKYPQL-APEFVYGEYKFNNA	-----HK---aNK	ATVKK--E-FYSNIM-KFFESD--	1048
WP_002312694	990	HHADAYLNGVAVIALKKYPQL-APEFVYGEYKFNNA	-----HK---aNK	ATVKK--E-FYSNIM-KFFESD--	1049
WP_002314015	990	HHADAYLNGVAVIALKKYPQL-APEFVYGEYKFNNA	-----HK---aNK	ATVKK--E-FYSNIM-KFFESD--	1049
WP_002320716	990	HHADAYLNGVAVIALKKYPQL-APEFVYGEYKFNNA	-----HK---aNK	ATVKK--E-FYSNIM-KFFESD--	1049

WP_002330729	989	HHADAYLNGVWALALLLKKYPQL-APEFVYGEYLKFN	----HK---aNK	ATVKK--E-FYSNIM-KFFESD--	1048
WP_002335161	990	HHADAYLNGVWALALLLKKYPQL-APEFVYGEYLKFN	----HK---aNK	ATVKK--E-FYSNIM-KFFESD--	1049
WP_002345439	990	HHADAYLNGVWALALLLKKYPQL-APEFVYGEYLKFN	----HK---aNK	ATVKK--E-FYSNIM-KFFESD--	1049
WP_034867970	980	HHADAYLNGVFIANVLLKKYPKL-APEFVYKVKYSL	----AR---eNK	ATAKK--E-FYSNIM-KFFESD--	1039
WP_047937432	990	HHADAYLNGVIALALLLKKYPQL-APEFVYGEYLKFN	----HK---aNK	ATVKK--E-FYSNIM-KFFESD--	1049
WP_010720994	980	HHADAYLNGVFIANVLLKKYPKL-APEFVYKVKYSL	----AR---eNK	ATAKK--E-FYSNIM-KFFESD--	1039
WP_010737004	980	HHADAYLNGVFIANVLLKKYPKL-APEFVYKVKYSL	----AR---eNK	ATAKK--E-FYSNIM-KFFESD--	1039
WP_034700478	980	HHADAYLNGVFIANVLLKKYPKL-APEFVYKVKYSL	----AR---eNK	ATAKK--E-FYSNIM-KFFESD--	1039
WP_007209003	978	HHADAYLNAVVALSLLRVYPQL-KPEFVYGEYKNS-	----IHDq--NK	ATIKK--qFYSNIT-RYFASK--	1037
WP_023519017	974	HHADAYLNGVWAMTLLKKYPKL-APEFVYGSYIKGDI	----NQ---iNK	ATAKK--E-FYSNIM-KFFESE--	1033
WP_010770040	986	HHGDAYLNGVWANSLLRVYPQL-QPEFVYGDYPKFNA	----YKA--NK	ATAKK--Q-LYTNIM-KFFAED--	1045
WP_048604708	983	HHADAYLNGVWATALLKIYPQL-EPEFVYGEFHRFNA	----FKE--NK	ATAKK--Q-FYSNLM-EFSKSD--	1042
WP_010750235	983	HHADAYLNAVVALALLLKKYPRL-APEFVYGSFAKFHL	----VK---eNK	ATAKK--E-FYSNIM-KFFEKE--	1042
AI116583	1021	HHADAYLNAVVGVALIKKYPKL-ESEFVYGDYKVDV	S---EQEi--GK	ATAKY--F-FYSNIM-NFFKTEIT	1090
WP_029073316	988	HHADAYIATILGTYIGHRFESL-DAKYIYGEYQKIFR	----NKNk--DK	---KDg---FILNSM-RNLYADK-	1052
WP_031589969	988	HHADAYIATILGTYIGHRFESL-DAKYIYGEYQKIFR	----QKNk--GK	---NDg---FILNSM-RNIYADK-	1052
KDA45870	973	HHADAYLNAVVALIMAKYPEL-KPYFVYGEYKTK-	----FKG1--GK	ATAKN--tLYANVL-YFLKENEV	1034
WP_039099354	991	HHADAYLTAFLGTYLRRFPTD-EMLLMNGEYKFKAS-	----QQ---DK	---RN--F---NFL-NGLKKD--	1043
AKP02966	1000	HHADAYLASFLGTYLRRFPTD-EMLLMNGEYKFKAS-	----KELysKK	-SRKN-gF-IISPLV-----	1062
WP_010991369	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGDYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_033838504	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGDYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
EHN60060	988	HHADAYLNGVWANTLLKVYPQL-EPEFVYGDYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1047
EFR89594	754	HHADAYLNGVWANTLLKVYPQL-EPEFVYGDYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	813
WP_038409211	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGDYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-RFFAKE--	1044
EFR95520	604	HHADAYLNGVWANTLLKVYPQL-EPEFVYGDYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-RFFAKE--	663
WP_003723650	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_003727705	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_003730785	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_003733029	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_003739838	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_014601172	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_023548323	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_031665337	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
WP_031669209	985	HHADAYLNGVWANTLLKVYPQL-EPEFVYGEYHQFDW	----FKA--NK	ATAKK--Q-FYTNIM-LFFAQK--	1044

WP_033920898	985	HHAHDAYLNGVWANTLLKVPQL-EPEFVYGEYHQFDW	----	FKA---	NK	ATAKK--Q-FYTNIM-LFFAQK--	1044
AKI42028	988	HHAHDAYLNGVWANTLLKVPQL-EPEFVYGEYHQFDW	----	FKA---	NK	ATAKK--Q-FYTNIM-LFFGQK--	1047
AKI50529	988	HHAHDAYLNGVWANTLLKVPQL-EPEFVYGEYHQFDW	----	FKA---	NK	ATAKK--Q-FYTNIM-LFFAQK--	1047
EFR83390	433	HHAHDAYLNGVWANTLLKVPQL-EPEFVYGEYHQFDW	----	FKA---	NK	ATAKK--Q-FYTNIM-LFFAQK--	492
WP_046323366	985	HHAHDAYLNCVVANTLLKVPQL-EPEFVYGDYHQFDW	----	FKA---	NK	ATAKK--Q-FYTNIM-LFFAKK--	1044
AKE81011	998	HHAHDAYLNAVGTALIKKYPKL-ESEFVYGDYKVDV	S---	EQEi---	GK	ATAKY--F-FYSNIM-NFFKTEIT	1067
CUO82355	981	HHAHDAYIVALIGGFMRRDRYPNMhDSKAVYSEYMKMFR	----	NKnd--	QK	-----g---FVINSM-NYPY-EV-	1042
WP_033162887	983	HHAHDAYIACIVGQFMHQNFHL-DAKIIYGQYK-----	----	KNY--	KK	---NYg---FILNSM-NHLQSDI-	1042
AGZ01981	1015	HHAHDAYLNAVGTALIKKYPKL-ESEFVYGDYKVDV	S---	EQEi---	GK	ATAKY--F-FYSNIM-NFFKTEIT	1084
AKA60242	982	HHAHDAYLNAVGTALIKKYPKL-ESEFVYGDYKVDV	S---	EQEi---	GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
AKS40380	982	HHAHDAYLNAVGTALIKKYPKL-ESEFVYGDYKVDV	S---	EQEi---	GK	ATAKY--F-FYSNIM-NFFKTEIT	1051
4UN5_B	986	HHAHDAYLNAVGTALIKKYPKL-ESEFVYGDYKVDV	S---	EQEi---	GK	ATAKY--F-FYSNIM-NFFKTEIT	1055

WP_010922251	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKTEVQT	GGFSK	ESIL-PKR-	1114
WP_039695303	1059	YAD-GTVFERPIIE	T-NAD-GE-IAWNKGIDFEKVRKVL-S-YPQVNIIVKKEVETQT	GGFSK	ESIL-PKG-	1120
WP_045635197	1056	YAD-GTIVKRENIE	Y-SKDtGE-IAWNKEKDFAI IKKVL-S-LPQVNIIVKKEVQT	GGFSK	ESIL-PKG-	1118
5AXW A	772	YKYsHRVDDKKNRE	VNNLN-GL- --YDKDND--KLKLINKSPEKLLIMYHHDpQT	--YQK	KLIMeQYgd	852
WP_009880683	736	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	798
WP_010922251	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_011054416	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_011284745	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_011285506	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_011527619	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_012560673	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_014407541	1051	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1113
WP_020905136	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_023080005	1051	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1113
WP_023610282	1051	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1113
WP_030125963	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_030126706	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_031488318	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_032460140	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_032461047	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_032462016	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_032462936	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_032464890	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_033888930	877	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	939
WP_038431314	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_038432938	1051	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1113
WP_038434062	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
BAQ51233	963	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1025
KGE60162	227	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	289
KGE60856	1	-----IE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	52
WP_002989955	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVL-S-MPQVNIIVKKEVQT	GGFSK	ESIL-PKR-	1114
WP_003030002	1042	-----DIQ	T-NED-GE-IAWNKEKHIKILKVL-S-YPQVNIIVKKEEQT	GGFSK	ESIL-PKR-	1093
WP_003065552	1059	YSN-GKVIVRPVVE	Y-SKD-TediAWDKSNFRTI CKVL-S-YPQVNIIVKKEVETQT	GGFSK	ESIL-PKG-	1121
WP_001040076	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKHKHFATVRKVL-S-YPQVNIIVKKEIQT	GGFSK	ESIL-AHG-	1112

WP_001040078	1058	LAD-GSIVVRPVIE	TGRYM-GK-TAWDKKKHFATVRKVLVLS-YPQVNIIVKKTEIQ	GGFSK	ESIL-AHG-	1120
WP_001040080	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040081	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040083	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040085	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040087	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040088	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040089	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040090	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040091	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040092	1050	LAD-ETAAVKDDIE	VNNET-GE-IAWDKKKHFATVRKVLVLS-YPQVNIIVKKTEVQ	GGFSK	ESIL-AHS-	1112
WP_001040094	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040095	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEVQ	GGFSK	ESIL-AHG-	1112
WP_001040096	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEVQ	GGFSK	ESIL-AHG-	1112
WP_001040097	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040098	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040099	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040100	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040104	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040105	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040106	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040107	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040108	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040109	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_001040110	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_015058523	1050	LAD-ETAAVKDDIE	VNNET-GE-IAWDKKKHFATVRKVLVLS-YPQVNIIVKKTEVQ	GGFSK	ESIL-AHS-	1112
WP_017643650	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_017647151	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_017648376	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_017649527	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_017771611	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
WP_017771984	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
CFQ25032	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112
CFV16040	1050	LAD-GTAAVKDDIE	VNNDT-GE-IVWDKKKHFATVRKVLVLS-YPQNNIVKKTEIQ	GGFSK	ESIL-AHG-	1112

KLJ37842	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
KLJ72361	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
KLL20707	1064	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1126
KLL42645	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
WP_047207273	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
WP_047209694	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
WP_050198062	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
WP_050201642	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
WP_050204027	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
WP_050881965	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
WP_050886065	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
AHN30376	1050	LAD-ETVAVKDDIE	VNNET-GE-IAWKKKHFAFVRKVL-YPQNNIVKKTEVQT	GGFSK	ESIL-AHS-	1112
EA078426	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
CCW42055	1050	LAD-GTVVVKDDIE	VNNDT-GE-IVWDKKKHFAFVRKVL-YPQNNIVKKTEIQT	GGFSK	ESIL-AHG-	1112
WP_003041502	1051	FAD-GTVVVERPDIE	T-SED-GE-IAWKKKHFAFVRKVL-YPQNNIVKKTEVQT	HGLDR	PSPK-PKP-	1122
WP_037593752	1043	-----DIQ	T-NED-GE-IAWKKKHFAFVRKVL-YPQNNIVKKTEEQT	GGFSK	ESIL-PKG-	1094
WP_049516684	1043	-----DIQ	T-NED-GE-IAWKKKHFAFVRKVL-YPQNNIVKKTEEQT	GGFSK	ESIL-PKG-	1094
GAD46167	1042	-----DIQ	T-NED-GE-IAWKKKHFAFVRKVL-YPQNNIVKKTEEQT	GGFSK	ESIL-PKG-	1093
WP_018363470	1063	YSN-GKVIVRPAVE	Y-SKDtGE-IAWKKKHFAFVRKVL-YPQNNIVKKVEVQT	GGFSK	ESIL-PKG-	1125
WP_003043819	1061	LAN-GEIRKRPLIE	TNGET-GE-IAWKKKHFAFVRKVL-YPQNNIVKKVEVQT	GGFSK	ESIL-SKR-	1123
WP_006269658	1042	-----DIQ	T-NED-GE-IAWKKKHFAFVRKVL-YPQNNIVKKTEEQT	GGFSK	ESIL-PKG-	1093
WP_048800889	1052	FAD-GTVVVERPDIE	T-SED-GE-IAWKKKHFAFVRKVL-YPQNNIVKKVEKQT	GRFSK	ESIL-PKG-	1113
WP_012767106	1051	LAN-GEIRKRPLIE	TNEET-GE-IVWDKGRDFATVRKVL-MPQNNIVKKTEVQT	GALTN	ESIL-ARG-	1113
WP_014612333	1051	LAN-GEIRKRPLIE	TNEET-GE-IVWDKGRDFATVRKVL-MPQNNIVKKTEVQT	GALTN	ESIL-ARG-	1113
WP_015017095	1051	LAN-GEIRKRPLIE	TNEET-GE-IVWNKGRDFATVRKVL-MPQNNIVKKTEVQT	GALTN	ESIL-ARG-	1113
WP_015057649	1051	LAN-GEIRKRPLIE	TNEET-GE-IVWDKGRDFATVRKVL-MPQNNIVKKTEVQT	GALTN	ESIL-ARG-	1113
WP_048327215	1051	LAN-GEIRKRPLIE	TNEET-GE-IVWDKGRDFATVRKVL-MPQNNIVKKTEVQT	GALTN	ESIL-ARG-	1113
WP_049519324	1051	LAN-GEIRKRPLIE	TNEET-GE-IVWDKGRDFATVRKVL-MPQNNIVKKTEVQT	GALTN	ESIL-ARG-	1113
WP_012515931	1044	L-----H	VNSD--GE-EIWNANKHLPIIKNLV-IPQNNIVKKTEVQT	GGFYK	ESIL-SKG-	1094
WP_021320964	1044	L-----H	VNSD--GE-EIWNANKHLPIIKNLV-IPQNNIVKKTEVQT	GGFYK	ESIL-SKG-	1094
WP_037581760	1044	L-----H	VNSD--GE-EIWNANKHLPIIKNLV-IPQNNIVKKTEVQT	GGFYK	ESIL-SKG-	1094
WP_004232481	1062	YAD-GRVFERPDIE	T-NAD-GE-VVWNKQDFNIVRKL-YPQNNIVKKVEVQT	GGFSK	ESIL-PKG-	1123
WP_009854540	1057	YAD-GTVVFERPIIE	T-NAD-GE-IAWKKKHFAFVRKVL-YPQNNIVKKVEVQT	GGFSK	ESIL-PKG-	1118
WP_012962174	1057	YSN-GKVVVRPVIE	C-SKDtGE-IAWKKKHFAFVRKVL-YPQNNIVKKVEVQT	GGFSK	ESIL-PKG-	1119

WP_039695303	1059	YAD-GTVFERPIIE	T-NAD-GE-IAWNKQIDFEKVRKVL-S-YPQVNIIVKKVEEQT	GGFSK	ESIL-PKG-	1120
WP_014334983	1062	YAD-GRVFERPDIE	T-NAD-GE-VAVNKQKDFDIVRKLVS-YPQVNIIVKKVEAQ	GGFSK	ESIL-SKG-	1123
WP_003099269	1052	LAD-DTIFTRPQIE	VNTET-GE-IVWDKVKDMQTIKRVMS-YPQVNIIVMKTEVQT	GGFSK	ESIW-PKG-	1114
AHY15608	1052	LAD-DTIFTRPQIE	VNTET-GE-IVWDKVKDMQTIKRVMS-YPQVNIIVMKTEVQT	GGFSK	ESIW-PKG-	1114
AHY17476	1052	LAD-DTIFTRPQIE	VNTET-GE-IVWDKVKDMQTIKRVMS-YPQVNIIVMKTEVQT	GGFSK	ESIW-PKG-	1114
ESR09100						
AGM98575	1052	LAD-DTIFTRPQIE	VNTET-GE-IVWDKVKDMQTIKRVMS-YPQVNIIVMKTEVQT	GGFSK	ESIW-PKG-	1114
ALF27331	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_018372492	1056	--K-----DVR	--DQEtGE-IVWDKKEIENIVKKVIY-SSPVNIIVKKREEQS	GALFK	QSNM-AVGY	1108
WP_045618028	1057	YAD-GTIVKRENIE	Y-SKDtGE-IAWNKEKDFATIKKVL-S-LPQVNIIVKKTEEQ	GGLFD	NNIV-SKKK	1124
WP_045635197	1056	YAD-GTIVKRENIE	Y-SKDtGE-IAWNKEKDFATIKKVL-S-LPQVNIIVKKREVT	GGFSK	ESIL-PKG-	1118
WP_002263549	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002263887	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002264920	1042		T-DKN-GE-IIWKKDEYISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002269043	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002269448	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002271977	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002272766	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002273241	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002275430	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002276448	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002277050	1047	LAD-DQIVERPMIE	VNDET-GE-IAWDKTKHITTVKKVLS-YPQVNIIVKKVEEQ	GGLFD	-----PKS-	1111
WP_002277364	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002279025	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002279859	1042		T-DKN-GE-IIWKKDEYISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002280230	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002281696	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002282247	1047	LAD-DQIVERPMIE	VNDET-GE-IAWDKTKHITTVKKVLS-YPQVNIIVKKVEEQ	GGLFD	-----PKS-	1111
WP_002282906	1042		I-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002283846	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002287255	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002288990	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002289641	1042		T-DKN-GE-IIWKKDEHISNIKKVLS-YPQVNIIVKKVEEQ	GGFSK	ESIL-PKG-	1093
WP_002290427	1042		T-DKN-GE-IIWKKDEYISNIKKVLS-YPQVNIIVKKVEEQ	GGFFK	ESIL-PKG-	1093

WP_002295753	1042	-----DVR	T-DKN-GE-I IWKKDEYI SNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_002296423	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_002304487	1056	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1107
WP_002305844	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_002307203	1042	-----DVR	T-DKN-GE-I IWKKDEYI SNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_002310390	1042	-----DVR	T-DKN-GE-I IWKKDEYI SNIKKVLS-YPQVNI VKKVEEQT	GGFFK	ESIL-PKG-	1093
WP_002352408	1042	-----DVR	T-DKN-GE-I IWKKDEYI SNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_012997688	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_014677909	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_019312892	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_019313659	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_019314093	1042	-----DVR	T-DKN-GE-I IWKKDEYI SNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_019315370	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_019803776	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_019805234	1042	-----DVR	T-DKN-GE-I IWKKDEYI SNIKKVLS-YPQVNI VKKVEEQT	GGFFK	ESIL-PKG-	1093
WP_024783594	1042	-----DVR	T-DKN-GE-I IWKKDEYI SNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_024784288	1047	LAD-DQIVERPMIE	VNDET-GE-I AWDKTKHITTVKKVLS-YPQVNI VKKVEEQT	GGLFD	-----PKS-	1111
WP_024784666	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_024784894	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_024786433	1047	LAD-DQIVERPMIE	VNDET-GE-I AWDKTKHITTVKKVLS-YPQVNI VKKVEEQT	GGLFD	-----PKS-	1111
WP_049473442	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
WP_049474547	1042	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1093
EMC03581	1035	-----DVR	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1086
WP_000428612	1059	YAD-GTIVKRENIE	T-DKN-GE-I IWKKDEHISNIKKVLS-YPQVNI VKKVEEQT	GGFSK	ESIL-PKG-	1121
WP_000428613	1057	YAD-GTIVKRENIE	Y-SKDtGE-I AWNKEKDFATIKKVLS-LPQVNI VKKREVQT	GGFSK	ESIL-PKG-	1119
WP_049523028	1052	YAD-GTIIQRGNVE	Y-SKDtGE-I AWNKRDFAI VRKVLS-YPQVNI VKKTEEQT	GGFSK	ESIL-PKG-	1114
WP_003107102	1021	LAD-GTIVTRPQIE	TNTEt-GE-I VWDKVKDIKTIRKVLS-IPQIN VVKTEVQT	GGFSK	ESIL-SKR-	1083
WP_054279288	1053	LAN-GNIIKRSPIE	VNEET-GE-I VWDKTKDFGTVRKLVS-APQVNI VKKTEIQT	GGFSN	ETIL-SKG-	1115
WP_049531101	1057	YAD-GTIVKRENIE	Y-SKDtGE-I AWNKEIDFATIRKILS-LSQVNI VKKTEEQT	GGLFD	NNIV-SKKk	1124
WP_049538452	1057	YAD-GTIVKRENIE	Y-SKDtGE-I AWNKEKDFATIKKILS-LPQVNI VKKTEEQT	GGLFD	NNIV-SKKk	1124
WP_049549711	1059	YAD-GTIVKRENIE	Y-SKDtGE-I AWNKEKDFATIKKVLS-YPQVNI VKKTEEQT	GGLFD	NNIV-SKEk	1126
WP_007896501	1058	LAD-GTLMKRPVIE	TNTEt-GE-VVWDKVKDFKTIRKVLS-YPQVNI VKKTEIQS	GAFSK	ESVL-SKG-	1120
EFR44625	1010	LAD-GTLMKRPVIE	TNTEt-GE-VVWDKVKDFKTIRKVLS-YPQVNI VKKTEIQS	GAFSK	ESVL-SKG-	1072
WP_002897477	1056	YAD-GTIRKRENIE	Y-SKDtGE-I AWDKTKDFATIKKVLS-YPQVNI VKKREVQT	GGFSK	ESIL-PKG-	1118

WP_002906454	1056	YAD-GTIKKRENIE	Y-SNDtGE-IAWNKEKDFATIKKVLs-LPQVNIvKKTEEQT	GGLFD	NNIV-SKkK	1123
WP_009729476	1057	YAD-GTIvKRENIE	Y-SKDtGE-IAWNKEKDFATIKKVLs-LPQVNIvKKREVQOT	GGFSK	ESIL-PKG-	1119
CQR24647	1047	LAD-GRVVEKPVIE	ANEET-GE-IAWDKTKHFANvKKVLs-YPQVSIvKKVEEQOT	GGFSK	ESIL-PKG-	1109
WP_000066813	1061	YAD-GTIvKRENIE	Y-SKDtGE-IAWNKEKDFATIKKVLs-LPQVNIvKKTEVQOT	GGFSK	ESIL-PKG-	1123
WP_009754323	1057	YAD-GTIvKRENIE	Y-SKDtGE-IAWNKEKDFATIKKVLs-LPQVNIvKKREVQOT	GGFSK	ESIL-PKG-	1119
WP_044674937	1049	YSKtGEVRIRPVIE	VNKET-GE-IVWDKKSDFTRvKVLs-YPQVNIvKKVEMQOT	GGFSK	ESIL-QHG-	1112
WP_044676715	1051	YSKtGEVRIRPVIE	VNKET-GE-IVWDKKSDFTRvKVLs-YPQVNIvKKVEMQOT	GGFSK	ESIL-QHG-	1114
WP_044680361	1051	YSKtGEVRIRPVIE	VNKET-GE-IVWDKKSDFTRvKVLs-YPQVNIvKKVEMQOT	GGFSK	ESIL-QHG-	1114
WP_044681799	1049	YSKtGEVRIRPVIE	VNKET-GE-IVWDKKSDFTRvKVLs-YPQVNIvKKVEMQOT	GGFSK	ESIL-QHG-	1112
WP_049533112	1051	FAD-GTIVVERPDIE	T-SED-GE-IAWNkQDFKIVRkVLs-YPQVNIvKKTEVQOT	HGLDR	PSPK-PKP-	1122
WP_029090905	1008	-KQ-----Q	--NSTtGE-VKWNPEVDIAKLKRILN-FKQCNIVRkVEEQS	GALFK	ETIY-PVEe	1061
WP_006506696	1039	--D-----	-----GK-LIWNP-DLINEIKKCFY-YKDCYCTTKLDQKS	GQLFN	-TVL-SNDa	1084
AIT42264	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATvRkVLs-MPQVNIvKKTEVQOT	GGFSK	ESIL-PKR-	1114
WP_034440723	1042	-----LA	--NPD-GE-IAWEKDKDLNTIRKVLs-SKQINIiKKAEEGK	GRLFK	ETIN-SRPs	1092
AKQ21048	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATvRkVLs-MPQVNIvKKTEVQOT	GGFSK	ESIL-PKR-	1114
WP_004636532	1043	-----	VNEET-GE-ILWdTERHLSTIKRVLs-WKQMIvKKVEKQK	GQLWK	ETIY-PKG-	1092
WP_002364836	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_016631044	999	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1049
EMS75795	783	--E-----Y	SYDEN-GE-IFWDKARHIPOIKKVIS-SHQVNIvKKVEVQOT	GGFYK	ETVN-PKG-	834
WP_002373311	1048	--E-----P	RFTKD-SE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_002378009	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_002407324	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_002413717	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_010775580	1050	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1100
WP_010818269	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_010824395	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_016622645	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_033624816	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_033625576	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_033789179	1048	--E-----P	RFTKD-GE-ILWSN-SYLTKIKKELN-YHQMIvKKVEVQK	GGFSK	ESIK-PKG-	1098
WP_002310644	1049	--T-----P	VCDEN-GE-IFWDKSKSIAQVKKVIN-HHHMNIvKKTEIQK	GGFSK	ETVE-PKK-	1100
WP_002312694	1050	--T-----P	VCDEN-GE-IFWDKSKSIAQVKKVIN-HHHMNIvKKTEIQK	GGFSK	ETVE-PKK-	1101
WP_002314015	1050	--T-----P	VCDEN-GE-IFWDKSKSIAQVKKVIN-HHHMNIvKKTEIQK	GGFSK	ETVE-PKK-	1101
WP_002320716	1050	--T-----P	VCDEN-GE-IFWDKSKSIAQVKKVIN-HHHMNIvKKTEIQK	GGFSK	ETVE-PKK-	1101

WP_002330729	1049	--T-----P	VCDEN-GE-IFWDKSKSIAQVKKVIN-HHHMNIIVKKTEIQK	GGFSE	ETVE-PKK-	1100
WP_002335161	1050	--T-----P	VCDEN-GE-IFWDKSKSIAQVKKVIN-HHHMNIIVKKTEIQK	GGFSK	ETVE-PKK-	1101
WP_002345439	1050	--T-----P	VCDEN-GE-IFWDKSKSIAQVKKVIN-HHHMNIIVKKTEIQK	GGFSK	ETVE-PKK-	1101
WP_034867970	1040	--E-----P	FCDEN-GE-IYWEKSHHLPRIKKVLS-SHQVNVVKKVEQQK	GGFYK	ETVN-SKE-	1091
WP_047937432	1050	--T-----P	VCDEN-GE-IFWDKSKSIAQVKKVIN-HHHMNIIVKKTEIQK	GGFSK	ETVE-PKK-	1101
WP_010720994	1040	--E-----P	FCDEN-GE-IYWEKSHHLPRIKKVLS-SHQVNVVKKVEQQK	GGFYK	ETVN-SKE-	1091
WP_010737004	1040	--E-----P	FCDEN-GE-IYWEKSHHLPRIKKVLS-SHQVNVVKKVEQQK	GGFYK	ETVN-SKE-	1091
WP_034700478	1040	--E-----P	FCDEN-GE-IYWEKSHHLPRIKKVLS-SHQVNVVKKVEQQK	GGFYK	ETVN-SKE-	1091
WP_007209003	1038	--D-----	IINDD-GE-ILWNKQETIAQVIKTLG-MHQVNVVKKVEIQK	GGFSK	ESIQ-PKG-	1089
WP_023519017	1034	--E-----I	ICDEQ-GE-VIWNKKRDLSTIKKTIG-AHQVNIIVKKVEKQK	GGFYK	ETIN-SKA-	1085
WP_010770040	1046	--A-----V	IIDEN-GE-ILWDK-KNIATVKKVMS-YPQMNIVKKPEIQT	GSFSK	ETIK-PKG-	1096
WP_048604708	1043	--K-----V	IIDEN-GE-ILWNQ-KKIIVTVKKVMN-YRQMNIVKKVEIQK	GGFSK	ESIL-PKG-	1093
WP_010750235	1043	--E-----Q	FCDEN-GE-IFWDRKRHIQIKKVIS-SHQVNIIVKKVEVQT	GSFYK	ETVN-TKE-	1094
AI116583	1091	LAN-GEIRKRLIE	TNGET-GE-IVWDKGRDFATVRKVLV-MPQVNIIVKKTEVQT	GGFSK	ESIL-PKR-	1153
WP_029073316	1053	--D-----	----T-GE-VVWDP-EWISRIKKCFY-YKDCFVTKKLEENN	GSFFN	-TVR-PNDe	1099
WP_031589969	1053	--D-----	----T-GE-IVWDP-NYIDRIKKCFY-YKDCFVTKKLEENN	GTFFN	-TVL-PNDt	1099
KDA45870	1035	YPF-----	-----WDKARDLPTIKRILY-RAQVNVKVRKAERQT	GGFSD	EMLV-PKS-	1078
WP_039099354	1044	-----E	LVDEN-TEaViWNKESGLAYLNKIYQ-FKKILVTRVHENS	GALFN	QTLYaAKDd	1097
AKP02966	1063	--N-----GTTQ	--DRNtGE-IIWNVG-FRDKILKIFN-YHQCNVTRKTEIKT	GQFYD	QTIYsPKNp	1118
WP_010991369	1045	--D-----R	IIDEN-GE-ILWDK-KYLDTVKKVMS-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1095
WP_033838504	1045	--D-----R	IIDEN-GE-ILWDK-KYLDTVKKVMS-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1095
EHN60060	1048	--D-----R	IIDEN-GE-ILWDK-KYLDTVKKVMS-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1098
EFR89594	814	--D-----R	IIDEN-GE-ILWDK-KYLDTVKKVMS-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	864
WP_038409211	1045	--N-----Q	IIDKN-GE-ILWDN-RYLDTIKKVLS-YRQMNIVKKTEIQK	GEFSN	ATVN-PKG-	1095
EFR95520	664	--N-----Q	IIDKN-GE-ILWDN-RYLDTIKKVLS-YRQMNIVKKTEIQK	GEFSN	ATVN-PKG-	714
WP_003723650	1045	--E-----R	IIDEN-GE-ILWDK-KYLETIKKVLd-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1095
WP_003727705	1045	--E-----R	IIDEN-GE-ILWDK-KYLETIKKVLd-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1095
WP_003730785	1045	--E-----R	IIDEN-GE-ILWDK-KYLETIKKVLd-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1095
WP_003733029	1045	--D-----R	IIDEN-GE-ILWDK-RYLETVKKVLG-YRQMNIVKKTEIQK	GEFSN	VTPN-PKG-	1095
WP_003739838	1045	--E-----R	IIDEN-GE-ILWDK-KYLETIKKVLd-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1095
WP_014601172	1045	--E-----R	IIDEN-GE-ILWDK-KYLETIKKVLd-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1095
WP_023548323	1045	--E-----R	IIDEN-GE-ILWDK-KYLETIKKVLN-YRQMNIVKKTEIQK	GEFSN	QNPk-PRG-	1095
WP_031665337	1045	--E-----R	IIDEN-GE-ILWDK-KYLETIKKVLd-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1095
WP_031669209	1045	--D-----R	IIDEN-GE-ILWDK-RYLETVKKVLG-YRQMNIVKKTEIQK	GEFSN	VTPN-PKG-	1095

WP_033920898	1045	--E-----R	I IDEN-GE-ILWDK-KYLETIKKVLN-YRQMNIVKKTEIQK	GEFSN	QNPk-PRG-	1095
AKI42028	1048	--E-----R	I IDEN-GE-ILWDK-KYLETIKKVLD-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	1098
AKI50529	1048	--E-----R	I IDEN-GE-ILWDK-KYLETIKKVLN-YRQMNIVKKTEIQK	GEFSN	QNPk-PRG-	1098
EFR83390	493	--E-----R	I IDEN-GE-ILWDK-KYLETIKKVLD-YRQMNIVKKTEIQK	GEFSK	ATIK-PKG-	543
WP_046323366	1045	--D-----R	I IDEN-GE-ILWDK-KYLDTIKKVLN-YRQMNIVKKTEIQK	GEFSN	ATAN-PKG-	1095
AKE81011	1068	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVLV-MPQVNIKKTEVQT	GGFSK	ESIL-PKR-	1130
CUO82355	1043	--D-----	-----GK-LIWNP-DLINEIKKCFY-YKDCYCTTKLDQKS	GQMFN	-TVL-PNDa	1088
WP_033162887	1043	--D-----	----T-GE-VMWDP-AKIGIKKSCFY-YKDVYVTKKLEQNS	GTLFN	-TVL-PNDa	1089
AGZ01981	1085	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVLV-MPQVNIKKTEVQT	GGFSK	ESIL-PKR-	1147
AKA60242	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVLV-MPQVNIKKTEVQT	GGFSK	ESIL-PKR-	1114
AKS40380	1052	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVLV-MPQVNIKKTEVQT	GGFSK	ESIL-PKR-	1114
4UN5_B	1056	LAN-GEIRKRPLIE	TNGET-GE-IVWDKGRDFATVRKVLV-MPQVNIKKTEVQT	GGFSK	ESIL-PKR-	1118

WP_010922251	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_039695303	1121	--DSD	KLI PRKTkKV-YW-DTKKYGGFDSPTVAYSV-FVWAD--VE--	KGKAKKLKTVKELAVGISIME	RSFFEE	1185	
WP_045635197	1119	--NSD	KLI PRKT-KDILL-DTKKYGGFDSPTVAYSV-LLIAD--IE--	KGKAKKLKTVKTLVGITIME	KAAFEE	1183	
5AXW_A	853	--EKN	-LYKYEeTGNYL--TKYSKKDNGPVIKKI-----	KYGNKLNNAHLDITDDYPNS	-VKLSL	912	
WP_009880683	799	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELVGITIME	RSSFEEK	860
WP_010922251	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_011054416	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_011284745	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_011285506	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_011527619	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_012560673	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_014407541	1114	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1175
WP_020905136	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_023080005	1114	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1175
WP_023610282	1114	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1175
WP_030125963	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_030126706	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_031488318	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_032460140	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_032461047	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_032462016	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_032462936	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_032464890	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_033888930	940	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1001
WP_038431314	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_038432938	1114	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1175
WP_038434062	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
BAQ51233	1026	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1087
KGE60162	290	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	351
KGE60856	53	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	114
WP_002989955	1115	--NSD	KLIA----	RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--	KGKSKKLKSVKELLGITIME	RSSFEEK	1176
WP_003030002	1094	--ESD	KLI PRKT-KNSYW-NPKKYGGFDSPTVAYSV-LVFAD--VE--	KGKSKKLKSVKELLGITIME	KKRFEK	1158	
WP_003065552	1122	--DSD	KLI PRKTkKA-YW-DTKKYGGFDSPTVAYSV-FVWAD--VE--	KGKAKKLKTVKELAVGISIME	RSFFEE	1186	
WP_001040076	1113	--NSD	KLI PRKT-KDIYL-DPKKYGGFDSPTVAYSV-LVWAD--IK--	KGKAQKLKTVTELLGITIME	RSRFEK	1177	

WP_001040078	1121	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSKF EK	1185
WP_001040080	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040081	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040083	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040085	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040087	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040088	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040089	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
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WP_001040091	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040092	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RERF EK	1177
WP_001040094	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040095	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040096	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040097	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040098	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040099	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040100	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040101	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040105	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_001040106	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RFRF EK	1177
WP_001040107	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RFRF EK	1177
WP_001040108	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RFRF EK	1177
WP_001040109	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RFRF EK	1177
WP_001040110	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RFRF EK	1177
WP_015058523	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RERF EK	1177
WP_017643650	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_017647151	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_017648376	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_017649527	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
WP_017771611	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RFRF EK	1177
WP_017771984	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
CFQ25032	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177
CFV16040	1113	--NSD	KLI PRKT-KDIYL-DPKKYGDFDSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRF EK	1177

KLJ37842	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
KLJ72361	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
KL20707	1127	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1191
KL42645	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RFRFEK	1177
WP_047207273	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
WP_047209694	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
WP_050198062	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
WP_050201642	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
WP_050204027	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RFRFEK	1177
WP_050881965	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
WP_050886065	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
AHN30376	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RERFEK	1177
EA078426	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
CCW42055	1113	--NSD	KLIPRKT-KDIYL-DPKKYGDFSPIVAYSV-LVAD--IK--KGKQKLTVTTELLGITIME	RSRFEK	1177
WP_003041502	1123	--DSS	ENLVGK-RNL--DPKKYGGYAGISNYSYV-LVKAI--IE--KGVKKETMVEFQGISILD	RITFEK	1185
WP_037593752	1095	--ESD	KLIPRKT-KNSYW-NPKKYGDFSPVAYSI-LVFAD--VE--KGSKKLRKQDMVGITIME	KKRFK	1159
WP_049516684	1095	--ESD	KLIPRKT-KNSYW-NPKKYGDFSPVAYSI-LVFAD--VE--KGSKKLRKQDMVGITIME	KKRFK	1159
GAD46167	1094	--ESD	KLIPRKT-KNSYW-NPKKYGDFSPVAYSI-LVFAD--VE--KGSKKLRKQDMVGITIME	KKRFK	1158
WP_018363470	1126	--DSD	KLIPRKT-KV-LW-EPKKYGGFSPVAYSV-LVAD--VE--KGTKKLTVKELVGSIME	RSFFEK	1190
WP_003043819	1124	--ESA	KLIP---RKKGW-DTRKYGGFSPVAYSI-LVAK--VE--KGAKKLKSVKLVGITIME	KGSYK	1185
WP_006269658	1094	--ESD	KLIPRKT-KNSYW-DPKKYGDFSPVAYSI-LVFAD--VE--KGSKKLRKQDMVGITIME	KKRFK	1158
WP_048800889	1114	--DSD	KLIPRKT-KNSYW-DPKKYGDFSPVAYSI-LVFAD--VE--KGSKKLRKQDMVGITIME	RPFFEK	1178
WP_012767106	1114	--SFD	KLIPRKT-KNSYW-DPKKYGDFSPVAYSV-LVAD--IK--KGKQKLTVKELVGSIME	KLVFEK	1177
WP_014612333	1114	--SFD	KLIS----RKHRF-ESSKYGGFSPVAYSV-LVAKskVQ--DGKVKIKITGKELIGITLLD	KLVFEK	1177
WP_015017095	1114	--SFD	KLIS----RKHRF-ESSKYGGFSPVAYSV-LVAKskVQ--DGKVKIKITGKELIGITLLD	KLVFEK	1177
WP_015057649	1114	--SFD	KLIS----RKHRF-ESSKYGGFSPVAYSV-LVAKskVQ--DGKVKIKITGKELIGITLLD	KLVFEK	1177
WP_048327215	1114	--SFD	KLIS----RKHRF-ESSKYGGFSPVAYSV-LVAKskVQ--DGKVKIKITGKELIGITLLD	KLVFEK	1177
WP_049519324	1114	--SFD	KLIS----RKHRF-ESSKYGGFSPVAYSV-LVAKskVQ--DGKVKIKITGKELIGITLLD	KLVFEK	1177
WP_012515931	1095	--NSD	KLIP----RKNW-DTRKYGGFSPVAYSV-LVIK--ME--KGAKVLKPKVEMVGITIME	RTAFEE	1156
WP_021320964	1095	--NSD	KLIP----RKNW-DTRKYGGFSPVAYSV-LVIK--ME--KGAKVLKPKVEMVGITIME	RTAFEE	1156
WP_037581760	1095	--NSD	KLIP----RKNW-DTRKYGGFSPVAYSV-LVIK--ME--KGAKVLKPKVEMVGITIME	RTAFEE	1156
WP_004232481	1124	--DSD	KLIPRKT-KL-QW-ETQKYGGFSPVAYSV-LVAD--VE--KGTTRKLTVKELVGSIME	RSSFEE	1188
WP_009854540	1119	--DSD	KLIPRKT-KV-YW-DTKKYGGFSPVAYSV-FVAD--VE--KGAKKLKTVKELVGSIME	RSFFEK	1183
WP_012962174	1120	--NSD	KLIPRKT-KF-RW-DTPKYGGFSPVAYSV-FVIAD--VE--KGAKKLKTVKELVGSIME	RSSFEE	1184

WP_039695303	1121	--DSD	KLIPRKTkKV-YW-DTKKYGDFDSPITVAYSV-FVWAD--VE--KGKAKKLTWKELVGVISIME	RSFFEE	1185
WP_014334983	1124	--DSD	KLIPRKTkKV-YW-DTKKYGDFDSPITVAYSV-LVWAD--IE--KGKAKKLTWKELVGVISIME	RSFFEE	1188
WP_003099269	1115	--DSD	KLIA----RKKS-DPKKYGDFDSPITVAYSV-LVWAK--IA--KGKTQKLTIKELVGVISIME	QDEFEK	1176
AHY15608	1115	--DSD	KLIA----RKKS-DPKKYGDFDSPITVAYSV-LVWAK--IA--KGKTQKLTIKELVGVISIME	QDEFEK	1176
AHY17476	1115	--DSD	KLIA----RKKS-DPKKYGDFDSPITVAYSV-LVWAK--IA--KGKTQKLTIKELVGVISIME	QDEFEK	1176
ESR09100	1	----	-----ME	QDEFEK	8
AGM98575	1115	--DSD	KLIA----RKKS-DPKKYGDFDSPITVAYSV-LVWAK--IA--KGKTQKLTIKELVGVISIME	QDEFEK	1176
ALF27331	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAN--IE--KGKSKLKTVDLVTGTIME	RTIFEK	1158
WP_018372492	1109	--NN	KLIP----RKKDW-SVDKYGDFDIEPAESYSLaIFYTD--IN----GKPKKSTIIASRME	KKDYEK	1167
WP_045618028	1125	vvdAS	KLTPIKS-G--L-SPEKYGGYARPTIAYSV-LVIAD--IE--KGKAKKLTWKELVGVITVD	KKKFEA	1188
WP_045635197	1119	--NSD	KLIPRKT-KDILL-DTKKYGDFDSPITVAYSV-LLIAD--IE--KGKAKKLTWKELVGVITIME	KAAFEF	1183
WP_002263549	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002263887	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002264920	1094	--DSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002269043	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002269448	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002271977	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002272766	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002273241	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002275430	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002276448	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002277050	1112	--PLE	KLVLKK---AL-NPEKYGGYQKPTTAYPI-LLIVD-----TKQLIPISVMD	KKRFEQ	1166
WP_002277364	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002279025	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002279859	1094	--DSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002280230	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002281696	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002282247	1112	--PLE	KLVLKK---AL-NPEKYGGYQKPTTAYPI-LLIVD-----TKQLIPISVMD	KKRFEQ	1166
WP_002282906	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002283846	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002287255	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002288990	1094	--NSY	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002289641	1094	--NSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002290427	1094	--DSD	KLIPRKT-KKFW-DTKKYGDFDSPITVAYSV-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158

WP_002295753	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002296423	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002304487	1108	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1172
WP_002305844	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002307203	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002310390	1094	--DSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_002352408	1094	--DSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_012997688	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_014677909	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_019312892	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_019313659	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_019314093	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_019315370	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_019803776	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_019805234	1094	--DSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_024783594	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KSKSKLKTVKALVGVTIME	KMTFER	1158
WP_024784288	1112	--PLE	KLVLKK---AL-NPEKYGQKPTTAYPI-LLIVD-----TKQLIPISVMD	KRFEQ	1166
WP_024784666	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_024784894	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_024786433	1112	--PLE	KLVLKK---AL-NPEKYGQKPTTAYPI-LLIVD-----TKQLIPISVMD	KRFEQ	1166
WP_049473442	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
WP_049474547	1094	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1158
EMC03581	1087	--NSD	KLI PRKT-KKFW-DTKKYGDFSPIVAYSI-LVIAD--IE--KGKSKLKTVKALVGVTIME	KMTFER	1151
WP_000428612	1122	--NSD	KLI PRKT-KDILW-DTKKYGDFSPIVAYSI-LLIAD--IE--KGKAKRLKTVKTLVGVTIME	KATFEK	1186
WP_000428613	1120	--NSD	KLI PRKT-KDILW-ETTKYGGDFSPVIAYSI-LLIAD--IE--KGKAKLKTVKTLVGVTIME	KAAFEE	1184
WP_049523028	1115	--NSD	KLI PRKT-KNVQL-DTKKYGDFSPVIAYSI-LLVAD--VE--KGKSKLKTVKSLIGITIME	KVKFEA	1179
WP_003107102	1084	--DSD	KLI P----RKNW-DPKKYGFGSPIIAYSV-LVAK--VT--KGKSKTCSVKELVGITIME	QNEFEK	1145
WP_054279288	1116	--KSS	KLI P----RKNW-DTKKYGDFSPVIAYSV-LVAK--VE--KGKAKLKPVKELVGITIME	RTKFEA	1178
WP_049531101	1125	VVDAS	KLI P I K S - G - - - L - S P E K Y G G Y A R P T I A Y S V - L V I A D - - I E - - K G A K K L K R I K E M V G I T I Q D	KKKFEA	1188
WP_049538452	1125	VVDAS	KLI P I K S - G - - - L - S P E K Y G G Y A R P T I A Y S V - L V I A D - - I E - - K G T K K L K R I K E M I G I T V Q D	KKIFES	1188
WP_049549711	1127	VVDAS	KLI P I K S - G - - - L - S P E K Y G G Y A R P T I A Y S V - L V I A D - - I E - - K G T K K L K R I K E M V G I T I Q D	KKKFEA	1190
WP_007896501	1121	--NSD	KLI E - - - - R K K G W - D P K K Y G G F D S P N T A Y S I - F V W A K - - V A - - K R K A Q K L K T V K E I V G I T I M E	QAEYEK	1182
EFR44625	1073	--NSD	KLI E - - - - R K K G W - D P K K Y G G F D S P N T A Y S I - F V W A K - - V A - - K R K A Q K L K T V K E I V G I T I M E	QAEYEK	1134
WP_002897477	1119	--NSD	KLI PRKT-KDILW-DTKKYGDFSPVIAYSI-LLIAD--IE--KGKAKLKTVKTLVGVTIME	KAAFEE	1183

WP_002906454	1124	vvDAS	KLIPKS-S--L-SPEKYGGYARPTIAYSV-LVIAD--IEkGKAKKAKKRIKEIVGITIQD	KKKFEF	1189
WP_009729476	1120	--NSD	KLIPRKT-KDILW-DTTKYGGFDSPIVAYSI-LLIAD--IE--KGKAKKLTVKTLVGITIME	KDAFEK	1184
CQR24647	1110	--GSD	KLIAARKT-KNNYL-STQKYGGFDSPTVAYSIMFVAD--IE--KGKSKRLKTKVKEMIGITIME	RSRFES	1174
WP_000066813	1124	--NSD	KLIPRKT-KEILW-DTTKYGGFDSPIVAYSI-LLIAD--IE--KGKAKKLTVKTLVGITIME	KATFEK	1188
WP_009754323	1120	--NSD	KLIPRKT-KDILW-DTTKYGGFDSPIVAYSI-LLIAD--IE--KGKAKKLTVKTLVGITIME	KAAFEK	1184
WP_044674937	1113	--DSD	KLIPRKT-EKFYL-DTKKYGGFDSPTIAYSV-LLIAD--IE--KGKAKKLRVKELIGITIME	RMAFEK	1177
WP_044676715	1115	--DSD	KLIPRKT-EKFYL-DTKKYGGFDSPTIAYSV-LLIAD--IE--KGKAKKLRVKELIGITIME	RMAFEK	1179
WP_044680361	1115	--DSD	KLIPRKT-EKFYL-DTKKYGGFDSPTIAYSV-LLIAD--IE--KGKAKKLRVKELIGITIME	RMAFEK	1179
WP_044681799	1113	--DSD	KLIPRKT-EKFYL-DTKKYGGFDSPTIAYSV-LLIAD--IE--KGKAKKLRVKELIGITIME	RMAFEK	1177
WP_049533112	1123	--DSS	ENLVGVK-RNL--DPKKYGGYAGISNSYAV-LVKAI-IE--KGVKKKETTMLVFQGISILD	RITFEK	1185
WP_029090905	1062	--SSS	KTIP----LKKHL-DTAIYGGYTAVNYASYA--LIQ-IE--FK--KGRK--REIIGIPLAV	QTRIDN	1117
WP_006506696	1085	haDKG	AVVP---vNKNRS-DVHKYGGFSG--LQYTI-IE--VA--IEgqKKKGTTELKVKISGVPLHL	KAASIN	1149
AIT42264	1115	--NSD	KLIA----RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--KGKSKLKSVKELLGITIME	RSSFEEK	1176
WP_034440723	1093	k-KTE	KRIP----IKNNL-DPNIYGGYIEEKMAYYI-IE--AINYLE--NGKTKK-IE--AIVGISIKD	KKDFEG	1149
AKQ21048	1115	--NSD	KLIA----RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--KGKSKLKSVKELLGITIME	RSSFEEK	1176
WP_004636532	1093	--DSS	KLIP----VKEGM-DPQKYGGLSQVSEAFV-VIT-IE--HE--KGKSKLKSVKELLGITIME	QKAYEQ	1150
WP_002364836	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--SDLISIPVD	KTRFEQ	1156
WP_016631044	1050	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1107
EMS75795	835	--KPD	KLIP----RKAGW-DVSKYGGFSGPVVAYAV-AFI-IE--YE--KPKAR--KKAKAIEGITIMK	QSLFEQ	892
WP_002373311	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_002378009	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_002407324	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_002413717	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_010775580	1101	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1158
WP_010818269	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_010824395	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_016622645	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_033624816	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_033625576	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_033789179	1099	--PSN	KLIP----VKNGL-DPQKYGGFDSPIVAYTV-LF-T-IE--HE--KGK-KPL-IE--KQEIILGITIME	KTRFEQ	1156
WP_002310644	1101	--DSS	KLIP----RKNW-DPAKYGGLSGNVAYTV-AFT-IE--YE--KPKAR--KRTNALEGITIME	REAFEQ	1158
WP_002312694	1102	--DSS	KLIP----RKNW-DPAKYGGLSGNVAYTV-AFT-IE--YE--KPKAR--KRTNALEGITIME	REAFEQ	1159
WP_002314015	1102	--DSS	KLIP----RKNW-DPAKYGGLSGNVAYTV-AFT-IE--YE--KPKAR--KRTNALEGITIME	REAFEQ	1159
WP_002320716	1102	--DSS	KLIP----RKNW-DPAKYGGLSGNVAYTV-AFT-IE--YE--KPKAR--KRTNALEGITIME	REAFEQ	1159

WP_002330729	1101	--DSS	KLIP----	RKNW-DPAKYGGGLGSPNVAYTV-AFT-----YE--KGKAR--KRTNALEGITIME	REAFEQ	1158
WP_002335161	1102	--DSS	KLIP----	RKNW-DPAKYGGGLGSPNVAYTV-AFT-----YE--KGKAR--KRTNALEGITIME	REAFEQ	1159
WP_002345439	1102	--DSS	KLIP----	RKNW-DPTKYGGGLGSPNVAYTV-AFT-----YE--KGKAR--KRTNALEGITIME	REAFEQ	1159
WP_034867970	1092	--KPD	KLIE----	RKNW-DVTKYGGFGSPVIAYAI-AFV-----YA--KGTQ--KKTRAIEGITIME	QAAFEK	1149
WP_047937432	1102	--DSS	KLIP----	RKNW-DPAKYGGGLGSPNVAYTV-AFT-----YE--KGKAR--KRTNALEGITIME	REAFEQ	1159
WP_010720994	1092	--KPD	KLIE----	RKNW-DVTKYGGFGSPVIAYAI-AFV-----YA--KGTQ--KKTKAIEGITIME	QAAFEK	1149
WP_010737004	1092	--KPD	KLIE----	RKNW-DVTKYGGFGSPVIAYAI-AFV-----YA--KGTQ--KKTKAIEGITIME	QAAFEK	1149
WP_034700478	1092	--KPD	KLIE----	RKNW-DVTKYGGFGSPVIAYAI-AFV-----YA--KGTQ--KKTKAIEGITIME	QAAFEK	1149
WP_007209003	1090	--ESQ	KLIR----	RKQW-NTKKYGGFGSPVAYAI--LLS-FD--KGK-RKARSK-IVGITIQD	RESFEG	1147
WP_023519017	1086	--NPE	KLIP----	RKASL-DPLKYGGYSPVAYTV-IFI---FE--KGKQK--KVTKIEGITIME	QLRFEQ	1143
WP_010770040	1097	--DSD	KLIS----	RKTNW-SPKLYGGFGSPVAYSV-II-T--YE--KGK-KKVR-AKIVGITIME	QSLFKK	1154
WP_048604708	1094	--DSD	KLIS----	RKKEW-DTTKYGGFGSPVAYSV-VI-R--YE--KGK-TRKLV-KTIVGITIME	RAAFEK	1151
WP_010750235	1095	--KPD	KLIK----	RKNW-DVTKYGGFGSPVAYAV-VFT-----YE--KGKNH--KKAKAIEGITIME	QALFEK	1152
AI116583	1154	--NSD	KLIA----	RKKDW-DPKKYGGFGSPVAYSV-LVWAK--VE--KGSKKLKSVKELLGITIME	RSSF EK	1215
WP_029073316	1100	hsEKG	AKVP----	vNKLRS-NVHKYGGFEG--LKYSI-----VA--IKgkKKKGGKKIIDVNLVGIPLMY	KNVDDE	1164
WP_031589969	1100	nsDKD	ATVP----	vNKYRS-NVKYGGFSG--VNSFI-----VA--IKgkKKKGGKKKIEVNLKGTGIPLMY	KNADEE	1164
KDA45870	1079	--DSG	KLIP----	RKGL-DPVKYGGYAKAVESYAV-LITAD-eVK--KGTKKVKT--LVNIPIID	SKKYEA	1138
WP_039099354	1098	k-ASG	QLIPAKQdRPTAL----	YGGYSGKTAYMC---IVR--IKnkKGDLYKVCVETSWLAQLKQ	KKAFK	1170
AKP02966	1119	k----	KLIA----	QKDM-DPNIYGGFGDNKSSIT--IVK-ID---NNKIKPVA--IPIRLIN	----DK	1172
WP_010991369	1096	--NSS	KLIP----	RKTNW-DPMKYGGLDSPNMAYAV-VI-E--YA--KGK-NKLVEFKKIIIRVTIME	RKAFEK	1154
WP_033838504	1096	--NSS	KLIS----	RKTNW-DPMKYGGLDSPNMAYAV-VI-E--YA--KGK-NKLVEFKKIIIRVTIME	RKAFEK	1154
EHN60060	1099	--NSS	KLIS----	RKTNW-DPMKYGGLDSPNMAYAV-VI-E--YA--KGK-NKLVEFKKIIIRVTIME	RKAFEK	1157
EFR89594	865	--NSS	KLIP----	RKTNW-DPMKYGGLDSPNMAYAV-VI-E--YA--KGK-NKLVEFKKIIIRVTIME	RKAFEK	923
WP_038409211	1096	--NSS	KLIS----	RKADW-NPIKYGGFGDGSNMAYSI-VI-E--YE--KRK-KKTVIKELLIQINIME	RVAF EK	1154
EFR95520	715	--NSS	KLIS----	RKADW-NPIKYGGFGDGSNMAYSI-VI-E--YE--KRK-KKTVIKELLIQINIME	RVAF EK	773
WP_003723650	1096	--NSS	KLIP----	RKENW-DPMKYGGLDSPNMAYAV-II-E--HA--KGK-KKIVIEKKLIQINIME	RKMFEK	1154
WP_003727705	1096	--NSS	KLIP----	RKENW-DPMKYGGLDSPNMAYAV-II-E--HA--KGK-KKIVIEKKLIQINIME	RKMFEK	1154
WP_003730785	1096	--NSS	KLIP----	RKENW-DPVKYGGLDSPNMAYAV-II-E--HA--KGK-KKIVIEKKLIQINIME	RKMFEK	1154
WP_003733029	1096	--KSN	KLIP----	RKDW-DPIKYGGFGSKMAYAI-II-E--YE--KQK-RKVRIEKKLIQINIME	REAF EK	1154
WP_003739838	1096	--NSS	KLIP----	RKENW-DPMKYGGLDSPNMAYAV-II-E--HA--KGK-KKIVIEKKLIQINIME	RKAF EK	1154
WP_014601172	1096	--NSS	KLIP----	RKENW-DPMKYGGLDSPNMAYAV-II-E--HA--KGK-KKIVIEKKLIQINIME	RKMFEK	1154
WP_023548323	1096	--DSS	KLIP----	KKTNL-NPIKYGGFEGSNMAYAI-II-E--HE--KRK-KKVTIEKKLIQINIME	RKAF EK	1154
WP_031665337	1096	--NSS	KLIP----	RKENW-DPMKYGGLDSPNMAYAV-II-E--HA--KGK-KRIVIEKKLIQINIME	RKMFEK	1154
WP_031669209	1096	--KSN	KLIP----	RKDW-DPIKYGGFGDGSNMAYAI-II-E--YE--KQK-RKVRIEKKLIQINIME	REAF EK	1154

WP_033920898	1096	--DSS	KLIP-----KKTNL-NPIKYGGFEGSNMAYAI-II-E--HE--KRK-KKVTIEKKLIQINIME	RKAFEK	1154
AKI42028	1099	--NSS	KLIP-----RKENW-DPMKYGGGLDSPNMAYAV-II-E--HA--KGG-KKLI FEKKIIRITIME	RKMFEK	1157
AKI50529	1099	--DSS	KLIP-----KKTNL-NPIKYGGFEGSNMAYAI-II-E--HE--KRK-KKVTIEKKLIQINIME	RKAFEK	1157
EFR83390	544	--NSS	KLIP-----RKENW-DPMKYGGGLDSPNMAYAV-II-E--HA--KGG-KKIVIEKKLIQINIME	RKMFEK	602
WP_046323366	1096	--NSS	KLIP-----RKADW-DPIKYGGFDGSMAYAI-VI-E--HE--KRK-KKTVIKKELIQINIME	RTAFEK	1154
AKE81011	1131	--NSD	KLIA-----RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--KGSKKLKSVKELLGITIME	RSSF EK	1192
CUO82355	1089	hsAKG	AVIP---vNKNRK-DVNKYGGFSG--LQYVI-----AA--IEgtKKKGGKLVKVRKLSGIPLYL	KQADIK	1153
WP_033162887	1090	hsEKG	ATVP---LNKYRA-DVHKYGGFGN--VQSI I-----VA--IEgkKKKGGKLI D VRKLT S I P L H L	KNAPVE	1154
AGZ01981	1148	--NSD	KLIA-----RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--KGSKKLKSVKELLGITIME	RSSF EK	1209
AKA60242	1115	--NSD	KLIA-----RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--KGSKKLKSVKELLGITIME	RSSF EK	1176
AKS40380	1115	--NSD	KLIA-----RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--KGSKKLKSVKELLGITIME	RSSF EK	1176
4UN5_B	1119	--NSD	KLIA-----RKKDW-DPKKYGGFDSPTVAYSV-LVWAK--VE--KGSKKLKSVKELLGITIME	RSSF EK	1180

WP_010922251	1177	NPI---	DFLE---	AKGYKE--	V-KKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_039695303	1186	NPV---	EFLE---	NKGYHN--	I-REDKLIK--	LPKYSLFE--	FEGRRRLLAS	1248	ASELQKGNEMVLPGLVELLYHA
WP_045635197	1184	NPI---	TFLE---	NKGYHN--	V-RKENILC--	LPKYSLFE--	LENGRRLLAS	1246	AKELQKGNELVLPVYLTTLLYHS
5AXW A	913	KPYrfd	VYLD---	NGVYKFvt	V-KNLDVIK---	KENYVE--	VNSKAYEEAKK	978	-KKSNOAEFIASFYNNDLJKIN
WP_009880683	861	DPV---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	923	-GELQKGNELALPSKYVNFYLA
WP_010922251	1177	NPI---	DFLE---	AKGYKE--	V-KKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_011054416	1177	DPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_011284745	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIVK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_011285506	1177	NPI---	DFLE---	AKGYKE--	V-KKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_011527619	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_012560673	1177	DPV---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_014407541	1176	NPI---	DFLE---	AKGYKE--	V-KKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1238	-GELQKGNELALPSKYVNFYLA
WP_020905136	1177	NPI---	DFLE---	AKGYKE--	V-KKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_023080005	1176	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1238	-GELQKGNELALPSKYVNFYLA
WP_023610282	1176	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1238	-GELQKGNELALPSKYVNFYLA
WP_030125963	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_030126706	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_031488318	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_032460140	1177	DPV---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_032461047	1177	DPV---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_032462016	1177	NPI---	DFLE---	AKGYKE--	V-KKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_032462936	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_032464890	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_033888930	1002	NPI---	DFLE---	AKGYKE--	V-KKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1064	-GELQKGNELALPSKYVNFYLA
WP_038431314	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_038432938	1176	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1238	-GELQKGNELALPSKYVNFYLA
WP_038434062	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
BAQ51233	1088	NPI---	DFLE---	AKGYKE--	V-KKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1150	-GELQKGNELALPSKYVNFYLA
KGE60162	352	DPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	414	-GELQKGNELALPSKYVNFYLA
KGE60856	115	DPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	177	-GELQKGNELALPSKYVNFYLA
WP_002989955	1177	NPI---	DFLE---	AKGYKE--	V-RKDLIIK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGNELALPSKYVNFYLA
WP_003030002	1159	HPV---	DFLE---	QRYRN--	V-RLEKIIK--	LPKYSLFE--	LENKRRLIAS	1221	ARELQKGNELVLPQRFVTTLLYHS
WP_003065552	1187	NPV---	EFLE---	NKGYHN--	I-REDKLIK--	LPKYSLFE--	FEGRRRLLAS	1249	ASELQKGNEMVPGHLVKLLYHA
WP_001040076	1178	NPS---	AFLE---	SKGYLN--	I-RTDKLII--	LPKYSLFE--	LENGRRLLAS	1240	AGELQKGNELALPTQFMKFLYLA

WP_001040078 1186 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040080 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040081 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGETIDRLQKGNELALPTQFMKFLYLA
 1240
 WP_001040083 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040085 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040087 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040088 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040089 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040090 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040091 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040092 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040094 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040095 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040096 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040097 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS ADELQKGNELALPTQFMKFLYLA 1240
 WP_001040098 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040099 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040100 1178 NPS---AFLE---SKGYLD--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040104 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040105 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040106 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040107 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040108 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040109 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_001040110 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_015058523 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_017643650 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS ADELQKGNELALPTQFMKFLYLA 1240
 WP_017647151 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_017648376 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_017649527 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_017771611 1178 NPS---AFLE---SKGYLN--I-RDDKLMII--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 WP_017771984 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240
 CFQ25032 1178 NPS---AFLE---SKGYLN--I-RADKLI--LPKYSLFE---LENGRRRLLIAS AGELQKGNELALPTQFMKFLYLA 1240

CFV16040	1178	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
KLJ37842	1178	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
KLJ72361	1178	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
KL20707	1192	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1254	AGELQKGNELALPTQFMKFLYLA	1254
KL42645	1178	NPS	---AFLE---	SKGYLN	--I	RDDKLM	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
WP_047207273	1178	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
WP_047209694	1178	NPS	---AFLE---	SKGYLN	--I	RDDKLM	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
WP_050198062	1178	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
WP_050201642	1178	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
WP_050204027	1178	NPS	---AFLE---	SKGYLN	--I	RDDKLM	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
WP_050881965	1178	NLS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
WP_050886065	1178	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
AHN30376	1178	NPS	---AFLE---	SKGYLN	--I	RTDKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
EAO78426	1178	NPS	---AFLE---	SKGYLN	--I	RADKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
CCW42055	1178	NPS	---AFLE---	SKGYLN	--I	RTDKLI	I--LPKYSLFE---	LENGRRRLIAS	1240	AGELQKGNELALPTQFMKFLYLA	1240
WP_003041502	1186	DKR	---AFLL---	GKGYKD	--I	K--KII	E--LPKYSLFE---	LENGRRRLIAS	1253	RGEIHKGNELFVQKFTTLLYHA	1253
WP_037593752	1160	NPV	---DFLE---	QRYRN	--V	RLEKII	K--LPKYSLFE---	LKDGSRRLIAS	1222	ARELQKGNELVIPQRFRTLLYHS	1222
WP_049516684	1160	HPV	---DFLE---	QRYRN	--V	RLEKII	K--LPKYSLFE---	LENKRRRLIAS	1222	ARELQKGNELVIPQRFRTLLYHS	1222
GAD46167	1159	NPV	---DFLE---	QRYRN	--V	RLEKII	K--LPKYSLFE---	LENKRRRLIAS	1221	ARELQKGNELVIPQRFRTLLYHS	1221
WP_018363470	1191	NPV	---EFLK---	NKGYQN	--V	QEDKLM	K--LPKYSLFE---	FEGRRRLIAS	1253	ATELQKGNELMISAHLVALLYHA	1253
WP_003043819	1186	DPI	---GFLE---	AGYKD	--I	KKELIF	K--LPKYSLFE---	FEGRRRLIAS	1248	--ELQKANEVLVPHLVRLLYYT	1248
WP_006269658	1159	NPV	---DFLE---	QRYRN	--V	RLEKII	K--LPKYSLFE---	LENKRRRLIAS	1221	AKELQKGNELVIPQRFRTLLYHS	1221
WP_048800889	1179	NPI	---MFLE---	SKGYRN	--I	QKDKLI	K--LPKYSLFE---	FEGRRRLIAS	1241	AVELQKGNEMVLPQYLNLLYHA	1241
WP_012767106	1178	NPL	---KFIE---	DKGYGN	--V	QIDKCI	K--LPKYSLFE---	FENGTRRLIAS	1245	RGDLQKANEMFLPAKLVTLTY--	1245
WP_014612333	1178	NPL	---KFIE---	DKGYGN	--V	QIDKCI	K--LPKYSLFE---	FENGTRRLIAS	1245	RGDLQKANEMFLPAKLVTLTY--	1245
WP_015017095	1178	NPL	---KFIE---	DKGYGN	--V	QIDKCI	K--LPKYSLFE---	FENGTRRLIAS	1245	RGDLQKANEMFLPAKLVTLTY--	1245
WP_015057649	1178	NPL	---KFIE---	DKGYGN	--V	QIDKCI	K--LPKYSLFE---	FENGTRRLIAS	1245	RGDLQKANEMFLPAKLVTLTY--	1245
WP_048327215	1178	NPL	---KFIE---	DKGYGN	--V	QIDKCI	K--LPKYSLFE---	FENGTRRLIAS	1245	RGDLQKANEMFLPAKLVTLTY--	1245
WP_049519324	1178	NPL	---KFIE---	DKGYGN	--V	QIDKCI	K--LPKYSLFE---	FENGTRRLIAS	1245	RGDLQKANEMFLPAKLVTLTY--	1245
WP_012515931	1157	NPV	---VFLE---	ARGYRE	--I	QEHKII	K--LPKYSLFE---	LENGRRRLIAS	1219	-SELQKGNELFLPVDYMTFLYLA	1219
WP_021320964	1157	NPV	---VFLE---	ARGYRE	--I	QEHKII	K--LPKYSLFE---	LENGRRRLIAS	1219	-SELQKGNELFLPVDYMTFLYLA	1219
WP_037581760	1157	NPV	---VFLE---	ARGYRE	--I	QEHKII	K--LPKYSLFE---	LENGRRRLIAS	1219	-SELQKGNELFLPVDYMTFLYLA	1219
WP_004232481	1189	NPV	---SFLE---	KKGYHN	--V	QEDKLI	K--LPKYSLFE---	FEGRRRLIAS	1251	ATELQKGNELVLPQYMNLLYHS	1251
WP_009854540	1184	NPV	---EFLE---	NKGYHN	--I	REDKLI	K--LPKYSLFE---	FEGRRRLIAS	1246	ASELQKGNEMVLPGYLVELLYHA	1246

WP_012962174	1185	NPV---VFLE---KKGYN--V-QEDNLIK--LPKYSLFE---FEGRRRLIAS	ASELQKGNVWLSRHLVELLYHA	1247
WP_039695303	1186	NPV---EFLE---NKGYN--I-REDKLIK--LPKYSLFE---FEGRRRLIAS	ASELQKGNEMVLPGLVELLYHA	1248
WP_014334983	1189	NPV---SFLE---KKGYN--V-QEDKLIK--LPKYSLFE---FEGRRRLIAS	ATELQKGNEMVLPahlVELLYHA	1251
WP_003099269	1177	DPI---AFLE---KGYQD--I-QTSSIIK--LPKYSLFE---LENGRKRLLAS	--ELQKGNELALPNKYVKFLYLA	1239
AHY15608	1177	DPI---AFLE---KGYQD--I-QTSSIIK--LPKYSLFE---LENGRKRLLAS	--ELQKGNELALPNKYVKFLYLA	1239
AHY17476	1177	DPI---AFLE---KGYQD--I-QTSSIIK--LPKYSLFE---LENGRKRLLAS	--ELQKGNELALPNKYVKFLYLA	1239
ESR09100	9	DPI---AFLE---KGYQD--I-QTSSIIK--LPKYSLFE---LENGRKRLLAS	-KELQKGNELALPNKYVKFLYLA	71
AGM98575	1177	DPI---AFLE---KGYQD--I-QTSSIIK--LPKYSLFE---LENGRKRLLAS	--ELQKGNELALPNKYVKFLYLA	1239
ALF27331	1159	NPV---AFLE---RKGYN--V-QEENIVK--LPKYSLFE---LENGRRRLIAS	ARELQKGNELALPNHLGTMLYHA	1221
WP_018372492	1168	EPEr---FLA---QKGFER--V-EKT--IK--LPKYSLFE---MEKRRRLIAS	SGELQKGNQVLLPEHLIRLLSYA	1228
WP_045618028	1189	NPI---AYLE---ECGYKN--I-NPNLIK--LPKYSLFE---FNNQRRRLIAS	SIELQKGNELIVPYHFTALLYHA	1251
WP_045635197	1184	NPI---TFLE---NKGYN--V-RKENILC--LPKYSLFE---LENGRRRLIAS	AKELQKGNELIVPVYLTLLYHS	1246
WP_002263549	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002263887	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002264920	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002269043	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002269448	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002271977	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002272766	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002273241	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002275430	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002276448	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002277050	1167	NPV---KFLK---DKGYQD--I-EKNNFVK--LPKYTLVD---INGIKRLWAS	SKEVHKGNQLVWSKKSQDLYHA	1229
WP_002277364	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002279025	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002279859	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002280230	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002281696	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002282247	1167	NPV---KFLK---DKGYQD--I-EKNNFVK--LPKYTLVD---INGIKRLWAS	SKEVHKGNQLVWSKKSQDLYHA	1229
WP_002282906	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002283846	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002287255	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002288990	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221
WP_002289641	1159	DPV---AFLE---RKGYN--V-QEENIIC--LPKYSLFE---LENGRKRLLAS	ARELQKGNELIVLPNHLGTMLYHA	1221

WP_002290427	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_002295753	1159	DPI---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_002296423	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_002304487	1173	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1235
WP_002305844	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_002307203	1159	DPI---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_002310390	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_002352408	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_012997688	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_014677909	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_019312892	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_019313659	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_019314093	1159	DPI---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_019315370	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_019803776	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_019805234	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_024783594	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_024784288	1167	NPV---KFLK---DKYQO--I-EKNFVK--LPKYTLVD---INGIKRLWAS	SKEVHKGQNLVSKKSQDLEYHA	1229
WP_024784666	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_024784894	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_024786433	1167	NPV---KFLK---DKYQO--I-EKNFVK--LPKYTLVD---INGIKRLWAS	SKEVHKGQNLVSKKSQDLEYHA	1229
WP_049473442	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
WP_049474547	1159	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1221
EMC03581	1152	DPV---AFLE---RKYRN--V-QEENIIK--LPKYSLFK---LENGRKRLLAS	ARELQKGNIEIVLPNHLGTLLEYHA	1214
WP_000428612	1187	SPI---AFLE---NKYHN--V-RKENILC--LPKYSLFE---LKNGRRRMLAS	AKELQKGNIEIVLPVHLTTLLEYHA	1249
WP_000428613	1185	NPI---TFLE---NKYHN--V-RKENILC--LPKYSLFE---LENGRRRLLAS	AKELQKGNIEIVLPVYLTTLLEYHS	1247
WP_049523028	1180	NPV---AFLE---KGYQN--V-VEENIIR--LPKYSLFE---LENGRRRMLAS	AKELQKGNEMVLPYLIALLYHA	1242
WP_003107102	1146	DRI---TFLE---KKYQD--I-QESLIK--LPKFSLFE---LENGRKRLLAS	--ELQKGNELSLPNKYIQFLYLA	1208
WP_054279288	1179	NPI---AFLE---SKYHD--I-QEHLMIT--LPKYSLFE---LENGRRRLLAS	--ELQKGNEMVLPQHLVTFYRV	1241
WP_049531101	1189	NPT---AYLE---EYGYK--I-NPNLIK--LPKYSLFE---FNDGQRRLLAS	SIELQKGNELILPYHFTTLLEYHA	1251
WP_049538452	1189	NPI---AYLE---ECGYK--I-NPNLIK--LPKYSLFE---FNGGQRRLLAS	SIELQKGNELILPYHFTALLYHT	1251
WP_049549711	1191	NPI---AYLE---ECGYK--I-NPNLIK--LPKYSLFE---FNGGQRRLLAS	SIELQKGNELILPYHFTALLYHA	1253
WP_007896501	1183	DNI---AFLE---KKYQD--I-QEKLIIK--LPKYSLFE---LENGRRRLLAS	--EFQKGNELALSGKYMFLYLA	1245
EFR44625	1135	DNI---AFLE---KKYQD--I-QEKLIIK--LPKYSLFE---LENGRRRLLAS	--EFQKGNELALSGKYMFLYLA	1197

WP_002897477	1184	NPI---	TFLE---	NKGYHN--	V-RKENILC--	LPKYSLFE--	LENGRRRLIAS	1246	AKELQKGN	EIVL	PVCL	TTLLYYHS
WP_002906454	1190	NPV---	TYLE---	ECGYKN--	I-NSNLIK--	LPKYSLFE--	FNDGQRRLLAS	1252	SIELQKGN	ELIL	PLPH	TALLYYHA
WP_009729476	1185	NPI---	AFLE---	NKGYHN--	V-CKENILC--	LPKYSLFE--	LENGRRRLIAS	1247	AKELQKCN	EIVL	PVYL	TTLLYYHS
CQR24647	1175	NSV---	TFLE---	EKGYRN--	I-RENTIHK--	FPKYSLFE--	LENGRRRLIAS	1237	AIELQKGN	EMFL	PQQF	VNLLYYHA
WP_000066813	1189	NPI---	TFLE---	NKGYHN--	V-RKENILC--	LPKYSLFE--	LESGRRMLAS	1251	AKELQKGN	EIVL	PVYL	TTLLYYHS
WP_009754323	1185	NPI---	TFLE---	NKGYHN--	V-RKENILC--	LPKYSLFE--	LENGRRRLIAS	1247	AKELQKGN	EIVL	PVYL	TTLLYYHS
WP_044674937	1178	NPI---	EFLE---	HKGYKN--	I-LEKNIHK--	LPKYSLFE--	LENGRRRLIAS	1240	AKELQKGN	EMIL	PPHL	VTLLYYHS
WP_044676715	1180	NPI---	EFLE---	HKGYKN--	I-LEKNIHK--	LPKYSLFE--	LENGRRRLIAS	1242	AKELQKGN	EMIL	PPHL	VTLLYYHS
WP_044680361	1180	NPI---	EFLE---	HKGYKN--	I-LEKNIHK--	LPKYSLFE--	LENGRRRLIAS	1242	AKELQKGN	EMIL	PPHL	VTLLYYHS
WP_044681799	1178	NPI---	EFLE---	HKGYKN--	I-LEKNIHK--	LPKYSLFE--	LENGRRRLIAS	1240	AKELQKGN	EMIL	PPHL	VTLLYYHS
WP_049533112	1186	DKR---	AFLL---	GKGYKD--	I-K-KIIE--	LPKYSLFE--	LKDGSRRLMAS	1253	RGEIHKGN	ELFV	PQKF	TTLLYYHA
WP_029090905	1118	SETsl	qAYIA--	EQIKSE--	VeILN--	grILKYQLIS--	NNGNRLYIAG	1181	--SERHNA	RQLIVS	DEAAKVIWLI	
WP_006506696	1150	EKI---	NYIE--	eKEGLSD--	VrIHK--	Dn-IPVNQMIEm--	DGGEYLLTS	1211	--EYVNAR	QVLV	NEKQCALIADI	
AIT42264	1177	NPI---	DFLE---	AKGYKE--	V-KKDLIHK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGN	ELAL	PSKYVNFVLYLA	
WP_034440723	1150	QTT---	EYLG---	KIGFNK--	AsIIN--	S-FKNYTLFE--	LENGSRMIVG	1217	KGELQKGN	QMYL	LPQNLLEFVYHL	
AKQ21048	1177	NPI---	DFLE---	AKGYKE--	V-KKDLIHK--	LPKYSLFE--	LENGRKRMLAS	1239	-GELQKGN	ELAL	PSKYVNFVLYLA	
WP_004636532	1151	HPT---	AYLE---	EAGYNN--	P-TV--	LHE--LPKYQLFE--	LEDGSRRLMAS	1211	AKEFQKGN	QMYL	PLLELVVELLYHA	
WP_002364836	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKEAQKGN	QMYL	PEHLLTLLYYHA	
WP_016631044	1108	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1168	AKEAQKGN	QMYL	PEHLLTLLYYHA	
EMS75795	893	DPI---	GFLS---	NKGYSN--	V-TKF--	IK--LSKYTYLFE--	LENGRRRMLAS	953	-KEAQKAN	SFIL	PEKLV	TTLLYYHA
WP_002373311	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_002378009	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_002407324	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_002413717	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_010775580	1159	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1219	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_010818269	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_010824395	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_016622645	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_033624816	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PERLLTLLYYHA	
WP_033625576	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_033789179	1157	NPI---	LFLE---	EKGFLR--	P-RV--	LMK--LPKYTYLFE--	FPEGRRRLIAS	1217	AKAQKGN	QMYL	PEHLLTLLYYHA	
WP_002310644	1159	SPV---	LFK---	NKGYEQ--	A-EIE--	MK--LPKYALFE--	LENGRKRMLAS	1219	-KEAQKAN	SFLL	PEHLLTLLYYHA	
WP_002312694	1160	SPV---	LFK---	NKGYEQ--	A-EIE--	MK--LPKYALFE--	LENGRKRMLAS	1220	-KEAQKAN	SFLL	PEHLLTLLYYHA	
WP_002314015	1160	SPV---	LFK---	NKGYEQ--	A-EIE--	MK--LPKYALFE--	LENGRKRMLAS	1220	-KEAQKAN	SFLL	PEHLLTLLYYHA	

WP_002320716	1160	SPV---	LFLK---	NKGYEQ--	A-EIE--	MK--	LPKYALFE--	LENGRKRMMVAS	1210	-KEAQKANSFLLPEHLVTLLEYHA	1210
WP_002330729	1159	SPV---	LFLK---	NKGYEQ--	A-EIE--	MK--	LPKYALFE--	LENGRKRMMVAS	1219	-KEAQKANSFLLPEHLVTLLEYHA	1219
WP_002335161	1160	SPV---	LFLK---	NKGYEQ--	A-EIE--	MK--	LPKYALFE--	LENGRKRMMVAS	1220	-KEAQKANSFLLPEHLVTLLEYHA	1220
WP_002345439	1160	SPV---	LFLK---	NKGYEQ--	A-EIE--	MK--	LPKYALFE--	LENGRKRMMVAS	1220	-KEAQKANSFLLPEHLVTLLEYHA	1220
WP_034867970	1150	DPT---	TFLK---	EKGFPO--	V-TEF--	IK--	LPKYTLFE--	FNGRRRFLAS	1210	-KESQKGNPFILSDQLVTLLEYHA	1210
WP_047937432	1160	SPV---	LFLK---	NKGYEQ--	A-EIE--	MK--	LPKYALFE--	LENGRKRMMVAS	1220	-KEAQKANSFLLPEHLVTLLEYHA	1220
WP_010720994	1150	DPT---	TFLK---	DKGFPQ--	V-TEF--	IK--	LPKYTLFE--	FNGRRRFLAS	1210	-KESQKGNPFILSDQLVTLLEYHA	1210
WP_010737004	1150	DPT---	TFLK---	EKGFPO--	V-TEF--	IK--	LPKYTLFE--	FNGRRRFLAS	1210	-KESQKGNPFILSDQLVTLLEYHA	1210
WP_034700478	1150	DPT---	TFLK---	DKGFPH--	V-TEF--	IK--	LPKYTLFE--	FNGRRRFLAS	1210	-KESQKGNPFILSDQLVTLLEYHA	1210
WP_007209003	1148	NPII---	YLS---	KKDYHN--	pKVEAI--		LPKYSLFE--	FENGRRRMMVAS	1208	-SETQKGNQLIIPGHLMELLYHS	1208
WP_023519017	1144	DPR---	EFLK---	TKGYEG--	V-KQW--	LJ--	LPKYILFE--	AQGGYRRMIAS	1204	-QETQKANSILPENLVTLLYYHA	1204
WP_010770040	1155	DPV---	SLE---	EKGYAN--	P-EV--	LJH--	LPKYTYLFE--	LENGRRRLLIAS	1215	ANEAQKGNQVLPAHLVTLLEYHA	1215
WP_048604708	1152	NER---	EFLK---	NKGYQN--	P-QI--	CMK--	LPKYSLYE--	FDDGRRRLLIAS	1212	AKEAQKGNQVLPALHVTFLYYHA	1212
WP_010750235	1153	DPI---	SFLI---	EKGYSN--	V-NQF--	IK--	LPKYTLFE--	LANGQRRMLAS	1213	-QELQKANSFILPEKLVTLLEYHA	1213
AI116583	1216	NPI---	DFLE---	AKGYKE--	V-KKDLI IK--		LPKYSLFE--	LENGRKRMLAS	1278	-GELQKGNELALPSKYVNFLLYA	1278
WP_029073316	1165	TKI---	NYIK--	eSEGLEE--	VkI IK--	E--	ILKNQLIEi--	NGGLFYVTS	1225	--EIVNARQLILDNFNCTRIIDGI	1225
WP_031589969	1165	IKI---	NYLK--	qAEDLEE--	VqIGK--	E--	ILKNQLIEk--	GGGLYYIVA	1225	--EIVNARQLILDNFNCTRIIDGI	1225
KD445870	1139	DPT---	AYLA--	SRGYTNvtNsFIL--			PKYSLLEd--	PEGRRRYLAS	1199	-KEFQKANELILPQHLVELLYWV	1199
WP_039099354	1171	QKI--	spQFTKv--	KKQKgtiV--	KVVEDFEV--	IAPHILINqr	FDNGQELTLGS		1241	---HNEQELILDKTAVKLLNGA	1241
AKP02966	1173	CTL--	qNWLE--	ENVKHKksI	qI IK--	Nn--	VPIGQIY--	SKKVGLLS	1237	-REIANRQQLILPPEHSALLRIL	1237
WP_010991369	1155	DEK---	AFLE--	EQGYRQ--	P-KV--	LAK--	LPKYTYLFE--	CEEGRRRMLAS	1215	ANEAQKGNQQLVLPNHLVTLLEYHA	1215
WP_033838504	1155	DEK---	AFLE--	EQGYRQ--	P-KV--	LAK--	LPKYTYLFE--	CEEGRRRMLAS	1215	ANEAQKGNQQLVLPNHLVTLLEYHA	1215
EHN60060	1158	DEK---	AFLE--	EQGYRQ--	P-KV--	LAK--	LPKYTYLFE--	CEEGRRRMLAS	1218	ANEAQKGNQQLVLPNHLVTLLEYHA	1218
EFR89594	924	DEK---	AFLE--	EQGYRQ--	P-KV--	LAK--	LPKYTYLFE--	CEEGRRRMLAS	984	ANEAQKGNQQLVLPNHLVTLLEYHA	984
WP_038409211	1155	DQK---	AFLE--	EKGYS--	P-KV--	LTK--	IPKYTYLFE--	CENGRRRMLGS	1215	ANEAQKGNQQLVLPNHLVTLLEYHA	1215
EFR95520	774	DQK---	AFLE--	EKGYS--	P-KV--	LTK--	IPKYTYLFE--	CENGRRRMLGS	834	ANEAQKGNQQLVLPNHLVTLLEYHA	834
WP_003723650	1155	DEE---	AFLE--	EKGYRH--	P-KV--	LTK--	LPKYTYLFE--	CEKRRRMLAS	1215	ANEAQKGNQQLVLSNHLVSLLEYHA	1215
WP_003727705	1155	DEE---	AFLE--	EKGYHQ--	P-KV--	LTK--	LPKYTYLFE--	CEKRRRMLSS	1215	ANEAQKGNQQLVLSNHLVSLLEYHA	1215
WP_003730785	1155	DEE---	AFLE--	EKGYHQ--	P-KV--	LTK--	LPKYTYLFE--	CEKRRRMLSS	1215	ANEAQKGNQQLVLSNHLVSLLEYHA	1215
WP_003733029	1155	DEK---	TFLF--	EKGYHQ--	P-KV--	LTK--	VPKYTYLFE--	CKNRRRMLGS	1215	ANEAHKGNQMLLPNHLMALLYHA	1215
WP_003739838	1155	DEK---	SFLE--	EKGYRQ--	P-KV--	LTK--	LPKYTYLFE--	CENGRRRMLAS	1215	ANEAQKGNQQLVLSNHLVSLLEYHA	1215
WP_014601172	1155	DEE---	AFLE--	EKGYRH--	P-KV--	LTK--	LPKYTYLFE--	CEKRRRMLAS	1215	ANEAQKGNQQLVLSNHLVSLLEYHA	1215
WP_023548323	1155	DEK---	VFLE--	GKGYHQ--	P-KV--	LTK--	LPKYALFE--	CENGRRRMLGS	1215	ANEVHKGNQMLLPNHLMTLLYYHA	1215
WP_031665337	1155	DEE---	AFLE--	EKGYRH--	P-KV--	LTK--	LPKYTYLFE--	CEKRRRMLAS	1215	ANEAQKGNQQLVLSNHLVSLLEYHA	1215

WP_031669209	1155	DEK---TFLE---EKGYHQ--P-KV--LIK--VPKYTYLIE---CENGRRRMLGS	ANEAHKGNOMLLPNHLMTLLYHA	1215
WP_033920898	1155	DEK---VFLE---GKGYHQ--P-KV--LTK--LPKYALYE---CENGRRRMLGS	ANEVHKGNOMLLPNHLMTLLYHA	1215
AKI42028	1158	DEE---AFLE---EKGYRH--P-KV--LTK--LPKYTYLIE---CEKGRRRMLAS	ANEAQKGNQMLVLSNHLVSLLYHA	1218
AKI50529	1158	DEK---VFLE---GKGYHQ--P-KV--LTK--LPKYALYE---CENGRRRMLGS	ANEVHKGNOMLLPNHLMTLLYHA	1218
EFR83390	603	DEE---AFLE---EKGYRH--P-KV--LTK--LPKYTYLIE---CEKGRRRMLAS	ANEAQKGNQMLVLSNHLVSLLYHA	663
WP_046323366	1155	DQK---EFLE---GKGYRN--P-KV--ITK--IPKYTYLIE---CENGRRRMLGS	ANEAQKGNQMLVPNHLMTLLYHA	1215
AKE81011	1193	NPI---DFLE---AKGYKE--V-KKDLIIK--LPKYSLFE---LENGRKRMLAS	-GELQKGNELALPSKYVNFYLA	1255
CUO82355	1154	EQI---EYVE--kEEKLSD--VkiIK--Nn-IPLNQLIei-----DGRQYLLITS	--ECVNAMQVLVNEEQCKLIADI	1215
WP_033162887	1155	EQL---SYIAspeHEDLID--VrIVK--E--ILKNQLIei-----DGGLYYVTS	--EYVTARQLSINEQSCCKLISEI	1217
AGZ01981	1210	NPI---DFLE---AKGYKE--V-KKDLIIK--LPKYSLFE---LENGRKRMLAS	-GELQKGNELALPSKYVNFYLA	1272
AKA60242	1177	NPI---DFLE---AKGYKE--V-KKDLIIK--LPKYSLFE---LENGRKRMLAS	-GELQKGNELALPSKYVNFYLA	1239
AKS40380	1177	NPI---DFLE---AKGYKE--V-KKDLIIK--LPKYSLFE---LENGRKRMLAS	-GELQKGNELALPSKYVNFYLA	1239
4UN5_B	1181	NPI---DFLE---AKGYKE--V-KKDLIIK--LPKYSLFE---LENGRKRMLAS	-GELQKGNELALPSKYVNFYLA	1243

WP_010922251	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_039695303	1249	HRAD----NFNS-TEYLN--YVSEHKKEFEKVLSCVEDFANLYVDVE--KNLSKIR-A	VAD-SM---DNFSIEE--	1308
WP_045635197	1247	KNVH----KLDE-PGHLE--YIQKHRNEFKDLLNLVSEFSQKYVLAD--ANLEKIK-S	LYA-DN---EQADIEI--	1306
5AXW_A	979	GELYRVIgVNNDLlNRIE---VNMDITYREYLENMNDKRPPRIIKTiaSKTQSIK-K	LYEvKsk--KHPQIIKkg	1056
WP_009880683	924	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	989
WP_010922251	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_011054416	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_011284745	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_011285506	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_011527619	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_012560673	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_014407541	1239	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1304
WP_020905136	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_023080005	1239	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1304
WP_023610282	1239	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1304
WP_030125963	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_030126706	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_031488318	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_032460140	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_032461047	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_032462016	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_032462936	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_032464890	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_033888930	1065	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1130
WP_038431314	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_038432938	1239	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1304
WP_038434062	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
BAQ51233	1151	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1216
KGE60162	415	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	480
KGE60856	178	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	243
WP_002989955	1240	SHYEKLGsPEDnEQKQL--FVEQHKHYLDEIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
WP_003030002	1222	YQIE---KNYE-PEHRE--YVEKHKDEFKELLEYSVFSRKYVLAD--NNLTKIE-M	AYN-KH---RDKPIREq-	1281
WP_003065552	1250	QRIN----SFNS-TKYLd--YVSAHKKEFEKVLSCVEDFANLYVDVE--KNLSKIR-A	LFS-KN---KDAEVSS--	1309
WP_001040076	1241	SRYNESKgpEEIEKKQE--FVNQHVSyFDDIIFQIINDFSKRVIILAD--ANLEKIN-R	VAD-SM---DNFSIEE--	1306

WP_001040078	1249	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1314
WP_001040080	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040081	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040083	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040085	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040087	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040088	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040089	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040090	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040091	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040092	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYS-DNK--DNTpVDE--	1306
WP_001040094	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_001040095	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_001040096	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_001040097	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_001040098	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_001040099	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_001040100	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_001040104	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040105	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040106	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040107	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040108	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040109	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_001040110	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_015058523	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYS-DNK--DNTpVDE--	1306
WP_017643650	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_017647151	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_017648376	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_017649527	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_017771611	1241	SRYNELKgpEEeiEQKQe--FVwQHVSyFDDiLQlINDFSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_017771984	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
CFQ25032	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
CFV16040	1241	SRYNESKgpEEeiEKKQe--FVNQHVSyFDDiLQlINDFSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306

KLJ37842	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
KLJ72361	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
KLl20707	1255	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1320
KLl42645	1241	SRYNELKgPEEiEQKQe--FVvQHVSyFDDiLQlINDfSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_047207273	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_047209694	1241	SRYNELKgPEEiEQKQe--FVvQHVSyFDDiLQlINDfSNRvILAD--ANLEKIN-K	LYQ-DNK--ENIPVDE--	1306
WP_050198062	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_050201642	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_050204027	1241	SRYNELKgPEEiEQKQe--FVvQHVSyFDDiLQlINDfSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_050881965	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_050886065	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
AHN30376	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYS-DNK--DNTPVDE--	1306
EA078426	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSKRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
CCW42055	1241	SRYNESKgpEEiEKKQe--FVNQHVSyFDDiLQlINDfSNRvILAD--ANLEKIN-K	LYQ-DNK--ENISVDE--	1306
WP_003041502	1254	KRIN----NPIN-KDHIE--YVKKHRDDfKELLNvLEfNEKYvGAT--KNGERLK-E	AVA-DF--DSKSNEE--	1313
WP_037593752	1223	YQIE----KNYE-PEHRE--YVEKHKDEFKELLEyIsvfSRKYvILAD--NNLTkIE-M	LFS-KN--KDAEVSS--	1282
WP_049516684	1223	YRIE----KDYE-PEHRE--YVEKHKDEFKELLEyIsvfSRKYvILAD--NNLTkIE-M	LFS-KN--KDAEVSS--	1282
GAD46167	1222	YQIE----KNYE-PEHRE--YVEKHKDEFKELLEyIsvfSRKYvILAD--NNLTkIE-M	LFS-KN--KDAEVSS--	1281
WP_018363470	1254	HRIG----NFNS-AEHLK--YVSEHKKEFEFVLSvENfANvYVDVE--KNLSKIR-A	AAD-SM--DNFSIEE--	1313
WP_003043819	1249	QNI SATTgSNNLg-----YIEQhREEFKEIfeKIIdfSEKYvILKN--KVNSNLK-S	SFD-EQfavSDSIL--l-	1310
WP_006269658	1222	YRIE----KDYE-PEHRE--YVEKHKDEFKELLEyIsvfSRKYvILAD--NNLTkIE-M	LFS-KN--KDAEVSS--	1281
WP_048800889	1242	HRID----NSDN-SEHLK--YITEHKKEFGKLLSYIErfASLYVDVE--KNISKvK-E	AVE-KI--DSFSVKE--	1301
WP_012767106	1246	-HAHKIEsSKE--LEHEA--YILDhYNDLYQLLSyIERfASLYVDVE--KNISKvK-E	LFS-NI--ESYSISEI-	1308
WP_014612333	1246	-HAHKIEsSKE--LEHEA--YILDhYNDLYQLLSyIERfASLYVDVE--KNISKvK-E	LFS-NI--ESYSISEI-	1308
WP_015017095	1246	-HAHKIEsSKE--LEHEA--YILDhYNDLYQLLSyIERfASLYVDVE--KNISKvK-E	LFS-NI--ESYSISEI-	1308
WP_015057649	1246	-HAHKIEsSKE--LEHEA--YILDhYNDLYQLLSyIERfASLYVDVE--KNISKvK-E	LFS-NI--ESYSISEI-	1308
WP_048327215	1246	-HAHKIEsSKE--LEHEA--YILDhYNDLYQLLSyIERfASLYVDVE--KNISKvK-E	LFS-NI--ESYSISEI-	1308
WP_049519324	1246	-HAHKIEsSKE--LEHEA--YILDhYNDLYQLLSyIERfASLYVDVE--KNISKvK-E	LFS-NI--ESYSISEI-	1308
WP_012515931	1220	AHYHELTgSSEdVLRKKY--FVDRHLHYFDDIiQmINDfAErHILAS--SNLEKIN-H	TYH-NN--SDLPVNER-	1285
WP_021320964	1220	AHYHELTgSSEdVLRKKY--FVERHLHYFDDIiQmINDfAErHILAS--SNLEKIN-H	TYH-NN--SDLPVNER-	1285
WP_037581760	1220	AHYHELTgSSEdVLRKKY--FVERHLHYFDDIiQmINDfAErHILAS--SNLEKIN-H	TYH-NN--SDLPVNER-	1285
WP_004232481	1252	QhVN----NSHK-PEHLN--YVQKHKDEFKDIfnLIiSIArINILKP--KVVDNL--	-IN-EF--TEYQGED--	1308
WP_009854540	1247	HRAD----NFNS-TEyLN--YVSEHKKEFEKvLScVEDfANLYVDVE--KNLSKIR-A	VAD-SM--DNFSIEE--	1306
WP_012962174	1248	HRVN----SFNN-SEHLK--YVSEHKKEfGEVLScVENfAKSvYVDVE--KNLGKIR-A	VAD-KI--DTFSIED--	1307

WP_039695303	1249	HRAD----	NFNS-TEYLN--YVSEHKKEFEKVLSCVEDFANLYVDVE--KNLSKIR-A	VAD-SM---	DNFSIEE--	1308
WP_014334983	1252	HRID----	SFNS-TEHLK--YVSEHKKEFEKVLSCVENFSNLVYDVE--KNLSKVR-A	AAE-SM---	TNFSLEE--	1311
WP_003099269	1240	SHYTKFTgKEEDrEKRS--	YVESHLYYFDEIMQIIVEYSNRYILAD--SNLIKIQ-N	LYK-Ekd---	NFSIEEQ-	1305
AHY15608	1240	SHYTKFTgKEEDrEKRS--	YVESHLYYFXEVKSSF-----	---	-----	1273
AHY17476	1240	SHYTKFTgKEEDrEKRS--	YVESHLYYFX-----	---	-----	1267
ESR09100	72	SHYTKFTgKEEDrEKRS--	YVESHLYYFDEIMQIIVEYSNRYILAD--SNLIKIQ-N	LYK-Ek---	DNFSIEEQ-	137
AGM98575	1240	SHYTKFTgKEEDrEKRS--	YVESHLYYFDVRLSQFRVTNVEF-----	---	-----	1281
ALF27331	1222	KNIH----	KVDE-PKHLd--YVKKHKDEFKELLDVSNFSKKNILAE--SNLEKIE-E	LYA-QN---	NNKDITE--	1281
WP_018372492	1229	KKVDVlvkSKDD--	DYD--LEEHRAEFAELLDCKIKFNDDMYILAS--SNMSKIE-E	IYQ-KNi---	DAPIEE--	1289
WP_045618028	1252	QRIN----	KISE-PIHKQ--YVETHQSEFKELLTAIISLSKKYI-QK--PNVESL---	LQQ-AF---	DQSDKDIYq	1310
WP_045635197	1247	KNVH----	KLDE-PGHLE--YIQKHRNEFKLLDVLVSEFSQYVLAD--ANLEKIK-S	LYA-DN---	EQADIEI--	1306
WP_002263549	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002263887	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002264920	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002269043	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002269448	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002271977	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002272766	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002273241	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002275430	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002276448	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002277050	1230	HHL-----	DN-DYSNE--YVKNHYQQFDILFNEITSFSSKCKLKG--EHIQKIE-E	AYSker---	DSASIEE--	1287
WP_002277364	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002279025	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002279859	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002280230	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002281696	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002282247	1230	HHL-----	DN-DYSNE--YVKNHYQQFDILFNEITSFSSKCKLKG--EHIQKIE-E	AYSker---	DFASIEE--	1287
WP_002282906	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002283846	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002287255	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002288990	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002289641	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281
WP_002290427	1222	KNIH----	KVDE-PKHLd--YVDKHKDEFKELLDVSNFSKKYTLAE--GNLEKIK-E	LYA-QN---	NGEDLKE--	1281

WP_002295753	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_002296423	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_002304487	1236	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1295	LYA-QN--	NGEDLKE--
WP_002305844	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_002307203	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_002310390	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_002352408	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_012997688	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_014677909	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_019312892	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_019313659	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_019314093	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_019315370	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_019803776	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_019805234	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_024783594	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_024784288	1230	HHL-----	DN-DYSNE--	YVKHYYQFDILFNEIT	SFKKCKLKG--	EHIQKIE-E	1287	AYSker---	DFASIEE--
WP_024784666	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_024784894	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_024786433	1230	HHL-----	DN-DYSNE--	YVKHYYQFDILFNEIT	SFKKCKLKG--	EHIQKIE-E	1287	AYSker---	DSASIEE--
WP_049473442	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
WP_049474547	1222	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1281	LYA-QN--	NGEDLKE--
EMC03581	1215	KNIH----	KVDE-PKHLD--	YVDKHKDEFKELLDVVS	NFSKKYTLAE--	GNLEKIK-E	1274	LYA-QN--	NGEDLKE--
WP_000428612	1250	KNIH----	RLDE-PEHLE--	YIQKHRNEFKGLLN	LVSEFSQYVLAD--	ANLEKIK-N	1309	LYA-DN--	EQADIEI--
WP_000428613	1248	KNVH----	KLDE-PEHLE--	YIQKHRNEFKDLN	LVSEFSQYVLAD--	ANLEKIQ-N	1307	LYA-DN--	EQADIEI--
WP_049523028	1243	KRIQ----	KKDE-PEHLE--	YIKQHHSEFN	DLNLFVSEFSQYVLAE--	SNLEKIK-N	1302	LYI-DN--	EQTNMEE--
WP_003107102	1209	SRYT	SFSgKEEDrEKHRH--	FVESHLHYFDEIKDI	IADFSRRYIILAD--	ANLEKIL-T	1274	LYN-EKn--	QFSIEEQ-
WP_054279288	1242	SKRDk--	gTQSEnME---	YISNHKEKFI	EIHYIIRYAENVIKP--	KVIERLN-D	1303	TFNqKF---	NDSDLTEL-
WP_049531101	1252	QRIN----	KISE-PIHKQ--	YVETHQSEFEEL	LTIIISLKKYI-QK--	PIVESL---	1310	LQQ-AF---	EQADKDIYq
WP_049538452	1252	QRIN----	KISE-PIHKQ--	YVEAHQNEFKEL	LTIIISLKKYI-QK--	PNVESL---	1310	LQQ-AF---	EQADKDIYq
WP_049549711	1254	QRIN----	KFSE-PIHKQ--	YVEAHQNEFKEL	LTIIISLKKYI-QK--	PNVESL---	1312	LHQ-AF---	EQADNDIYq
WP_007896501	1246	SRYDKLs	KIESeQKKL--	FVEQHLHYFDEIL	DIVKHTATCYIKAE--	NNLKKIIS	1311	LYK-KK---	EAYSINEq-
EFR44625	1198	SRYDKLs	KIESeQKKL--	FVEQHLHYFDEIL	DIVKHTATCYIKAE--	NNLKKIIS	1263	LYK-KK---	EAYSINEq-
WP_002897477	1247	KNLH----	KLDE-PEHLE--	YIQKHRNEFKDLN	LVSEFSQYIILAE--	ANLEKIK-D	1306	LYA-DN--	EQADIEI--

WP_002906454	1253	QRIN----	KISE-PIHKQ--	YVEAHQNEFKELLTTIISLSKKYI-QK--PNVELL---	LQQ-AF---	DQADKDIYq	1311
WP_009729476	1248	KNVH----	KLDE-PGHLE--	YIQKHRNEFKDLLNLVSEFSQKYVLAD--ANLEKIK-N	LYA-DN---	EQADIEI--	1307
CQR24647	1238	QHAN----	KEDS-----	VI--YLEKRRHELSELFHIIIGVSEKTIKLP--KVENTLN-E	AFE-KHf--	EFDEVSE--	1295
WP_000066813	1252	KNVH----	KLDE-PEHLE--	YIQKHYEFKDLLNLVSEFSQKYVLAD--ANLEKIK-N	LYA-DN---	EQADIEI--	1311
WP_009754323	1248	KNVH----	KLDE-PEHLE--	YIQKHYEFKDLLNLVSEFSQKYVLAE--ANLEKIK-S	LYV-DN---	EQADIEI--	1307
WP_044674937	1241	SNIH----	KITE-PIHLN--	YVNKNKHEFKELLRHISDFSTRYILAQ--DRLSKIE-E	LYD-KN---	DGDDISD--	1300
WP_044676715	1243	SNIH----	KITE-PIHLN--	YVNKNKHEFKELLRHISDFSTRYILAQ--DRLSKIE-E	LYD-KN---	DGDDISD--	1302
WP_044680361	1243	SNIH----	KITE-PIHLN--	YVNKNKHEFKELLRHISDFSTRYILAQ--DRLSKIE-E	LYD-KN---	DGDDISD--	1302
WP_044681799	1241	SNIH----	KITE-PIHLN--	YVNKNKHEFKELLRHISDFSTRYILAQ--DRLSKIE-E	LYD-KN---	DGDDISD--	1300
WP_049533112	1254	KRIN----	NPIN-KDHIE--	YVKKHRDDFKELLNLYLEFNEKYVGAT--KNGERLK-E	AVA-DF--	DSKNEE--	1313
WP_029090905	1182	STKQA----	DE-AMFLKyrrLEHLEAVFEEL--	IRKQAADYQIFE--KLIKKIEVn	FYS-----	c-----TYNEK-	1240
WP_006506696	1212	YNAIYKQ--	DYDNLDDILMI-----	QLYIELTNKMKVLYPAY-rGIAEKFE-S	YVW-----	i-----SKEEK-	1268
AIT42264	1240	SHYEKLKgsPEdneEQQL--	FVEQHKHYLDEIIIEQISEFSKRVILAD--ANLdkVL-S		AYN-KH---	RDkPIREq-	1305
WP_034440723	1218	KHYNE----	DE--TSHK--	FIVEHKAYFDELLNYIVEFANKYLELE--NSIEKIK-D	LYH-----	gKGPdVEEke	1276
AKQ21048	1240	SHYEKLKgsPEdneEQQL--	FVEQHKHYLDEIIIEQISEFSKRVILAD--ANLdkVL-S		AYN-KH---	RDkPIREq-	1305
WP_004636532	1212	NRYDKVK--	----fpDSIE--	YVHDNLAKFDDLLLEYVIDFSNKYINAD--KNVQKIQ-K	IYK-EH---	GTEdVEL--	1271
WP_002364836	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_016631044	1169	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1228
EMS75795	954	QHYDEIAhKESF--	---D--	YVNDHLSEFEILDQVIDFSNRYTIAA--KNTEKIA-E	LFE-QN---	QESTVQS--	1013
WP_002373311	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_002378009	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_002407324	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_002413717	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_010775580	1220	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1279
WP_010818269	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_010824395	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_016622645	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_033624816	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_033625576	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_033789179	1218	KQCLL----	PNQ-SESLA--	YVEQHQPefQeILERVWdFAEVHTLAK--SKVQIIV-K	LFE-AN---	QTADVKE--	1277
WP_002310644	1220	KQYDEIshKESF--	---D--	YVNEHhKefSEVfARVLEFAGKYTLAE--KNIEKLE-K	IYK-EN---	QTDDLAK--	1279
WP_002312694	1221	KQYDEIshKESF--	---D--	YVNEHhKefSEVfARVLEFAGKYTLAE--KNIEKLE-K	IYK-EN---	QTDDLAK--	1280
WP_002314015	1221	KQYDEIshKESF--	---D--	YVNEHhKefSEVfARVLEFAGKYTLAE--KNIEKLE-K	IYK-EN---	QTDDLAK--	1280
WP_002320716	1221	KQYDEIshKESF--	---D--	YVNEHhKefSEVfARVLEFAGKYTLAE--KNIEKLE-K	IYK-EN---	QTDDLAK--	1280

WP_002330729	1220	KQYDEIshKESF-----D--YVNEHHKFESEVFARVLEFAGKYTLAE--KNI EKLE-K	IYK-EN---QTDDLAK--	1279
WP_002335161	1221	KQYDEIshKESF-----D--YVNEHHKFESEVFARVLEFAGKYTLAE--KNI EKLE-K	IYK-EN---QTDDLAK--	1280
WP_002345439	1221	KQYDEIshKESF-----D--YVNEHHKFESEVFARVLEFAGKYTLAE--KNI EKLE-K	IYK-EN---QTDDLAK--	1280
WP_034867970	1211	QHYDKITyQESF-----D--YVNTHLSDFSAILTEVLFAFAEKYTLAD--KNI ERIO-E	LYE-EN---KYGETSM--	1270
WP_047937432	1221	KQYDEIshKESF-----D--YVNEHHKFESEVFARVLEFAGKYTLAE--KNI EKLE-K	IYK-EN---QTDDLAK--	1280
WP_010720994	1211	QHYDKITyQESF-----D--YVNTHLSDFSAILTEVLFAFAEKYTLAD--KNI ERIO-E	LYE-EN---KYGEISM--	1270
WP_010737004	1211	QHYDKITyQESF-----D--YVNTHLSDFSAILTEVLFAFAEKYTLAD--KNI ERIO-E	LYE-EN---KYGETSM--	1270
WP_034700478	1211	QHYDKITyQESF-----D--YVNTHLSDFSAILTEVLFAFAEKYTLAD--KNI ERIO-E	LYE-EN---KYGEISM--	1270
WP_007209003	1209	KKIIN--gKNSD--SVS--YIQNNKFKREIFEYIVDFSSKYISAD--ANLNKIE-K	IFE-NNfh---KASEqe	1269
WP_023519017	1205	RHYDEINhKVSF-----D--YVNAHKEGFNDIFDFISDFGVRYIILAP--QHL EKIK-V	AYE-KN---KEVDLKE--	1264
WP_010770040	1216	KQVDE-----DS-GKSEE--YVREHRAEFAEILNYVQAFSETKILAN--KNLQTIL-K	LYE-EN---KEADIKI--	1274
WP_048604708	1213	KHCNE-----KP-D-SLK--YVTEHQSGFSEIMAHVVKDFAEKYTLVD--KNLEKIL-S	LYA-KN---MDSEVKE--	1270
WP_010750235	1214	NHYDEIAYKDSY-----D--YVNEHFSNFQDILDKVIIFAEKYTSAP--QKLNQII-A	TYE-KN---QEADRKI--	1273
AII16583	1279	SHYEKLKgsPEDnEQQL--FVEQHKHYLDEIEIQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1344
WP_029073316	1226	YKAMKYK-NYDNLDSKII-----NVYDIFVEKCLKLYPTY--kNIATNFE-N	FEN-----i-----SDEEK-	1282
WP_031589969	1226	YKAMKYK-NYDNLDSKII-----DLYRLLINKMELYPEYrkQLVKKFE-D	LKV-----i-----SIEEK-	1283
KDA45870	1200	NAKDG-----EQKLE-----DHKAEFKELFDKIMEFADKYVWAP--KNSEKIR-R	LYE-ENq-----DATPme	1253
WP_039099354	1242	LPLTQ-----SeEIAEQV-----YDEILDQVMHYFFPLYDNTQrAKLSAGKaA	DGN-KMv-----QVGQqv	1306
AKP02966	1238	QIPDE-----DpDQIIaf---YDKNILVEILQELITKMKKFFPFY--KNEQEFLaS	FNQ-----ATTSEk-	1296
WP_010991369	1216	ANCEV-----SD-GKSLD--YIESNREMFaELLAHVSEFAKRYTLAE--ANLNKIN-Q	LFE-QN---KEGDICA--	1274
WP_033838504	1216	ANCEV-----SD-GKSLD--YIESNREMFaELLAHVSEFAKRYTLAE--ANLNKIN-Q	LFE-QN---KEGDICA--	1274
EHN60060	1219	ANCEV-----SD-GKSLD--YIESNREMFaELLAHVSEFAKRYTLAE--ANLNKIN-Q	LFE-QN---KEGDICA--	1277
EFR89594	985	ANCEV-----SD-GKSLD--YIESNREMFaELLAHVSEFAKRYTLAE--ANLNKIN-Q	LFE-QN---KEGDICA--	1043
WP_038409211	1216	KNCEA-----ND-GESLA--YIEMHREMFaELLAHVSEFAKRYTLAN--DRLEKIN-M	FFE-QN---KKGDIKV--	1274
EFR95520	835	KNCEA-----ND-GESLA--YIEMHREMFaELLAHVSEFAKRYTLAN--DRLEKIN-M	FFE-QN---KKGDIKV--	893
WP_003723650	1216	KNCEA-----YIEAHRETFSELLAQVSEFATRYTLAD--ANLSKIN-N	LFE-QN---KEGDICA--	1274
WP_003727705	1216	KNCEA-----YIEAHRETFSELLAQVSEFATRYTLAD--ANLSKIN-N	LFE-QN---KEGDICA--	1274
WP_003730785	1216	KNCEA-----YIEAHRETFSELLAQVSEFATRYTLAD--ANLSKIN-N	LFE-QN---KEGDICA--	1274
WP_003733029	1216	EKYEA-----ID-GESLA--YIEVHRALFDELLAYISEFARKYTLAN--DRLEIN-M	LYE-RN---KDGDKS--	1274
WP_003739838	1216	KNCEA-----SD-GKSLD--YIESNREMFGELLAHVSEFAKRYTLAD--ANLSKIN-Q	LFE-QN---KDN DIKV--	1274
WP_014601172	1216	KNCEA-----SD-GKSLK--YIEAHRETFSELLAQVSEFATRYTLAD--ANLSKIN-N	LFE-QN---KEGDIQA--	1274
WP_023548323	1216	EKREA-----ID-GESLA--YIEAHKAVFGELLAHVSEFAKRYTLAN--DKLDEIN-M	LYE-RN---KDGDKS--	1274
WP_031665337	1216	KNCEA-----SD-GKSLK--YIEAHRETFSELLAQVSEFATRYTLAD--ANLSKIN-N	LFE-QN---KEGDIKA--	1274
WP_031669209	1216	EKYEA-----ID-GESLA--YIEVHRALFDELLAYISEFARKYTLAN--DRLEIN-M	LYE-RN---KDGDKS--	1274

WP_033920898	1216	EKREA-----ID-GESLA--YIEAHKAVFGELLAHISEFARKYTLAN--DKLDEIN-M	LYE-RN---KGDVKS--	1274
AKI42028	1219	KNCEA-----SD-GKSLK--YIEAHRETFSELLAQVSEFATRYTLAD--ANLSKIN-N	LFE-QN---KEGDIQA--	1277
AKI50529	1219	EKREA-----ID-GESLA--YIEAHKAVFGELLAHISEFARKYTLAN--DKLDEIN-M	LYE-RN---KGDVKS--	1277
EFR83390	664	KNCEA-----SD-GKSLK--YTEAHRETFSELLAQVSEFATRYTLAD--ANLSKIN-N	LFE-QN---KEGDIKX--	722
WP_046323366	1216	KNCEA-----SD-GKSLA--YIESHREMFAEILLDSISEFASRYTLAD--ANLEKIN-T	IFE-QN---KSGDVKV--	1274
AKE81011	1256	SHYEKLKgsPEDnEQKQL--FVEQHKHYLDEIIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1321
CUO82355	1216	YNAIYKQ-DFDGLDNMLMi-----QLYLQLIDKLTLYPIY-mGIVEKFE-K	FVS-----i-----SKEEK-	1272
WP_033162887	1218	YAAMLKK-RYEYLDDEEIf-----DLYLQLLQKMDTLYPAY-kGIAKRFF-D	FKN-----i-----DVVEK-	1274
AGZ01981	1273	SHYEKLKgsPEDnEQKQL--FVEQHKHYLDEIIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1338
AKA60242	1240	SHYEKLKgsPEDnEQKQL--FVEQHKHYLDEIIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
AKS40380	1240	SHYEKLKgsPEDnEQKQL--FVEQHKHYLDEIIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1305
4UN5_B	1244	SHYEKLKgsPEDnEQKQL--FVEQHKHYLDEIIIEQISEFSKRVIILAD--ANLDKVL-S	AYN-KH---RDKPIREq-	1309

WP_010922251	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_039695303	1309	ISN---SFI	NLLTTLTALGAP-ADFNFLG--EKI--PRK--R-YTSTKECL	NATLIHQSIITGLYETRIDLSKL--	1369
WP_045635197	1307	LAN---SFI	NLLTFTALGAP-AAFKFFG--KDI--DRK--R-YTTVSEIL	NATLIHQSIITGLYETWIDLKSL--	1367
5AXW A					
WP_009880683	990	-AE---NII	HLFTLTNLGAP-AAFKCFD--TTI--GRN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1049
WP_010922251	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_011054416	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_011284745	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATFIHQSIITGLYETRIDLSQL--	1365
WP_011285506	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_011527619	1306	-AE---NII	HLFTLTNLGAP-TAFKYFD--TTI--DRK--R-YTSTKEVL	DATFIHQSIITGLYETRIDLSQL--	1365
WP_012560673	1306	-AE---NII	HLFTLTNLGAP-AAFKCFD--TTI--GRN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_014407541	1305	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1364
WP_020905136	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_023080005	1305	-AK---NII	HLFTLTNLGAP-AAFKYFD--TTI--ERN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1364
WP_023610282	1305	-AK---NII	HLFTLTNLGAP-AAFKYFD--TTI--ERN--R-YKSIKEVL	DATLIHQSIITGLYEIRIDLSQL--	1364
WP_030125963	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--GRN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_030126706	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_031488318	1306	-AE---NII	HLFTLTNLGAP-AAFIYFD--TTI--GRN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_032460140	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--GRN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_032461047	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--GRN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_032462016	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_032462936	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_032464890	1306	-AE---NII	HLFTLTNLGAP-TAFKYFD--TTI--DRK--R-YTSTKEVL	DATFIHQSIITGLYETRIDLSQL--	1365
WP_033888930	1131	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1190
WP_038431314	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
WP_038432938	1305	-AK---NII	HLFTLTNLGAP-AAFKYFD--TTI--ERN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1364
WP_038434062	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--GRN--R-YKSIKEVL	DATLIHQSIITGLYETRIDLSQL--	1365
BAQ51233	1217	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1276
KGE60162	481	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	540
KGE60856	244	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	303
WP_002989955	1306	-AE---NII	HLFTLTNLGAP-TAFKYFD--TTI--DRK--R-YTSTKEVL	DATFIHQSIITGLYETRIDLSQL--	1365
WP_003030002	1282	LAK---SFI	SLLTFTAFGAP-AAFNFFG--ENI--DRK--R-YTSVTECL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_003065552	1310	ISN---SFI	NLLTTLTALGAP-ADFNFLG--EKI--PRK--R-YTSTKECL	NATLIHQSIITGLYETRIDLSKI--	1370
WP_001040076	1307	LAN---NII	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367

KLJ37842	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
KLJ72361	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
KL20707	1321	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1381
KL42645	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
WP_047207273	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
WP_047209694	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
WP_050198062	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
WP_050201642	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
WP_050204027	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
WP_050881965	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
WP_050886065	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
AHN30376	1307	LAK---	NLFTFTSLGAP-AAFKFFD--KSV--DRK--R-YTSTKEVL	DSTLIHQSIITGLYETRIDLGKL--	1367
EA078426	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
CCW42055	1307	LAN---	NLFTFTSLGAP-AAFKFFD--KIV--DRK--R-YTSTKEVL	NSTLIHQSIITGLYETRIDLGKL--	1367
WP_003041502	1314	ICT---	GLFELTSLGSA-SDFEFLG--VKI--PRY--RdYTPSSLLK	DSTLIHQSIITGLYETRIDLSKL--	1383
WP_037593752	1283	LAK---	SLLTFTAFGAP-AAFNFFG--ENI--DRK--R-YTSVTECL	NATLIHQSIITGLYETRIDLSKL--	1343
WP_049516684	1283	LAK---	SLLTFTAFGAP-AAFNFFG--ENI--DRK--R-YTSVTECL	NATLIHQSIITGLYETRIDLSKL--	1343
GAD46167	1282	LAK---	SLLTFTAFGAP-AAFNFFG--ENI--DRK--R-YTSVTECL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_018363470	1314	ISD---	LLLTALGAP-ADFNFLG--EKI--PRK--R-YNSTKECL	NATLIHQSIITGLYETRIDLSKL--	1374
WP_003043819	1311	SN---	SLLKYTSFGAS-GGTFDLD--LDVkgGRL--R-YQTVTEVL	DATLIYQSIITGLYETRIDLSQL--	1372
WP_006269658	1282	LAK---	SLLTFTAFGAP-AAFNFFG--ENI--DRK--R-YTSVTECL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_048800889	1302	ISN---	HLTLTALGAP-ADFNFLG--EKI--PRK--R-YTSTKECL	NATLIHQSIITGLYETQTDLSKL--	1362
WP_012767106	1309	CS---	LLTLTASGAP-ADFKFLG--TTI--PRK--R-YGSPQSIL	SSTLIHQSIITGLYETRIDLSQL--	1368
WP_014612333	1309	CS---	LLTLTASGAP-ADFKFLG--TTI--PRK--R-YGSPQSIL	SSTLIHQSIITGLYETRIDLSQL--	1368
WP_015017095	1309	CS---	LLTLTASGAP-ADFKFLG--TTI--PRK--R-YGSPQSIL	SSTLIHQSIITGLYETRIDLSQL--	1368
WP_015057649	1309	CS---	LLTLTASGAP-ADFKFLG--TTI--PRK--R-YGSPQSIL	SSTLIHQSIITGLYETRIDLSQL--	1368
WP_048327215	1309	CS---	LLTLTASGAP-ADFKFLG--TTI--PRK--R-YGSPQSIL	SSTLIHQSIITGLYETRIDLSQL--	1368
WP_049519324	1309	CS---	LLTLTASGAP-ADFKFLG--TTI--PRK--R-YGSPQSIL	SSTLIHQSIITGLYETRIDLSQL--	1368
WP_012515931	1286	AE---	NVFTFVALGAP-AAFKFFD--ATI--DRK--R-YTSTKEVL	NATLIHQSVTGLYETRIDLSQL--	1345
WP_021320964	1286	AE---	NVFTFVALGAP-AAFKFFD--ATI--DRK--R-YTSTKEVL	NATLIHQSVTGLYETRIDLSQL--	1345
WP_037581760	1286	AE---	NVFTFVALGAP-AAFKFFD--ATI--DRK--R-YTSTKEVL	NATLIHQSVTGLYETRIDLSQL--	1345
WP_004232481	1309	ISSlseSFI	LLKFI SF GAP-GAFKFLK--LDV--KQSn1R-YKSTTEAL	SATLIHQSVTGLYETRIDLSKL--	1374
WP_009854540	1307	ISN---	LLTLTALGAP-ADFNFLG--EKI--PRK--R-YTSTKECL	TATLIHQSIITGLYETRIDLSKL--	1367
WP_012962174	1308	ISI---	LLTLTALGAP-ADFNFLG--EKI--PRK--R-YTSTKECL	NATLIHQSIITGLYETRIDLSKL--	1368

WP_039695303	1309	ISN---SFI	NLTLTALGAP-ADFNFLG--EKI--PRK--R-YTSTKECL	NATLIHQSIITGLYETRIDLSKL--	1369
WP_014334983	1312	ISA---SFI	NLTLTALGAP-ADFNFLG--EKI--PRK--R-YTSTKECL	SATLIHQSVTGLYETRIDLSKL--	1372
WP_003099269	1306	-AI---NML	NLFTFDLGA-PAFKAFFN--GDI--DRK--R-YSSTNEII	NSTLIYQSPTGLYETRIDLSKL--	1365
AHY15608					
AHY17476					
ESR09100	138	-AI---NML	NLFTFDLGA-PAFKAFFNg--DI--DRK--R-YSSTNEII	NSTLIYQSPTGLYETRIDLSKL--	197
AGM98575					
ALF27331	1282	LAS---SFI	NLLTFTAIGAP-AAFKFFD--NNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_018372492	1290	VAR---SFV	-LLNFTMMGAA-TDFKFFG--QII--PRK--R-YPSTTECL	KSTLIHQSVTGLYETRIDLSKL--	1350
WP_045618028	1311	LSE---SFI	SLLKLI SF GAP-GTFKFLG--VEI--SQSnvR-YQSVSSCF	NATLIHQSIITGLYETRIDLSKL--	1373
WP_045635197	1307	LAN---SFI	NLLTFTAIGAP-AAFKFFG--KDI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETWIDLSKL--	1367
WP_002263549	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLNLK--	1342
WP_002263887	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLNLK--	1342
WP_002264920	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002269043	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLNLK--	1342
WP_002269448	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002271977	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002272766	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002273241	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002275430	1282	LSS---SFI	NLLTFTAIGAP-AAFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002276448	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002277050	1288	LAD---GFI	KLLGFTQLGAT-SPFSFLG--IKL--NQK--Q-YTGKKDYL	EATLIHQSIITGLYETRIDLNLK--	1352
WP_002277364	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002279025	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002279859	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002280230	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002281696	1282	LSS---SFI	NLLTFTAIGAP-AAFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002282247	1288	LAD---GFI	KLLGFTQLGAT-SPFSFLG--IKL--NQK--Q-YTGKKDYL	EATLIHQSIITGLYETRIDLSKL--	1352
WP_002282906	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002283846	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002287255	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002288990	1282	LAS---SFI	NLLTFTAIGAP-AAFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002289641	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342
WP_002290427	1282	LAS---SFI	NLLTFTAIGAP-ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSIITGLYETRIDLSKL--	1342

WP_002295753	1282	LAS---SFI	LLLTFTAI GAP -AAFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_002296423	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_002304487	1296	LAS---SFI	LLLTFTAI GAP -AAFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLNKL--	1356
WP_002305844	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_002307203	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_002310390	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_002352408	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_012997688	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_014677909	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_019312892	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_019313659	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLNKL--	1342
WP_019314093	1282	LAS---SFI	LLLTFTAI GAP -AAFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_019315370	1282	LSS---SFI	LLLTFTAI GAP -AAFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_019803776	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLNKL--	1342
WP_019805234	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_024783594	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLNKL--	1342
WP_024784288	1288	LAD---GFI	KLGGFTQLGAT -SPFSFLG--IKL--NQK--Q-YTGKDDYL	EATLIHQSI TGLYETRIDLSKL--	1352
WP_024784666	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_024784894	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_024786433	1288	LAD---GFI	KLGGFTQLGAT -SPFSFLG--IKL--NQK--Q-YTGKDDYL	EATLIHQSI TGLYETRIDLSKL--	1352
WP_049473442	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
WP_049474547	1282	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1342
EMC03581	1275	LAS---SFI	LLLTFTAI GAP -ATFKFFD--KNI--DRK--R-YTSTTEIL	NATLIHQSI TGLYETRIDLSKL--	1335
WP_000428612	1310	LAN---SFI	LLLTFTALGAP -AAFKFFG--KDV--DRK--R-YTTVSEIL	NATLIHQSI TGLYETRIDLSKL--	1370
WP_000428613	1308	LAN---SFI	LLLTFTALGAP -AAFKFFG--KDI--DRK--R-YTTVSEIL	NATLIHQSI TGLYETRIDLSKL--	1368
WP_049523028	1303	IAN---SFI	LLLTFTAFGAP -AVFKFFG--KDI--ERK--R-YSTVTEIL	KATLIHQSL TGLYETRIDLSKL--	1363
WP_003107102	1275	-AT---NML	NLFTFTGLGAP -ATLKFFN--VDI--DRK--R-YTSSTEIL	NSTLIHQSI TGLYETRIDLSKI--	1334
WP_054279288	1304	-SI---SFL	NLFKFTSFGAP -EKFTFLN--SEIkqDDV--R-YRSTKECL	NSTLIHQSV TGLYETRIDLSQF--	1365
WP_049531101	1311	LSE---SFI	SLLKLT SFGAP -GAFRFLG--VEI--SQSsvR-YQSVSSCF	NATLIHQSI TGLYETRIDLSKL--	1373
WP_049538452	1311	LSE---SFI	SLLKLT SFGAP -GAFKFLG--VEI--SQSsvR-YKPN SQFL	DATLIHQSI TGLYETRIDLSKL--	1373
WP_049549711	1313	LSE---SFI	SLLKLT SFGAP -GAFKFLG--AEI--SQSsvR-YKPN SQFL	DTTLIHQSI TGLYETRIDLSKL--	1375
WP_007896501	1312	-AL---NML	NLFI FTSLGAP -STFVFFD--ETI--DRK--R-YTTSSDVL	NGILIQSITGLYETRIDLSRF--	1371
EFR44625	1264	-AL---NML	NLFI FTSLGAP -STFVFFD--ETI--DRK--R-YTTSSDVL	NGILIQSITGLYETRIDLSRF--	1323
WP_002897477	1307	LAN---SFI	LLLTFTALGAP -AAFKFFG--KDV--DRK--R-YTTVSEIL	NATLIHQSI TGLYETRIDLSKL--	1367

WP_002906454	1312	LSE---SFI	SLKLTSTFGAP-GAFKFLG--VEI--SQSsvr-YKPNsqFL	DTTLIHQSITGLYETRIDLskL--	1374
WP_009729476	1308	LAN---SFI	NLLTFTALGAP-AAFKFFG--KDV--DRK--R-YTTVSEIL	NATLIHQSVTGLYETRIDLskL--	1368
CQR24647	1296	LAQ---SFI	SLLKFTAFGAP-GGFKFLD--ADI--KQsnLR-YQTVTEVL	SSTLIHQSVTGLYETRIDLskL--	1358
WP_000066813	1312	LAN---SFI	NLLTFTALGAP-AAFKFLG--KDV--DRK--R-YTTVSEIL	NATLIHQSVTGLYETRIDLskL--	1372
WP_009754323	1308	LAN---SFI	NLLTFTALGAP-AAFKFFG--KDV--DRK--R-YTTVSEIL	NATLIHQSVTGLYETRIDLskL--	1368
WP_044674937	1301	LTS---SFV	NLLTFTAIGAP-AAFKFLG--SVI--DRK--R-YTSIAEIL	EATLIHQSVTGLYETRIDLskL--	1361
WP_044676715	1303	LTS---SFV	NLLTFTAIGAP-AAFKFLG--SVI--DRK--R-YTSIAEIL	EATLIHQSVTGLYETRIDLskL--	1363
WP_044680361	1303	LTS---SFV	NLLTFTAIGAP-AAFKFLG--SVI--DRK--R-YTSIAEIL	EATLIHQSVTGLYETRIDLskL--	1363
WP_044681799	1301	LTS---SFV	NLLTFTAIGAP-AAFKFLG--SVI--DRK--R-YTSIAEIL	EATLIHQSVTGLYETRIDLskL--	1361
WP_049533112	1314	ICT---SFL	GLFELTSLGSA-SDFEFLG--VKI--PRY--RdYTPSSLLK	DSTLIHQSVTGLYETRIDLskL--	1383
WP_029090905	1241	-VK---VI	ELLKITQANATnGDLKLLK---M-sNReg-R-LGSVSVAl	DFKIINQSVTGLYQSIEdYNN--	1300
WP_006506696	1269	-AN---II	QMLIVMHRGPQnGNIvYDDf--KI--sDRIg-R-LKTKNHNL	NIVFISQSPtGIYTKKYKL-----	1329
AIT42264	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSVTGLYETRIDLskL--	1365
WP_034440723	1277	LVE---SFI	NLLAITKCGPA-ADITFLG--EKI--SRK--R-YRSTNCLW	GSEVIFQSPtGLYETRLRL-----	1335
AKQ21048	1306	-AE---NII	HLFTLTNLGAP-AAFKYFD--TTI--DRK--R-YTSTKEVL	DATLIHQSVTGLYETRIDLskL--	1365
WP_004636532	1272	TVE---SFV	NLMFTAMGAP-ATFKFYG--ESI--TRS--R-YTSITEFR	GSTLIHQSVTGLYETRYKL-----	1329
WP_002364836	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_016631044	1229	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1286
EMS75795	1014	LSQ---SFI	NLMQLNAMGAP-ADFKFFD--VII--PRK--R-YPSLITEIW	ESTIIYQSVTGLRETRRMATLwd	1076
WP_002373311	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_002378009	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_002407324	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_002413717	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_010775580	1280	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1337
WP_010818269	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_010824395	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_016622645	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_033624816	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_033625576	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_033789179	1278	IAA---SFI	QLMQFNAMGAP-STFKFFQ--KDI--ERA--R-YTSIKEIF	DATIIYQSTTGLYETRRKV-----	1335
WP_002310644	1280	LAS---SFV	NLMQFNAMGAP-ADFKFFD--VTI--PRK--R-YTSLITEIW	QSTIIHQSVTGLYETRRKV-----	1339
WP_002312694	1281	LAS---SFV	NLMQFNAMGAP-ADFKFFD--VTI--PRK--R-YTSLITEIW	QSTIIHQSVTGLYETRRKV-----	1340
WP_002314015	1281	LAS---SFV	NLMQFNAMGAP-ADFKFFD--VTI--PRK--R-YTSLITEIW	QSTIIHQSVTGLYETRRKV-----	1340
WP_002320716	1281	LAS---SFV	NLMQFNAMGAP-ADFKFFD--VTI--PRK--R-YTSLITEIW	QSTIIHQSVTGLYETRRKV-----	1340

WP_002330729	1280	LAS---SfV	NLMQFNAMGAP-ADFKFFD-VTI--PRK--R-YTSLTEIW	QSTIIHQSIITGLYETRIRMGK---	1339
WP_002335161	1281	LAS---SfV	NLMQFNAMGAP-ADFKFFD-VTI--PRK--R-YTSLTEIW	QSTIIHQSIITGLYETRIRMGK---	1340
WP_002345439	1281	LAS---SfV	NLMQFNAMGAP-ADFKFFD-VTI--PRK--R-YTSLTEIW	QSTIIHQSIITGLYETRIRMGK---	1340
WP_034867970	1271	IAQ---SfL	QLLQFNAGAP-ADFKFFG-VTI--PRK--R-YTSLTEIW	DATIIYQSVTGLYETRIRMGDLwa	1333
WP_047937432	1281	LAS---SfV	NLMQFNAMGAP-ADFKFFD-VTI--PRK--R-YTSLTEIW	QSTIIHQSIITGLYETRIRMGK---	1340
WP_010720994	1271	IAQ---SfL	QLLQFNAGAP-ADFKFFG-VTI--PRK--R-YTSLTEIW	DATIIYQSVTGLYETRIRMGDLwa	1333
WP_010737004	1271	IAQ---SfL	QLLQFNAGAP-ADFKFFG-VTI--PRK--R-YTSLTEIW	DATIIYQSVTGLYETRIRMGDLwa	1333
WP_034700478	1271	IAQ---SfL	QLLQFNAGAP-ADFKFFG-VTI--PRK--R-YTSLTEIW	DATIIYQSVTGLYETRIRMGDLwa	1333
WP_007209003	1270	IAK---SfI	NLLTFTAMGAP-ADFEFFG-EKI--PRK--R-YVSISEII	DAVFIHQSIITGLYETRIRLTEV--	1330
WP_023519017	1265	MID---AIL	SLLKFTLFGAS-VEFKFFD-IKI--LK--R-YKSLTDIWI	EATIIYQSVTGLYERRVEVRKLwd	1326
WP_010770040	1275	IAE---SfV	NLMKFSAYGAP-MDFKFFG-KTI--PRS--R-YTSVGEILL	SATIIINQSIITGLYETRRKL-----	1332
WP_048604708	1271	IAQ---SfV	DLMQFNAGAP-ADFKFFG-ETI--PRK--R-YTSVNEILL	EATIIINQSIITGLYETRRRL-----	1328
WP_010750235	1274	MAH---SfV	NLMQFNALGAP-ADFKFFD-TTI--TRK--R-YTSLTEIW	QSTIIYQSVTGLYETRRRMADLwd	1336
AI116583	1345	-AE---NII	HLFTLTNLGAP-AAFKYFD-TTI--DRK--R-YTSTKEVL	DATLIHQSIITGLYETRIDLSQL--	1404
WP_029073316	1283	-CE---VI	QMLVVMHAGPQnGNITFDDf--KL--sNRLg--R-LNCKTISL	TTVFIADSPtGMYSKKYKL-----	1343
WP_031589969	1284	-CN---II	QIILATLHCNSSiGKIMYSDF--KI--sTTIgr--LNGRTISL	DISFIAESPtGMYSKKYKL-----	1344
KDA45870	1254	LGK---NFV	ELLRYTDGAA-SDFKFFG-ENI--PRK--R-YNSAGSLL	NGTLIYQSKTGLYETRIDLGLK--	1314
WP_039099354	1307	ILDr---V	-LIGLHANAaV-SDLGVLKisTPL--GKM--Q---QPSGIS	DTQIIYQSPtGLFERRVALRDL--	1368
AKP02966	1297	INSl-eELI	TLLHANSTSAH-LIFNNIE-kKAF--GRK-----THGLT	DTDFIYQSVTGLYETRIHIE----	1356
WP_010991369	1275	IAQ---SfV	DLMAFNAMGAP-ASFkFFE-TTI--ERK--R-YNNLKELL	NSTIIYQSIITGLYESRKRl-----	1332
WP_033838504	1275	IAQ---SfV	DLMAFNAMGAP-ASFkFFE-TTI--ERK--R-YNNLKELL	NSTIIYQSIITGLYESRKRl-----	1332
EHN60060	1278	IAQ---SfV	DLMAFNAMGAP-ASFkFFE-TTI--ERK--R-YNNLKELL	NSTIIYQSIITGLYESRKRl-----	1335
EFR89594	1044	IAQ---SfV	DLMAFNAMGAP-ASFkFFE-TTI--ERK--R-YNNLKELL	NSTIIYQSIITGLYESRKRl-----	1332
WP_038409211	1275	IAK---SfD	DLMAFNAMGAP-ASFkFFE-TTI--ERK--R-YNNLKELL	NSTIIYQSIITGLYESRKRl-----	1101
EFR95520	894	IAK---SfD	KLKVFNAFGAP-RDFEFfE-TTI--KRK--R-YNIKELL	NATIIYQSIITGLYEARKRL-----	1332
WP_003723650	1275	IAQ---SfV	KLKVFNAFGAP-RDFEFfE-TTI--KRK--R-YNIKELL	NATIIYQSIITGLYEARKRL-----	951
WP_003727705	1275	IAQ---SfV	DLMAFNAMGAP-ASFkFFE-ATI--DRK--R-YTNLKELL	SSTIIYQSIITGLYESRKRl-----	1332
WP_003730785	1275	IAQ---SfV	DLMAFNAMGAP-ASFkFFE-ATI--DRK--R-YTNLKELL	SSTIIYQSIITGLYESRKRl-----	1332
WP_003733029	1275	IAE---SfV	DLMAFNAMGAP-ASFkFFE-ATI--DRK--R-YTNLKELL	SSTIIYQSIITGLYESRKRl-----	1332
WP_003739838	1275	IAQ---SfV	SLKKFNafGVH-QDFsFFG-TKI--ERK--R-DRKLNELL	NSTIIYQSIITGLYESRKRl-----	1332
WP_014601172	1275	IAQ---SfV	NLMAFNAMGAP-ASFkFFE-ATI--ERK--R-YTNLKELL	SATIIYQSIITGLYEARKRL-----	1332
WP_023548323	1275	IAE---SfV	DLMAFNAMGAP-ASFkFFE-ATI--DRK--R-YTNLKELL	SSTIIYQSIITGLYESRKRl-----	1332
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WP_000066813	1373	GED	1375
WP_009754323	1369	GED	1371
WP_044674937	1362	GGD	1364
WP_044676715	1364	GGD	1366
WP_044680361	1364	GGD	1366
WP_044681799	1362	GGD	1364
WP_049533112	1384	GED	1386
WP_029090905	---	---	---
WP_006506696	---	---	---
AIT42264	1366	GGD	1389
WP_034440723	---	---	---
AKQ21048	1366	GGD	1384
WP_004636532	1330	-ED	1332
WP_002364836	1336	-VD	1337
WP_016631044	1287	-VD	1288
EMS75795	1077	GEQ	1079
WP_002373311	1336	-VD	1337
WP_002378009	1336	-VD	1337
WP_002407324	1336	-VD	1337
WP_002413717	1336	-VD	1337
WP_010775580	1338	-VD	1339
WP_010818269	1336	-VD	1337
WP_010824395	1336	-VD	1337
WP_016622645	1336	-VD	1337
WP_033624816	1336	-VD	1337
WP_033625576	1336	-VD	1337
WP_033789179	1336	-VD	1337
WP_002310644	---	---	---
WP_002312694	---	---	---
WP_002314015	---	---	---
WP_002320716	---	---	---

WP_002330729	---			
WP_002335161	---			
WP_002345439	---			
WP_034867970	GEQ	1334	1336	1336
WP_047937432	---			
WP_010720994	GEQ	1334	1336	1336
WP_010737004	GEQ	1334	1336	1336
WP_034700478	GEQ	1334	1336	1336
WP_007209003	---			
WP_023519017	GER	1327	1330	1330
WP_010770040	-VD	1333	1334	1334
WP_048604708	-GD	1329	1330	1330
WP_010750235	GVQ	1337	1339	1339
AI116583	GGD	1405	1424	1424
WP_029073316	---			
WP_031589969	---			
KDA45870	---			
WP_039099354	---			
AKP02966	---			
WP_010991369	-DD	1333	1334	1334
WP_033838504	-DD	1333	1334	1334
EHN60060	-DD	1336	1337	1337
EFR89594	-DD	1102	1103	1103
WP_038409211	-ED	1333	1334	1334
EFR95520	-ED	952	953	953
WP_003723650	-DD	1333	1334	1334
WP_003727705	-DD	1333	1334	1334
WP_003730785	-DD	1333	1334	1334
WP_003733029	-DN	1333	1334	1334
WP_003739838	-DG	1333	1334	1334
WP_014601172	-DD	1333	1334	1334
WP_023548323	-DS	1333	1334	1334
WP_031665337	-DD	1333	1334	1334
WP_031669209	-DN	1333	1334	1334

WP_033920898	1333	-DS	1334
AKI42028	1336	-DD	1337
AKI50529	1336	-DS	1337
EFR83390	781	-DD	782
WP_046323366	1333	-DD	1334
AKE81011	1382	GGD	1400
CUO82355	---	---	
WP_033162887	---	---	
AGZ01981	1399	GGD	1417
AKA60242	1366	GGD	1368
AKS40380	1366	GGD	1376
4UN5_B	1370	GGD	1372

Table 2. T to C changes with NGG PAM. Table 2 shows a list of T to C mutations that may be corrected using any of the base editors provided herein. GRNAs and gRNAall indicate the protospacer and PAM sequence, where the PAM sequence is the last 3 nucleotides of each of the sequences in GRNAs and gRNAall.

Name	Gene ID	Gene Sympo l	SEQ ID NO:	Flanks	SEQ ID NO:	GRNAs	SEQ ID NO:	gRNAall
NM_000071.2(CBS):c.833T>C (p.Ile278Thr)	875	CBS	2540	['CTGAAGCCGC GCCCTCTGCAG ATCAYTGGGGT GGATCCCGAA GGGTCCATC']	2703 - 2704	['ATCAYTGGGGTG GATCCCGAAGG', 'TCAYTGGGGTGGGA TCCCGAAGGG']	2907 - 2908	['ATCAYTGGGGTG GATCCCGAAGG', 'TCAYTGGGGTGG ATCCCGAAGGG']
NM_001385.2(DPYS):c.1078T>C (p.Trp360Arg)	1807	DPYS	2541	['TGTTGAAGAT CGGATGTCCGT AATAYGGGAA AAAGGCGTGG TGGGTTTCAC']	2705 - 2707	['CGTAATAYGGGA AAAAGGCGTGG', 'AATAYGGGAAAA AGGCGTGGTGG', 'ATAYGGGAAAAA GGCCTGGTGGG']	2909 - 2911	['CGTAATAYGGGA AAAAGGCGTGG', 'AATAYGGGAAAA AGGCGTGGTGG', 'ATAYGGGAAAAA GGCCTGGTGGG']
NM_000027.3(AGA):c.916T>C (p.Cys306Arg)	175	AGA	2542	['TCCAGAATTC TTTGGGGCTGT TATAYGTGCCA ATGTGACTGGA AGTTACGG']	2708	['GTTATAYGTGCC AATGTGACTGG']	2912	['GTTATAYGTGCC AATGTGACTGG']
NM_000035.3(ALDOB):c.442T>C (p.Trp148Arg)	229	ALDO B	2543	['GAAAGATGGT GTTGACTTTGG GAAGYGGCGT GCTGTGCTGAG GATTGCCGA']	2709	['GGAAGYGGCGTG CTGTGCTGAGG']	2913	['GGAAGYGGCGT GCTGTGCTGAGG']
NM_173560.3(RFX6):c.380+2T>C	2225 46	RFX6	2544	['GCAGACACAG CTCACGCTGCA GTGGYGAGAC TCGCCCGCAGG GTACACTGA']	2710 - 2711	['CAGTGGYGAGAC TCGCCCGCAGG', 'AGTGGYGAGACTC GCGCCGCAGGG']	2914 - 2915	['CAGTGGYGAGAC TCGCCCGCAGG', 'AGTGGYGAGACT CCGCCCGCAGGG']
NM_153704.5(TM67):c.1843T>C (p.Cys615Arg)	9114 7	TM67	2545	['AGAACGTTTT GTCACCTATGT TGGAHGTGCCT TTGCTCTGAAG GTAAGTTT']	2712	['TGGAHGTGCCTTT GCTCTGAAGG']	2916	['TGGAHGTGCCTT TGCTCTGAAGG']

NM_000124.3(ERCC6):c.2960 T>C (p.Leu987Pro)	2074	ERCC6	2546	['AAGCAGTTTT TGACAAATAG AGTGCYAAAA GACCCAAAAC AAAGGCGGTTT ']	2713	['TGCYAAAAGACC CAAACAAAGG']	2917	['TGCYAAAAGACC CAAACAAAGG']
NM_020435.3(GJC2):c.857T> C (p.Met286Thr)	5716 5	GJC2	2547	['TGCCTGCTGC TCAACCTCTGT GAGAYGGCCC ACCTGGGCTTG GGCAGCGCG']	2714	['TGAGAYGGCCCA CCTGGGCTTGG']	2918 - 2919	['TGAGAYGGCCCA CCTGGGCTTGG', 'GAGAYGGCCCAC CTGGGCTTGGG']
NM_000920.3(PC):c.434T>C (p.Val145Ala)	5091	PC	2548	['CGGTTTATTG GGCCAAGCCC AGAAGBGGTC CGCAAGATGG GAGACAAGGT G']	2715	['CCAGAAGBGGTC CGCAAGATGGG']	2920	['CCAGAAGBGGTC CGCAAGATGGG']
NM_000026.2(ADSL):c.674T >C (p.Met225Thr)	158	ADSL	2549	['TCCAAGGTAG AGCAGCTTGAC AAGAYGGTGA CAGAAAAGGC AGGATTTAAG']	2716	['AAGAYGGTGACA GAAAAGGCAGG']	2921	['AAGAYGGTGAC AGAAAAGGCAGG']
NM_000391.3(TPP1):c.1093T >C (p.Cys365Arg)	1200	TPP1	2550	['TCTCTCAGGT GACAGTGGGG CCGGGYGTTG GTCTGTCTCTG GAAGACACCA']	2717	['GCCGGGYGTTGG TCTGTCTCTGG']	2922	['GCCGGGYGTTGG TCTGTCTCTGG']
NM_004183.3(BEST1):c.704T >C (p.Val235Ala)	7439	BEST1	2551	['TACGACTGGA TTAGTATCCCA CTGGYGTATAC ACAGGTGAGG ACTAGGCTG']	2718	['CACTGGYGTATA CACAGGTGAGG']	2923	['CACTGGYGTATA CACAGGTGAGG']
NM_000019.3(ACAT1):c.935 T>C (p.Ile312Thr)	38	ACAT 1	2552	['CTCAATGTTA CACCCTGGCA AGAAYAGTAG GTAAGGCCAG GCGAGGTGGC']	2719	['CAAGAAYAGTAG GTAAGGCCAGG']	2924	['CAAGAAYAGTA GGTAAGGCCAGG']
NM_000543.4(SMPD1):c.911 T>C (p.Leu304Pro)	6609	SMPD 1	2553	['CGGGCCCTGA CCACCGTCACA GCACYTGTGA GGAAGTTCTG GGGCCAGTG']	2720	['CACYTGTGAGGA AGTTCCTGGGG']	2925 - 2927	['AGCACYTGTGAG GAAGTTCCTGG', 'GCACYTGTGAGG AAGTTCCTGGG', 'CACYTGTGAGGA AGTTCCTGGGG']
NM_000527.4(LDLR)	3949	LDLR	2554	['ACAAATCTGA	2721	['CGGYATGGGCGG	2928	['ACTGCGGYATGG

LDLR):c.694+2 T>C				CGAGGAAAAC TGCGGYATGG GCGGGGCCAG GGTGGGGGCG G']		GGCCAGGGTGG']	- 2930	GCGGGGCCAGG', 'CTGCGGYATGGG CGGGGCCAGGG', 'CGGYATGGGCGG GGCCAGGGTGG']
NM_012464.4(TLL1):c.713T> C (p.Val238Ala)	7092	TLL1	2555	['AAGAACTGTG ATAAATTTGGG ATTGYTGTTCA TGAATTGGGTC ATGTGATA']	2722	['GGGATTGYTGTT ATGAATTGGG']	2931	['GGGATTGYTGTT CATGAATTGGG']
NM_000112.3(SLC26A2):c.- 26+2T>C	1836	SLC26 A2	2556	['CCTGCAGCGG CCCGGACCCG AGAGGYGAGA AGAGGGAAGC GGACCAGGGA A']	2723	['GAGAGGYGAGAA GAGGGAAGCGG']	2932	['GAGAGGYGAGA AGAGGGAAGCGG']
NM_00100574 1.2(GBA):c.751 T>C (p.Tyr251His)	2629	GBA	2557	['CATCTACCAC CAGACCTGGG CCAGAYACTTT GTGAAGTAAG GGATCAGCAA']	2724	['GCCAGAYACTTT GTGAAGTAAGG']	2933 - 2934	['GCCAGAYACTTT GTGAAGTAAGG', 'CCAGAYACTTTGT GAAGTAAGGG']
NM_020365.4(EIF2B3):c.1037 T>C (p.Ile346Thr)	8891	EIF2B3	2558	['CCACCAGTCC ATTCGTCAGCC CAGAYTGTCAG GCAAACACCT GGTAAGTGCT']	2725	['CCAGAYTGTCAG CAAACACCTGG']	2935	['CCAGAYTGTCAG CAAACACCTGG']
NM_022041.3(GAN):c.1268T >C (p.Ile423Thr)	8139	GAN	2559	['TGCTATGCAG CTATGAAAAA GAAAAYCTAC GCCATGGGTG GAGGCTCCTAC ']	2726	['AAGAAAAYCTAC GCCATGGGTGG']	2936 - 2937	['AAGAAAAYCTAC GCCATGGGTGG', 'AAAAYCTACGCC ATGGGTGGAGG']
NM_054027.4(ANKH):c.143T >C (p.Met48Thr)	5617 2	ANKH	2560	['GCTGTCAAGG AGGATGCAGT CGAGAYGCTG GCCAGCTACG GGCTGGCGTAC ']	2727 - 2728	['GTCGAGAYGCTG GCCAGCTACGG', 'TCGAGAYGCTGGC CAGCTACGGG']	2938 - 2939	['GTCGAGAYGCTG GCCAGCTACGG', 'TCGAGAYGCTGG CCAGCTACGGG']
NM_006329.3(FBLN5):c.506T >C (p.Ile169Thr)	1051 6	FBLN5	2561	['TTGCTTGCAT TTCTGTTTCCA GACAYTGATG AATGTCGCTAT GGTTACTGC']	2729	['GACAYTGATGAA TGTCGCTATGG']	2940	['GACAYTGATGAA TGTCGCTATGG']
NM_004086.2(-1	-1	-	2562	['GCACCTCTGG	2730	['AGATAYGGCTTC	2941	['AGATAYGGCTTC

COCH):c.1535 T>C (p.Met512Thr)				ATGACCTGAA AGATAYGGCTT CTAAACCGAA GGAGTCTCAT']		TAAACCGAAGG']		TAAACCGAAGG']
NM_002942.4(ROBO2):c.283 4T>C (p.Ile945Thr)	6092	ROBO 2	2563	['AATAGCAACA GTGGCCCAAAT GAGAYTGGA ATTTTGGCCGT GGAGGTAAG']	2731	['GAGAYTGGA ATTTTGGCCGTGG']	2942	['GAGAYTGGA ATTTTGGCCGTGG']
NM_001300.5(KLF6):c.190T> C (p.Trp64Arg)	1316	KLF6	2564	['CAAATTTGAC AGCCAGGAAG ATCTGYGGACC AAAATCATTCT GGCTCGGGA']	2732	['TCTGYGGACCA AATCATTCTGG']	2943	['TCTGYGGACCA AATCATTCTGG']
NM_030653.3(DDX11):c.2271 +2T>C	1663	DDX11	2565	['CTGGCATATT CCAGGTGCATC CAGGYGCGGG CGTCATGCTGG GCTTGGGTC']	2733	['TCCAGGYGCGGG CGTCATGCTGG']	2944 - 2945	['TCCAGGYGCGGG CGTCATGCTGG', 'CCAGGYGCGGGC GTCATGCTGGG']
NM_001451.2(FOXF1):c.1138 T>C (p.Ter380Arg)	2294	FOXF1	2566	['CCAAGACATC AAGCCTTGCGT GATGYGAGGC TGCCGCCGAG GCCCTCCTG']	2734	['TGATGYGAGGCT GCCGCCGAGG']	2946	['TGATGYGAGGCT GCCGCCGAGG']
NM_000435.2(NOTCH3):c.13 63T>C (p.Cys455Arg)	4854	NOTC H3	2567	['CCTCGACCGC ATAGGCCAGTT CACCYGTATCT GTATGGCAGGT GGGTGGTG']	2735	['ACCYGTATCTGTA TGGCAGGTGG']	2947 - 2948	['TTCACCYGTATC TGATGGCAGG', 'ACCYGTATCTGTA TGGCAGGTGG']
NM_002427.3(MMP13):c.272 T>C (p.Met91Thr)	4322	MMP1 3	2568	['CTTGACGATA ACACCTTAGAT GTCAYGAAAA AGCCAAGATG CGGGGTTCT']	2736 - 2737	['GTCAYGAAAAAG CCAAGATGCGG', 'TCAYGAAAAAGCC AAGATGCGGG']	2949 - 2950	['GTCAYGAAAA GCCAAGATGCGG', 'TCAYGAAAAAGC CAAGATGCGGG']
NM_000211.4(ITGB2):c.446T >C (p.Leu149Pro)	3689	ITGB2	2569	['GATGACCTCA GGAATGTCAA GAAGCYAGGT GGCGACCTGCT CCGGGCCCTC']	2738	['AGCYAGGTGGCG ACCTGCTCCGG']	2951	['AGCYAGGTGGCG ACCTGCTCCGG']
NM_005502.3(ABCA1):c.442 9T>C (p.Cys1477Arg)	19	ABCA 1	2570	['CAAAATCAAG AAGATGCTGCC TGTGYGTCCCC CAGGGGCAGG GGGGCTGCC']	2739 - 2740	['CCTGTGYGTCCCC CAGGGGCAGG', 'CTGTGYGTCCCC AGGGGCAGGG']	2952 - 2955	['CCTGTGYGTCCC CCAGGGGCAGG', 'CTGTGYGTCCCC AGGGGCAGGG', 'TGTGYGTCCCCA

								GGGGCAGGGG', 'GTGYGTCCCCCA GGGGCAGGGGG']
m.12297T>C	4568	MT- TL2	2571	['AAAGGATAAC AGCTATCCATT GGTCYTAGGCC CCAAAAATTTT GGTGCAAC']	2741	['GTCYTAGGCCCC AAAAATTTTGG']	2956	['GTCYTAGGCCCC AAAAATTTTGG']
m.4290T>C	4565	MT-TI	2572	['AAATATGTCT GATAAAAGAG TACTYTGATA GAGTAAATAA TAGGAGCTTA']	2742	['ACTYTGATAGAG TAAATAATAGG']	2957	['ACTYTGATAGAG TAAATAATAGG']
m.4291T>C	4565	MT-TI	2573	['AATATGTCTG ATAAAAGAGT TACTTYGATAG AGTAAATAAT AGGAGCTTAA']	2743	['ACTTYGATAGAG TAAATAATAGG']	2958	['ACTTYGATAGAG TAAATAATAGG']
m.3394T>C	4535	MT- ND1	2574	['GCTTACCGAA CGAAAAATTCT AGGCYATATA CAACTACGCA AAGGCCCAA']	2744	['GGCYATATACAA CTACGCAAAGG']	2959	['GGCYATATACAA CTACGCAAAGG']
NM_002764.3(PRPS1):c.344T >C (p.Met115Thr)	5631	PRPS1	2575	['ATCTCAGCCA AGCTTGTTGCA AATAYGCTATC TGTAGCAGGTG CAGATCAT']	2745	['GCAAATAYGCTA TCTGTAGCAGG']	2960	['GCAAATAYGCTA TCTGTAGCAGG']
NM_000132.3(F8):c.5372T>C (p.Met1791Thr)	2157	F8	2576	['AGAGCAGAA GTTGAAGATA ATATCAYGGTG AGTTAAGGAC AGTGGAATTAC ']	2746	['TCAYGGTGAGTT AAGGACAGTGG']	2961	['TCAYGGTGAGTT AAGGACAGTGG']
NM_000132.3(F8):c.1754T>C (p.Ile585Thr)	2157	F8	2577	['CCTTTC AATA TATGTAATTAA CAGAYAATGT CAGACAAGAG GAATGTCATC']	2747	['AACAGAYAATGT CAGACAAGAGG']	2962	['AACAGAYAATGT CAGACAAGAGG']
NM_000133.3(F9):c.1328T>C (p.Ile443Thr)	2158	F9	2578	['TGTGCAATGA AAGGCAAATA TGGAAYATAT ACCAAGGTATC CCGGTATGTC']	2748	['GAAYATATACCA AGGTATCCCGG']	2963	['GAAYATATACCA AGGTATCCCGG']

NM_000169.2(GLA):c.806T> C (p.Val269Ala)	-1	-	2579	['TTATTTTCATTC TTTTTCTCAGT TAGYGATTGGC AACTTTGGCCT CAGCTGG']	2749	['CAGTTAGYGATT GGCAACTTTGG']	2964	['CAGTTAGYGATT GGCAACTTTGG']
NM_000116.4(TAZ):c.352T> C (p.Cys118Arg)	6901	TAZ	2580	['CTCCCACTTC TTCAGCTTGGG CAAGYGTGTG CCTGTGTGCCG AGGTGAGCT']	2750	['AAGYGTGTGCCT GTGTGCCGAGG']	2965	['AAGYGTGTGCCT GTGTGCCGAGG']
NM_000061.2(BTK):c.2T>C (p.Met1Thr)	695	BTK	2581	['GGTGAACTCC AGAAAAGAAGA AGCTAYGGCC GCAGTGATTCT GGAGAGCATC']	2751	['AGCTAYGGCCGC AGTGATTCTGG']	2966	['AGCTAYGGCCGC AGTGATTCTGG']
NM_000061.2(BTK):c.1223T> C (p.Leu408Pro)	695	BTK	2582	['AAGGACCTGA CCTTCTGAAG GAGCYGGGGA CTGGACAATTT GGGGTAGTG']	2752	['AGCYGGGGACTG GACAATTTGGG']	2967 - 2968	['GAGCYGGGGACT GGACAATTTGG', 'AGCYGGGGACTG GACAATTTGGG']
NM_000061.2(BTK):c.1741T> C (p.Trp581Arg)	695	BTK	2583	['CAAGTTCAGC AGCAAATCTG ACATTYGGGCT TTTGGTAAGTG GATAAGATT']	2753	['ACATTYGGGCTTT TGGTAAGTGG']	2969	['ACATTYGGGCTT TTGGTAAGTGG']
NM_014009.3(FOXP3):c.970T >C (p.Phe324Leu)	5094 3	FOXP3	2584	['GATTCATCCC CACCTCTGAC AGAGYTCCTCC ACAACATGGA CTACTTCAA']	2754	['GACAGAGYTCCT CCACAACATGG']	2970	['GACAGAGYTCCT CCACAACATGG']
NM_003688.3(CASK):c.2740 T>C (p.Trp914Arg)	8573	CASK	2585	['TGAGCTCGTG TGCACAGCCCC ACAGYGGGTC CCTGTCTCCTG GGTCTATTA']	2755 - 2756	['CACAGYGGGTCC CTGTCTCCTGG', 'ACAGYGGGTCCCT GTCTCCTGGG']	2971 - 2972	['CACAGYGGGTCC CTGTCTCCTGG', 'ACAGYGGGTCCC TGTCTCCTGGG']
NM_004992.3(MECP2):c.464 T>C (p.Phe155Ser)	4204	MECP 2	2586	['GACACATCCC TGGACCCTAAT GATTBTGACTT CACGGTAACTG GGAGAGGG']	2757	['GATTBTGACTTCA CGGTAACTGG']	2973 - 2974	['GATTBTGACTTC ACGGTAACTGG', 'ATTBTGACTTCAC GGTAACTGGG']
NM_000431.3(MVK):c.803T> C (p.Ile268Thr)	4598	MVK	2587	['ATCGTGGCCC CCCTCCTGACC TCAAYAGATG CCATCTCCCTG']	2758	['CTCAAYAGATGC CATCTCCCTGG']	2975	['CTCAAYAGATGC CATCTCCCTGG']

				GAGTGTGAG']				
NM_021961.5(TEAD1):c.1261 T>C (p.Tyr?His)	7003	TEAD1	2588	['TGAACACGGA GCACAACATC ATATTYACAGG CTTGTAAAGGA CTGAACATG']	2759	['TCATATTYACAG GCTTGTAAAGG']	2976	['TCATATTYACAG GCTTGTAAAGG']
NM_005633.3(SOS1):c.1294T >C (p.Trp432Arg)	6654	SOS1	2589	['CGAGATTCAG AAGAATATTG ATGGTYGGGA GGGAAAAGAC ATTGGACAGTG ']	2760	['GGTYGGGAGGGA AAAGACATTGG']	2977	['GGTYGGGAGGG AAAAGACATTGG']
NM_006920.4(SCN1A):c.3577 T>C (p.Trp1193Arg)	-1	-	2590	['TGTGGAAGAA GGCAGAGGAA AACAAAYGGTG GAACCTGAGA AGGACGTGTTT ']	2761	['AACAAAYGGTGG AACCTGAGAAGG']	2978	['AACAAAYGGTGG AACCTGAGAAGG']
NM_000141.4(FGFR2):c.1018 T>C (p.Tyr340His)	2263	FGFR2	2591	['TGTAACTTTT GAGGACGCTG GGGAAAYATAC GTGCTTGCCGG GTAATTCTAT']	2762 - 2763	['TGGGGAAAYATAC GTGCTTGCCGG', 'GGGGAAYATACGT GCTTGGC'GGG']	2979 - 2980	['TGGGGAAAYATAC GTGCTTGCCGG', 'GGGGAAYATACG TGCTTGGC'GGG']
NM_000174.4(GP9):c.70T>C (p.Cys24Arg)	2815	GP9	2592	['GGCCACCAAG GACTGCCCCAG CCCAYGTACCT GCCGCGCCCTG GAAACCAT']	2764	['CCCAYGTACCTG CCGCGCCCTGG']	2981	['CCCAYGTACCTG CCGCGCCCTGG']
NM_000175.3(GPI):c.1574T>C (p.Ile525Thr)	2821	GPI	2593	['CTGGGAAAGC AGCTGGCTAA GAAAABAGAG CCTGAGCTTGA TGGCAGTGCT']	2765	['AAAABAGAGCCT GAGCTTGATGG']	2982	['AAAABAGAGCCT GAGCTTGATGG']
NM_000315.2(PTH):c.52T>C (p.Cys18Arg)	5741	PTH	2594	['AGTTATGATT GTCATGTTGGC AATTYGTTTTT TTACAAAATCG GATGGGAA']	2766	['AATTYGTTTTCTT ACAAATCGG']	2983	['AATTYGTTTTCTT ACAAATCGG']
NM_000222.2(KIT):c.1676T>C (p.Val559Ala)	3815	KIT	2595	['CCCATGTATG AAGTACAGTG GAAGGNTGTT GAGGAGATAA ATGGAAACAA T']	2767	['AAGGNTGTTGAG GAGATAAATGG']	2984	['AAGGNTGTTGAG GAGATAAATGG']

NM_016835.4(MAPT):c.1839 T>C (p.Asn613=)	4137	MAPT	2596	['AGTCCAAGTG TGGCTCAAAG GATAAYATCA AACACGTCCCG GGAGGCGGCA']	2768 - 2769	['GGATAAYATCAA ACACGTCCCGG', 'GATAAYATCAAAC ACGTCCCGGG']	2985 - 2986	['GGATAAYATCAA ACACGTCCCGG', 'GATAAYATCAAA CACGTCCCGGG']
NM_170707.3(LMNA):c.1139 T>C (p.Leu380Ser)	4000	LMNA	2597	['GAGATCCACG CCTACCGCAAG CTCTYGGAGG GCGAGGAGGA GAGGTGGGCT']	2770	['TCTYGGAGGGCG AGGAGGAGAGG']	2987	['TCTYGGAGGGCG AGGAGGAGAGG']
NM_000424.3(KRT5):c.20T> C (p.Val7Ala)	3852	KRT5	2598	['GCCACCATGT CTCGCCAGTCA AGTGYGTCCTT CCGGAGCGGG GGCAGTCGT']	2771 -- 2773	['TCAAGTGYGTCCT TCCGGAGCGG', 'CAAGTGYGTCCTT CCGGAGCGGG', 'AAGTGYGTCCTT CCGGAGCGGG']	2988 - 2991	['TCAAGTGYGTCCT TCCGGAGCGG', 'CAAGTGYGTCCTT CCGGAGCGGG', 'AAGTGYGTCCTT CCGGAGCGGG', 'AGTGYGTCCTTCC GGAGCGGGGG']
NM_000184.2(HBG2):c.125T >C (p.Phe42Ser)	3048	HBG2	2599	['GTTGTCTACC CATGGACCCA GAGGTYCTTTG ACAGCTTTGGC AACCTGTCC']	2774	['CAGAGGTYCTTT GACAGCTTTGG']	2992	['CAGAGGTYCTTT GACAGCTTTGG']
NM_000515.4(GHI):c.291+6T >C	2688	GHI	2600	['AGGAAACAC AACAGAAATC CGTGAGYGGA TGCCTTCTCCC CAGGCGGGGA T']	2775	['TGAGYGGATGCC TTCTCCCCAGG']	2993	['TGAGYGGATGCC TTCTCCCCAGG']
NM_002087.3(GRN):c.2T>C (p.Met1Thr)	2896	GRN	2601	['TCCTTGGTAC TTTGCAGGCAG ACCAYGTGGA CCCTGGTGAGC TGGGTGGCC']	2776	['CCAYGTGGACCC TGGTGAGCTGG']	2994	['CCAYGTGGACCC TGGTGAGCTGG']
NM_00108311 2.2(GPD2):c.19 04T>C (p.Phe635Ser)	2820	GPD2	2602	['AGGTATAAGA AGAGATTTTAT AAGTYTGATGC AGACCAGAAA GGCTTTATT']	2777	['AAGTYTGATGCA GACCAGAAAGG']	2995	['AAGTYTGATGCA GACCAGAAAGG']
NM_00101807 7.1(NR3C1):c.1 712T>C (p.Val571Ala)	2908	NR3C1	2603	['CTCAACATGT TAGGAGGGCG GCAAGYGATT GCAGCAGTGA	2778	['AAGYGATTGCAG CAGTCAAATGG']	2996	['AAGYGATTGCAG CAGTCAAATGG']

				AATGGGCAAA G']				
NM_000138.4(FBN1):c.3793T >C (p.Cys1265Arg)	2200	FBN1	2604	['TATCCCTGGA GAGTACAGGT GCTTGYGTTAT GATGGATTCAT GGCATCTGA']	2779	['CTTGYGTTATGAT GGATTCATGG']	2997	['CTTGYGTTATGA TGGATTCATGG']
NM_000129.3(F13A1):c.728T >C (p.Met243Thr)	2162	F13A1	2605	['ATCCTGGACA CTTGCCTGTAT GTGAYGGACA GAGCACAAAT GGACCTCTCT']	2780	['TGTGAYGGACAG AGCACAAATGG']	2998	['TGTGAYGGACAG AGCACAAATGG']
NM_031226.2(CYP19A1):c.74 3+2T>C	-1	-	2606	['ATACAAAAAG TATGAGAAGTC TGTGYAAGTA ATACAACCTTTG GAAGATTTA']	2781	['CTGTGYAAGTAA TACAACCTTTGG']	2999	['CTGTGYAAGTAA TACAACCTTTGG']
NM_000416.2(IFNGR1):c.260 T>C (p.Ile87Thr)	3459	IFNGR 1	2607	['AATATTTCTC ATCATTATTGT AATAYTTCTGA TCATGTTGGTG ATCCATCA']	2782	['TGTAATAYTTCTG ATCATGTTGG']	3000	['TGTAATAYTTCT GATCATGTTGG']
NM_000018.3(ACADVL):c.84 8T>C (p.Val283Ala)	37	ACAD VL	2608	['GTGAAGGAG AAGATCACAG CTTTTGYGGTG GAGAGGGGCT TCGGGGGCATT ']	2783 - 2784	['TTTGYGGTGGAG AGGGGCTTCGG', 'TTGYGGTGGAGAG GGGCTTCGGG']	3001 - 3002	['TTTGYGGTGGAG AGGGGCTTCGG', 'TTGYGGTGGAGA GGGGCTTCGGG']
NM_000195.4(HPS1):c.716T> C (p.Leu239Pro)	3257	HPS1	2609	['GCTGGGGTAG AGGTCCTGAAC CAGGRGGATG AGGGCAAGCA GGTCGGCCGG']	2785	['CCAGGRGGATGA GGGCAAGCAGG']	3003	['CCAGGRGGATGA GGGCAAGCAGG']
NM_000352.4(ABCC8):c.257 T>C (p.Val86Ala)	6833	ABCC 8	2610	['ACCTTCATGC TGCTCTTCGTC CTGGBGTGTGA GATTGCAGAG GGCATCCTG']	2786 - 2787	['CCTGGBGTGTGA GATTGCAGAGG', 'CTGGBGTGTGAGA TTGCAGAGGG']	3004 - 3005	['CCTGGBGTGTGA GATTGCAGAGG', 'CTGGBGTGTGAG ATTGCAGAGGG']
NM_000528.3(MAN2B1):c.24 26T>C (p.Leu809Pro)	4125	MAN2 B1	2611	['GGCAGCAGCC TGAGAGATGG CTCGCYGGAG CTCATGGTGAG TGGGTCAGAG']	2788	['CGCYGGAGCTCA TGGTGAGTGGG']	3006 - 3007	['TCGCYGGAGCTC ATGGTGAGTGG', 'CGCYGGAGCTCA TGGTGAGTGGG']
NM_002863.4(PYGL	5836	PYGL	2612	['CTCCAGTGAC	2789	['AAGAAYATGCC	3008	['AAGAAYATGCC

PYGL):c.2461 T>C (p.Tyr821His)				CGAACAATTA AAGAAAYATGC CCAAAACATCT GGAACGTGGA']		AAAACATCTGG']		AAAACATCTGG']
NM_000495.4(COL4A5):c.46 99T>C (p.Cys1567Arg)	1287	COL4 A5	2613	['CTTTTCCTTTA CCAGATGTGCA GTAYGTGAAG CTCCAGCTGTG GTGATCGC']	2790	['AGTAYGTGAAGC TCCAGCTGTGG']	3009	['AGTAYGTGAAGC TCCAGCTGTGG']
NM_000155.3(GALT):c.460T >C (p.Trp154Arg)	2592	GALT	2614	['TGAGATCCGG GCTGTTGTTGA TGCABGGGCCT CAGTCACAGA GGAGCTGGG']	2791	['TGCABGGGCCTC AGTCACAGAGG']	3010	['TGCABGGGCCTC AGTCACAGAGG']
NM_000155.3(GALT):c.482T >C (p.Leu161Pro)	2592	GALT	2615	['GCATGGGCCT CAGTCACAGA GGAGCYGGGT GCCCAGTACCC TTGGGTGCAG']	2792	['AGCYGGGTGCCC AGTACCCTTGG']	3011	['AGCYGGGTGCCC AGTACCCTTGG']
NM_000155.3(GALT):c.499T >C (p.Trp167Arg)	2592	GALT	2616	['AGAGGAGCTG GGTGCCAGTA CCCTYGGGTGC AGGTTTGTGAG GTCGCCCC']	2793	['CCCTYGGGTGCA GGTTTGTGAGG']	3012	['CCCTYGGGTGCA GGTTTGTGAGG']
NM_000155.3(GALT):c.509T >C (p.Ile170Thr)	2592	GALT	2617	['GAGCTCCGTA TCCCTATCTGA TAGAHCTTTGA AAACAAAGGT GCCATGATG']	2794	['TGATAGAHCTTTG AAACAAAGG']	3013	['TGATAGAHCTTT GAAAACAAAGG']
NM_004523.3(KIF11):c.2547+ 2T>C	3832	KIF11	2618	['CAGGAACTTC ACAACCTATTG GAGGYAATAA CTTTGTAAGTG GAACTTACT']	2795	['GGAGGYAATAAC TTTGTAAGTGG']	3014	['GGAGGYAATAA CTTTGTAAGTGG']
NM_005211.3(CSF1R):c.2381 T>C (p.Ile794Thr)	1436	CSF1R	2619	['TTGACCAATG GTCATGTGGCC AAGAYTGGGG ACTTCGGGCTG GCTAGGGAC']	2796	['CAAGAYTGGGGA CTTCGGGCTGG']	3015	['CAAGAYTGGGG ACTTCGGGCTGG']
NM_005188.3(CBL):c.1150T> C (p.Cys384Arg)	867	CBL	2620	['CTCCACATTC CAACTATGTAA AATABGTGCTG AAAATGATAA GGATGTAAA']	2797	['AATABGTGCTGA AAATGATAAGG']	3016	['AATABGTGCTGA AAATGATAAGG']

NM_006306.3(SMC1A):c.235 IT>C (p.Ile784Thr)	8243	SMC1 A	2621	['GTGTTTGAAG AGTTTTGTCGG GAGAYTGGTG TGCGCAACATC CGGGAGTTT']	2798	['AGAYTGGTGTGC GCAACATCCGG']	3017	['AGAYTGGTGTGC GCAACATCCGG']
NM_002242.4(KCNJ13):c.722 T>C (p.Leu241Pro)	-1	-	2622	['TGGTGTAATG GAGTGATAGT ACGTTDGTGGA AAGATGAAGA ATGGACATTC']	2799	['GTTDGTGGAAAG ATGAAGAATGG']	3018	['GTTDGTGGAAAG ATGAAGAATGG']
NM_000199.3(SGSH):c.892T >C (p.Ser298Pro)	6448	SGSH	2623	['CCCCAGCGTT TTGGGTGCTCC GGGGRTGACA CCAGTAAGGG TTCAGCAGTG']	2800	['TCCGGGGRTGAC ACCAGTAAGGG']	3019	['TCCGGGGRTGAC ACCAGTAAGGG']
NM_020191.2(MRPS22):c.644 T>C (p.Leu215Pro)	5694 5	MRPS2 2	2624	['CCAATAATTT TCAAGGAAGA AAATCYTAGG GTAAGGTGACT TAGGTTTTAT']	2801	['ATCYTAGGGTAA GGTGACTTAGG']	3020	['ATCYTAGGGTAA GGTGACTTAGG']
NM_017882.2(CLN6):c.200T> C (p.Leu67Pro)	5498 2	CLN6	2625	['CCCATTCTTC CATTTGCTCCG CAGCYGGTATT CCCTCTCGAGT GGTTTCCA']	2802	['AGCYGGTATTCC CTCTCGAGTGG']	3021	['AGCYGGTATTCC CTCTCGAGTGG']
NM_014874.3(MFN2):c.1392 +2T>C	9927	MFN2	2626	['GTAGTCCTCA AGGTTTATAAG AATGWGAGTC ATGGAGCAAC AGGTCTCTTT']	2803	['AATGWGAGTCAT GGAGCAACAGG']	3022	['AATGWGAGTCA TGGAGCAACAGG']
NM_024599.5(RHBDF2):c.55 7T>C (p.Ile186Thr)	7965 1	RHBD F2	2627	['GCTTACCGCC CCCCTCCCTTC CAGAYTGTGG ATCCGCTGGCC CGGGGCCGG']	2804	['AGAYTGTGGATC CGCTGGCCCCGG']	3023	['AGAYTGTGGATC CGCTGGCCCCGG']
NM_020894.2(UVSSA):c.94T >C (p.Cys32Arg)	5765 4	UVSS A	2628	['GAAAATGAA GGAAGTGAAG AAAATTYGCA AGTATGTCTTA GGGTTTCAGTAA ']	2805	['AAAATTYGCAAG TATGTCTTAGG']	3024 - 3025	['AAAATTYGCAAG TATGTCTTAGG', 'AAATTYGCAAGT ATGTCTTAGGG']
NM_00116158 1.1(POC1A):c. 398T>C	2588 6	POC1A	2629	['GCCAGTGATG ACAAGACTGTT AAGCYGTGGG	2806	['AGCYGTGGGACA AGAGCAGCCGG']	3026	['AGCYGTGGGACA AGAGCAGCCGG']

(p.Leu133Pro)				ACAAGAGCAG CCGGGAATGT']				
NM_005340.6(HINT1):c.250T >C (p.Cys84Arg)	3094	HINT1	2630	['ACACTTAATG ATTGTTGGCAA GAAAYGTGCT GCTGATCTGGG CCTGAATAA']	2807 - 2808	['CAAGAAAYGTGC TGCTGATCTGG', 'AAGAAAYGTGCTG CTGATCTGGG']	3027 - 3028	['CAAGAAAYGTGC TGCTGATCTGG', 'AAGAAAYGTGCT GCTGATCTGGG']
NM_000495.4(COL4A5):c.43 8+2T>C	1287	COL4 A5	2631	['TTTCCTGGTTT ACAGGGTCCTC CAGYAAGTTAT AAAATTTGGG ATTATGAT']	2809	['TCCAGYAAGTTA TAAAATTTGGG']	3029 - 3030	['CTCCAGYAAGTT ATAAAATTTGG', 'TCCAGYAAGTTA TAAAATTTGGG']
NM_000344.3(SMN1):c.388T >C (p.Tyr130His)	6606	SMN1	2632	['AACCTGTGTT GTGGTTTACAC TGGAYATGGA AATAGAGAGG AGCAAAATCT']	2810	['CACTGGAYATGG AAATAGAGAGG']	3031	['CACTGGAYATGG AAATAGAGAGG']
NM_005334.2(HCFC1):c.- 970T>C	3054	HCFC1	2633	['TTAGTTGTTA CTTCTCACAC AAGAYGGCGG CTCCCAGGGA GGAGGCATGA']	2811	['CAAGAYGGCGGC TCCCAGGGAGG']	3032	['CAAGAYGGCGG CTCCCAGGGAGG']
NM_000431.3(MVK):c.1039+ 2T>C	4598	MVK	2634	['GTGGCATCAC ACTCCTCAAGC CAGGYATCCC GGGGGTAGGT GGGCCAGGCT']	2812	['CCAGGYATCCC GGGGTAGGTGG']	3033 - 3034	['CCAGGYATCCC GGGGTAGGTGG', 'CAGGYATCCC GGGTAGGTGGG']
NM_018344.5(SLC29A3):c.60 7T>C (p.Ser203Pro)	5531 5	SLC29 A3	2635	['TATGAGGAAC TCCCAGGCACT GATAYCAGGT GAGAGCCAGG GTCCGGCAG']	2813	['ACTGATAYCAGG TGAGAGCCAGG']	3035 - 3036	['ACTGATAYCAGG TGAGAGCCAGG', 'CTGATAYCAGGT GAGAGCCAGGG']
NM_000108.4(DLD):c.140T> C (p.Ile47Thr)	1738	DLD	2636	['GTAGTTGATG CTGATGTAACA GTTAYAGGTTT TGGTCTGGAG GATATGTT']	2814 - 2815	['ACAGTTAYAGGT TCTGGTCTGG', 'GTTAYAGGTTCTG GTCCTGGAGG']	3037 - 3038	['ACAGTTAYAGGT TCTGGTCTGG', 'GTTAYAGGTTCTG GTCCTGGAGG']
NM_004333.4(BRAF):c.1403 T>C (p.Phe468Ser)	673	BRAF	2637	['GGACAAAGA ATTGGATCTGG ATCATYTGGAA CAGTCTACAAG GGAAAGTGG']	2816 - 2817	['ATCATYTGGAAC AGTCTACAAGG', 'TCATYTGGAACAG TCTACAAGGG']	3039 - 3040	['ATCATYTGGAAC AGTCTACAAGG', 'TCATYTGGAAC GTCTACAAGGG']
NM_000540.2(RYR1):c.1205T	6261	RYR1	2638	['CAGGAGGAGT CCCAGGCCGCC	2818	['CGCAYGATCCAC AGCACCAATGG']	3041	['CGCAYGATCCAC AGCACCAATGG']

>C (p.Met402Thr)				CGCAYGATCC ACAGACCAA TGGCCTATAC']				
NM_000256.3(MYBPC3):c.13 51+2T>C	4607	MYBP C3	2639	['GTAGCACGGA GCTCTTTGTGA AAGGYGGGCC TGGGACCTGA GGATGTGGGA']	2819	['AAAGGYGGGCCT GGGACCTGAGG']	3042	['AAAGGYGGGCCT GGGACCTGAGG']
NM_000256.3(MYBPC3):c.82 1+2T>C	4607	MYBP C3	2640	['CCTCCTATCA GCCTTCCGCCG CACGYGAGTG GCCATCCTCAG GGCCTGGGG']	2820	['CACGYGAGTGGC CATCCTCAGGG']	3043 - 3044	['GCACGYGAGTGG CCATCCTCAGG', 'CACGYGAGTGGC CATCCTCAGGG']
NM_000257.3(MYH7):c.2546 T>C (p.Met849Thr)	4625	MYH7	2641	['AAGAGTGCAG AAAGAGAGAA GGAGAYGGCC TCCATGAAGG AGGAGTTCAC A']	2821	['GGAGAYGGCCTC CATGAAGGAGG']	3045	['GGAGAYGGCCTC CATGAAGGAGG']
NM_206933.2(USH2A):c.160 6T>C (p.Cys536Arg)	7399	USH2 A	2642	['CGACACAACA AGCCAGCCAT ATAGAYGCCTC TGCTCCCAGGA GAGCTTCAC']	2822	['ATATAGAYGCCT CTGCTCCCAGG']	3046	['ATATAGAYGCCT CTGCTCCCAGG']
NM_000059.3(BRCA2):c.316 +2T>C	675	BRCA 2	2643	['TAGATAAATT CAAATTAGACT TAGGYAAGTA ATGCAATATGG TAGACTGGG']	2823	['CTTAGGYAAGTA ATGCAATATGG']	3047	['CTTAGGYAAGTA ATGCAATATGG']
NM_007294.3(BRCA1):c.529 1T>C (p.Leu1764Pro)	672	BRCA 1	2644	['CTCTTCTTCC AGATCTTCAGG GGGCYAGAAA TCTGTTGCTAT GGGCCCTTC']	2824	['GGCYAGAAATCT GTTGCTATGGG']	3048 - 3049	['GGGCYAGAAATC TGTTGCTATGG', 'GGCYAGAAATCT GTTGCTATGGG']
NM_00113008 9.1(KARS):c.5 17T>C (p.Tyr173His)	3735	KARS	2645	['AGCTTCTGGG GGAAAGCTCA TCTTCYATGAT CTTCGAGGAG AGGGGGTGAA']	2825	['TTCYATGATCTTC GAGGAGAGGG']	3050 - 3051	['CTTCYATGATCT TCGAGGAGAGG', 'TTCYATGATCTTC GAGGAGAGGG']
NM_00128300 9.1(RTEL1):c.3 730T>C (p.Cys1244Arg)	-1	-	2646	['CGGGCCCCTC TCAGCAGGCTG TGTGYGCCAG GGCTGTGGGG']	2826	['CTGTGTGYGCCA GGGCTGTGGGG']	3052	['CTGTGTGYGCCA GGGCTGTGGGG']

)				CAGAGGACGT']				
NM_005554.3(KRT6A):c.140 6T>C (p.Leu469Pro)	3853	KRT6 A	2647	['GAGATCGCCA CCTACCGCAAG CTGCBGGAGG GTGAGGAGTG CAGGTGGGTA']	2827	['TGCBGGAGGGTG AGGAGTGCAGG']	3053	['TGCBGGAGGGTG AGGAGTGCAGG']
NM_000218.2(KCNQ1):c.550 T>C (p.Tyr184His)	3784	KCNQ I	2648	['CTGGTCCGCC GGCTGCCGCA GCAAGBACGT GGGCCTCTGGG GGCGCTGCG']	2828	['AGCAAGBACGTG GGCCTCTGGGG']	3054 - 3056	['CAGCAAGBACGT GGGCCTCTGGG', 'AGCAAGBACGTG GGCCTCTGGGG', 'GCAAGBACGTGG GCCCTCTGGGG']
NM_198056.2(SCN5A):c.5624 T>C (p.Met1875Thr)	6331	SCN5A	2649	['GAGATGGACG CCCTGAAGATC CAGAHGGAGG AGAAGTTCATG GCAGCCAAC']	2829	['CCAGAHGGAGGA GAAGTTCATGG']	3057	['CCAGAHGGAGG AGAAGTTCATGG']
NM_006920.4(SCN1A):c.269 T>C (p.Phe90Ser)	6323	SCN1A	2650	['TGTTGTGTTC CTGTCTTACAG ACTTYTATAGT ATTGAATAAA GGGAAGGCC']	2830 - 2831	['ACTTYTATAGTAT TGAATAAAGG', 'CTTYTATAGTATT GAATAAAGGG']	3058 - 3059	['ACTTYTATAGTA TTGAATAAAGG', 'CTTYTATAGTATT GAATAAAGGG']
NM_006920.4(SCN1A):c.272 T>C (p.Ile91Thr)	6323	SCN1A	2651	['TGTGTTCCCTG TCTTACAGACT TTTAYAGTATT GAATAAAGGG AAGGCCATC']	2832 - 2833	['ACTTTTAYAGTAT TGAATAAAGG', 'CTTTTAYAGTATT GAATAAAGGG']	3060 - 3061	['ACTTTTAYAGTA TTGAATAAAGG', 'CTTTTAYAGTATT GAATAAAGGG']
NM_006514.3(SCN10A):c.166 IT>C (p.Leu554Pro)	6336	SCN10 A	2652	['GGAGTCAGGG TTGCTGGGTTG AGGARGAGGG CTTCTAGGGAG GGGGCCTTG']	2834 - 2836	['GAGGARGAGGGC TTCTAGGGAGG', 'AGGARGAGGGCTT CTAGGGAGGG', 'GGARGAGGGCTTC TAGGGAGGGG']	3062 - 3064	['GAGGARGAGGG CTTCTAGGGAGG', 'AGGARGAGGGCT TCTAGGGAGGG', 'GGARGAGGGCTT CTAGGGAGGGG']
NM_000251.2(MSH2):c.2005 +2T>C	4436	MSH2	2653	['AACAGATGTT CCACATCATTA CTGGYAAAAA ACCTGGTTTTT GGGCTTTGT']	2837 - 2838	['CTGGYAAAAAAC CTGGTTTTTGG', 'TGGYAAAAAACCT GGTTTTTGGG']	3065 - 3066	['CTGGYAAAAAAC CTGGTTTTTGG', 'TGGYAAAAAACCT TGGTTTTTGGG']
NM_000251.2(MSH2):c.595T >C (p.Cys199Arg)	4436	MSH2	2654	['CCTCATCCAG ATTGGACCAA AGGAAYGTGT TTTACCCGGAG GAGAGACTGC']	2839	['AAGGAAYGTGTT TTACCCGGAGG']	3067	['AAGGAAYGTGTT TTACCCGGAGG']
NM_00100574	2629	GBA	2655	['TTCACCGCTC	2840	['GCCRAGTGGGTG	3068	['GAGCCRAGTGGG

1.2(GBA):c.667 T>C (p.Trp223Arg)				CATTGGTCTTG AGCCRAGTGG GTGATGTCCAG GGGCTGGCA']		ATGTCCAGGGG']	- 3070	TGATGTCCAGG', 'AGCCRAGTGGGT GATGTCCAGGG', 'GCCRAGTGGGTG ATGTCCAGGGG']
NM_003494.3(DYSF):c.1284+ 2T>C	8291	DYSF	2656	['GAGGTCAGCT TTGCGGGGAA AATGGYAAGG AGCAAGGGAG CAGGAGGGTT C']	2841	['ATGGYAAGGAGC AAGGGAGCAGG']	3071	['ATGGYAAGGAG CAAGGGAGCAGG']
NM_012463.3(ATP6V0A2):c. 825+2T>C	2354 5	ATP6V 0A2	2657	['ACCCGCATCC AGGATCTCTAC ACTGYGAGTA AGCTGGAAGT GGATTGCCTC']	2842	['CACTGYGAGTAA GCTGGAAGTGG']	3072	['CACTGYGAGTAA GCTGGAAGTGG']
NM_016725.2(FOLR1):c.493+ 2T>C	2348	FOLR1	2658	['ACAAGGGCTG GAACTGGACTT CAGGYGAGGG CTGGGGTGGG CAGGAATGGA']	2843	['AGGYGAGGGCTG GGGTGGGCAGG']	3073 - 3074	['CTTCAGGYGAGG GCTGGGGTGGG', 'AGGYGAGGGCTG GGGTGGGCAGG']
NM_003764.3(STX11):c.173T >C (p.Leu58Pro)	8676	STX11	2659	['GACATTCAGG ATGAAAACCA GCTGCGYGGTG GCCGACGTGA AGCGGCTGGG A']	2844	['TGCYGGTGGCCG ACGTGAAGCGG']	3075	['TGCYGGTGGCCG ACGTGAAGCGG']
NM_014714.3(IFT140):c.4078 T>C (p.Cys1360Arg)	9742	IFT140	2660	['GGACCCCAAG GAGTCCATCAA GCAGYGTGAG CTGCTCCTGGA GGAACCAGA']	2845	['GCAGYGTGAGCT GCTCCTGGAGG']	3076 - 3077	['CAAGCAGYGTGA GCTGCTCCTGG', 'GCAGYGTGAGCT GCTCCTGGAGG']
NM_000531.5(OTC):c.1005+2 T>C	5009	OTC	2661	['GAAAACAGA AAGTGGACAA TCATGGYAAG CAAGAAACAA GGAATGGAGG AT']	2846	['ATCATGGYAAGC AAGAAACAAGG']	3078	['ATCATGGYAAGC AAGAAACAAGG']
NM_000531.5(OTC):c.158T> C (p.Ile53Thr)	5009	OTC	2662	['CTAAAAAACT TTACCGGAGA AGAAABTAAA TATATGCTATG GCTATCAGCA']	2847	['AAGAAABTAAAT ATATGCTATGG']	3079	['AAGAAABTAAAT ATATGCTATGG']
NM_000531.5(OTC):c.158T> C (p.Ile53Thr)	5009	OTC	2663	['GAGAAAAGA GCTATCAGCA']	2848	['ACAAGATYGTCT GCTATCAGCA']	3080	['ACAAGATYGTCT GCTATCAGCA']

OTC):c.284T> C (p.Leu95Ser)				AGTACTCGAAC AAGATYGTCTA CAGAAACAGG TAAGTCCACT']		ACAGAAACAGG']		ACAGAAACAGG']
NM_000531.5(OTC):c.2T>C (p.Met1Thr)	5009	OTC	2664	['CGTCCTTTAC ACAATTA GAAGAYGCTG TTTAATCTGAG GATCCTGTTA']	2849	['AGAAGAYGCTGT TTAATCTGAGG']	3081	['AGAAGAYGCTGT TTAATCTGAGG']
NM_000531.5(OTC):c.526T> C (p.Tyr176His)	5009	OTC	2665	['CCATCCTATC CAGATCCTGGC TGATYACCTCA CGCTCCAGGTT GGTTTATT']	2850	['GGCTGATYACCT CACGCTCCAGG']	3082 - 3083	['GGCTGATYACCT CACGCTCCAGG', 'GATYACCTCACG CTCCAGGTTGG']
NM_000531.5(OTC):c.779T> C (p.Leu260Ser)	5009	OTC	2666	['GAAGCAGCGC ATGGAGGCAA TGTATYAATTA CAGACACTTGG ATAAGCATG']	2851	['ATGTATYAATTAC AGACACTTGG']	3084	['ATGTATYAATTA CAGACACTTGG']
NM_000322.4(PRPH2):c.736T >C (p.Trp246Arg)	5961	PRPH2	2667	['CCACCAGACG GAGGAGCTCA ACCTGYGGGT GCGTGGCTGCA GGGCTGCCCT']	2852 - 2853	['ACCTGYGGGTGC GTGGCTGCAGG', 'CCTGYGGGTGCGT GGCTGCAGGG']	3085 - 3086	['ACCTGYGGGTGC GTGGCTGCAGG', 'CCTGYGGGTGCG TGGCTGCAGGG']
NM_000211.4(ITGB2):c.1877 +2T>C	3689	ITGB2	2668	['CCCCTCACCC TGTGGCAAGTA CATGYGAGTG CAGGCGGAGC AGGCAGGGCG']	2854	['CATGYGAGTGCA GGCGGAGCAGG']	3087	['CATGYGAGTGCA GGCGGAGCAGG']
NM_015474.3(SAMHD1):c.11 06T>C (p.Leu369Ser)	2593 9	SAMH D1	2669	['TTTGTGTTGA TAAGCTCTACG GTGTRAAGAGT TGCGAGTGTGG AACATGTC']	2855	['GGTGTTRAAGAGT TGCGAGTGTGG']	3088	['GGTGTTRAAGAGT TGCGAGTGTGG']
NM_001101.3(ACTB):c.356T >C (p.Met119Thr)	60	ACTB	2670	['AACCCCAAGG CCAACCGCGA GAAGAYGACC CAGGTGAGTG GCCCCGTACCT']	2856	['GAGAAGAYGACC CAGGTGAGTGG']	3089	['GAGAAGAYGAC CCAGGTGAGTGG']
NM_015713.4(RRM2B):c.368 T>C	5048 4	RRM2 B	2671	['CTCGATGAGA ATTTGAAAGCC ATAGRAACAG	2857	['CCATAGRAACAG CGAGCCTCTGG']	3090	['CCATAGRAACAG CGAGCCTCTGG']

(p.Phe123Ser)				CGAGCCTCTGG AACCTGCAC']				
NM_015599.2(PGM3):c.248T >C (p.Leu83Ser)	5238	PGM3	2672	['TTGGTTGATC CTTTGGGTGAA ATGTYGGCACC ATCCTGGGAG GAACATGCC']	2858	['AATGTYGGCACC ATCCTGGGAGG']	3091	['AATGTYGGCACC ATCCTGGGAGG']
NM_002136.2(HNRNPA1):c.8 17T>C (p.Phe273Leu)	3178	HNRN PA1	2673	['GAATTACAAC AATCAGTCTTC AAATBTTGGAC CCATGAAGGG AGGAAATTT']	2859 - 2861	['TTCAAATBTTGGA CCCATGAAGG', 'TCAAATBTTGGAC CCATGAAGGG', 'AATBTTGGACCCA TGAAGGGAGG']	3092 - 3094	['TTCAAATBTTGG ACCCATGAAGG', 'TCAAATBTTGGAC CCATGAAGGG', 'AATBTTGGACCC ATGAAGGGAGG']
NM_002136.2(HNRNPA1):c.8 41T>C (p.Phe281Leu)	3178	HNRN PA1	2674	['TTTTGGACCC ATGAAGGGAG GAAATYTTGG AGGCAGAAGC TCTGGCCCCTA']	2862	['AATYTTGGAGGC AGAAGCTCTGG']	3095	['AATYTTGGAGGC AGAAGCTCTGG']
NM_022552.4(DNMT3A):c.27 05T>C (p.Phe902Ser)	1788	DNMT 3A	2675	['CGCAAAATAC TCCTTCAGCGG AGCGRAGAGG TGGCGGATGA CTGGCACGCT']	2863	['GCGRAGAGGTGG CGGATGACTGG']	3096	['GCGRAGAGGTGG CGGATGACTGG']
NM_000076.2(CDKN1C):c.*5 +2T>C	1028	CDKN 1C	2676	['GCGCAAGAG GCTGCGGTGA GCCAAGYGAG TACAGCGCACC TGGGGGGGCG C']	2864 - 2866	['CCAAGYGAGTAC AGCGCACCTGG', 'CAAGYGAGTACA GCGCACCTGGG', 'AAGYGAGTACAG CGCACCTGGGG']	3097 - 3099	['CCAAGYGAGTAC AGCGCACCTGG', 'CAAGYGAGTACA GCGCACCTGGG', 'AAGYGAGTACAG CGCACCTGGGG']
NC_012920.1: m.9478T>C	4514	MT- CO3	2677	['ATAATCCTAT TTATTACCTCA GAAGYTTTTTT CTTCGCAGGAT TTTTCTGA']	2867	['TCAGAAGYTTTTT TCTTCGCAGG']	3100	['TCAGAAGYTTTTT TCTTCGCAGG']
NM_002049.3(GATA1):c.2T> C (p.Met1Thr)	2623	GATA 1	2678	['CGCAGGTAA TCCCCAGAGGC TCCAYGGAGTT CCCTGGCCTGG GGTCCCTG']	2868 - 2869	['TCCAYGGAGTTC CCTGGCCTGGG', 'CCAYGGAGTTCCC TGGCCTGGGG']	3101 - 3103	['CTCCAYGGAGTT CCCTGGCCTGG', 'TCCAYGGAGTTC CCTGGCCTGGG', 'CCAYGGAGTTCC CTGGCCTGGGG']
NM_005740.2(DNAL4):c.153 +2T>C	1012 6	DNAL 4	2679	['GAGAAATTCT CCAACAACAA CGAGGYATTG	2870	['CGAGGYATTGCC AGCAGTGCAGG']	3104	['CGAGGYATTGCC AGCAGTGCAGG']

				CCAGCAGTGC AGGCGGCCCT ']				
NM_00128722 3.1(SCN11A):c .1142T>C (p.Ile381Thr)	1128 0	SCN11 A	2680	['GGGCTCTACT CAGTCTTCTTC TTCAYTGTGGT CATTTTCCTGG GCTCCTTC']	2871	['TTCAYTGTGGTCA TTTTCTGGG']	3105 - 3106	['CTTCAYTGTGGT CATTTTCCTGG', 'TTCAYTGTGGTCA TTTTCTGGG']
NM_00130294 6.1(TRNT1):c.4 97T>C (p.Leu166Ser)	5109 5	TRNT1	2681	['TAATGAATAG GTTTTGATGGC ACTTYATTTGA CTACTTTAATG GTTATGAA']	2872	['ACTTYATTTGACT ACTTTAATGG']	3107	['ACTTYATTTGAC TACTTTAATGG']
NM_178151.2(DCX):c.2T>C (p.Met1Thr)	1641	DCX	2682	['AGGTCTCTGA GGTCCACCAA AATAYGGAAC TTGATTTTGGGA CACTTTGAC']	2873	['CAAAATAYGGAA CTTGATTTTGG']	3108	['CAAAATAYGGA ACTTGATTTTGG']
NM_000169.2(GLA):c.758T> C (p.Ile253Thr)	-1	-	2683	['TGGACATCTT TTAACCAGGA GAGAAYTGTT GATGTTGCTGG ACCAGGGGGT']	2874	['GAGAGAAYTGTT GATGTTGCTGG']	3109	['GAGAGAAYTGTT GATGTTGCTGG']
NM_170707.3(LMNA):c.710T >C (p.Phe237Ser)	4000	LMNA	2684	['ATTGACAATG GGAAGCAGCG TGAGTYTGAG AGCCGGCTGG CGGATGCGCTG ']	2875	['TGAGTYTGAGAG CCGGCTGGCGG']	3110	['TGAGTYTGAGAG CCGGCTGGCGG']
NM_000256.3(MYBPC3):c.33 30+2T>C	4607	MYBP C3	2685	['CAGAAAGCCG ACAAGAAGAC CATGGBGAGC CCAGGGTCTGG GGTCCCCACG']	2876 - 2878	['ACCATGGBGAGC CCAGGGTCTGG', 'CCATGGBGAGCCC AGGGTCTGGG', 'CATGGBGAGCCCA GGGTCTGGGG']	3111 - 3113	['ACCATGGBGAGC CCAGGGTCTGG', 'CCATGGBGAGCC CAGGGTCTGGG', 'CATGGBGAGCCC AGGGTCTGGGG']
NM_005957.4(MTHFR):c.153 0+2T>C	4524	MTHF R	2686	['AGCGGGGGCT ATGTCTTCCAG AAGGYGTGGT AGGGAGGCAC GGGGTGCCCC']	2879 - 2881	['GAAGGYGTGGTA GGGAGGCACGG', 'AAGGYGTGGTAG GGAGGCACGGG', 'AGGYGTGGTAGG GAGGCACGGGG']	3114 - 3116	['GAAGGYGTGGTA GGGAGGCACGG', 'AAGGYGTGGTAG GGAGGCACGGG', 'AGGYGTGGTAGG GAGGCACGGGG']
NM_000264.3(PTCH1):c.3168 +2T>C	5727	PTCH1	2687	['AACCCCTGGA CGGCCGGGAT CATTGYGAGTG	2882 - 2884	['ATCATTGYGAGT GTATTATAAGG', 'TCATTGYGAGTGT	3117 - 3119	['ATCATTGYGAGT GTATTATAAGG', 'TCATTGYGAGTGT

				TATTATAAGGG GCTTTGTGG']		ATTATAAGGG', 'CATTGYGAGTGTA TTATAAGGGG']		ATTATAAGGG', 'CATTGYGAGTGT ATTATAAGGGG']
NM_000030.2(AGXT):c.322T >C (p.Trp108Arg)	189	AGXT	2688	['CTTCCTGGTT GGGGCCAATG GCATTYGGGG GCAGCGAGCC GTGGACATCG G']	2885	['CATTYGGGGGCA GCGAGCCGTGG']	3120	['CATTYGGGGGCA GCGAGCCGTGG']
NM_000023.2(SGCA):c.371T >C (p.Ile124Thr)	6442	SGCA	2689	['ACTCGGCAGA GGCTGGTGCTG GAGAYTGGGG ACCCAGAAGG TACCTCTAGC']	2886	['CTGGAGAYTGGG GACCCAGAAGG']	3121	['CTGGAGAYTGGG GACCCAGAAGG']
NM_001103.3(ACTN2):c.683 T>C (p.Met228Thr)	88	ACTN 2	2690	['GAGAAGCACCC TGGATATTCCT AAAAYGTTGG ATGCTGAAGGT GAGATGAAA']	2887	['CCTAAAAYGTTG GATGCTGAAGG']	3122	['CCTAAAAYGTTG GATGCTGAAGG']
NM_00116596 3.1(SCN1A):c. 4055T>C (p.Leu1352Pro)	-1	-	2691	['ATTCCATCCA TCATGAATGTG CTTCYGGTTTG TCTTATATTCT GGCTAATT']	2888	['TTCYGGTTTGTCT TATATTCTGG']	3123	['TTCYGGTTTGTCT TTATATTCTGG']
NM_00116596 3.1(SCN1A):c. 1265T>C (p.Val422Ala)	6323	SCN1A	2692	['CTAATAAATT TGATCCTGGCT GTGGHGGCCA TGGCCTACGAG GAACAGAAT']	2889	['TGTGGHGGCCAT GGCCTACGAGG']	3124	['TGTGGHGGCCAT GGCCTACGAGG']
NM_000426.3(LAMA2):c.828 2T>C (p.Ile2761Thr)	3908	LAMA 2	2693	['GCAGAATCAG AACCAGCTCTT TTGAYAGGGA GCAAGCAGTTC GGGCTTTCA']	2890 - 2891	['TTGAYAGGGAGC AAGCAGTTCGG', 'TGAYAGGGAGCA AGCAGTTCGGG']	3125 - 3126	['TTGAYAGGGAGC AAGCAGTTCGG', 'TGAYAGGGAGCA AGCAGTTCGGG']
NM_000257.3(MYH7):c.5117 T>C (p.Leu1706Pro)	-1	-	2694	['TCCCGGAAGC TGGCGGAGCA GGAGCYGATT GAGACTAGTG AGCGGGTGCA G']	2892	['AGCYGATTGAGA CTAGTGAGCGG']	3127	['AGCYGATTGAGA CTAGTGAGCGG']
NM_001399.4(EDA):c.396+2 T>C	1896	EDA	2695	['TCTGACTCCC AGGACGGGCA CCAGGKGAGT CACCTAGTAGG']	2893 - 2894	['ACCAGGKGAGTC ACCTAGTAGGG', 'CCAGGKGAGTCAC CTAGTAGGGG']	3128 - 3130	['CACCTAGTAGG', 'ACCAGGKGAGTC ACCTAGTAGGG',

				GGCGGCGGCG']				'CCAGGKGAGTCA CCTAGTAGGGG']
NM_001848.2(COL6A1):c.95 7+2T>C	1291	COL6 A1	2696	['TCCAGGGGAC CCAAGGGCTA CAAGGYGAGC GTGGGCTGCTG GGAGGGGGGA']	2895 - 2896	['ACAAGGYGAGCG TGGGCTGCTGG', 'CAAGGYGAGCGT GGGCTGCTGGG']	3131 - 3132	['ACAAGGYGAGC GTGGGCTGCTGG', 'CAAGGYGAGCGT GGGCTGCTGGG']
NM_000238.3(KCNH2):c.194 5+6T>C	3757	KCNH 2	2697	['CTGCGTCATG CTCATTGGCTG TGAGYGTGCC AGGGGCGGGC GGCGGGGAG']	2897 - 2898	['CTGTGAGYGTGC CCAGGGGCGGG', 'TGAGYGTGCCAG GGGCGGGCGG']	3133 - 3134	['CTGTGAGYGTGC CCAGGGGCGGG', 'TGAGYGTGCCA GGGGCGGGCGG']
NM_021007.2(SCN2A):c.1271 T>C (p.Val424Ala)	6326	SCN2A	2698	['CTAATAAATT TGATCTTGCT GTGGYGGCCA TGGCCTATGAG GAACAGAAT']	2899	['TGTGGYGGCCAT GGCCTATGAGG']	3135	['TGTGGYGGCCAT GGCCTATGAGG']
NM_021007.2(SCN2A):c.4308 +2T>C	6326	SCN2A	2699	['TATGCAGCTG TTGATTCACGA AATGYAAGTCT AGTTAGAGGG AAATTGTTT']	2900 - 2901	['CGAAATGYAAGT CTAGTTAGAGG', 'GAAATGYAAGTCT AGTTAGAGGG']	3136 - 3137	['CGAAATGYAAGT CTAGTTAGAGG', 'GAAATGYAAGTC TAGTTAGAGGG']
NM_000083.2(CLCN1):c.1283 T>C (p.Phe428Ser)	1180	CLCN1	2700	['CCCCGCGAAG CCATCAGTACT TTGTYTGACAA CAATACATGG GTGAAACAC']	2902 - 2903	['CTTTGTYTGACAA CAATACATGG', 'TTTGTYTGACAAC AATACATGGG']	3138 - 3139	['CTTTGTYTGACA ACAATACATGG', 'TTTGTYTGACAAC AATACATGGG']
NM_004550.4(NDUFS2):c.87 5T>C (p.Met292Thr)	4720	NDUF S2	2701	['CATTATGCTC TCCACAGTGGA GTGAYGCTTCG GGGCTCAGGC ATCCAGTGG']	2904	['GGAGTGAYGCTT CGGGGCTCAGG']	3140	['GGAGTGAYGCTT CGGGGCTCAGG']
NM_000546.5(TP53):c.584T> C (p.Ile195Thr)	7157	TP53	2702	['CACACGCAAA TTTCCTCCAC TCGGRTAAGAT GCTGAGGAGG GGCCAGACC']	2905 - 2906	['CTCGGRTAAGAT GCTGAGGAGGG', 'TCGGRTAAGATGC TGAGGAGGGG']	3141 - 3143	['ACTCGGRTAAGA TGCTGAGGAGG', 'CTCGGRTAAGAT GCTGAGGAGGG', 'TCGGRTAAGATG CTGAGGAGGGG']

Table 3. A to G with NGG PAM. Table 2 shows a list of A to G mutations that may be corrected using any of the base editors provided herein. GRNAs and gRNAall indicate the protospacer and PAM sequence, where the PAM sequence is the last 3 nucleotides of each of the sequences in GRNAs and gRNAall.

Name	Gene ID	Gene Symbol	SEQ ID NO:	Flanks	SEQ ID NO:	gRNAs	SEQ ID NO:	gRNAall
NM_017547.3(FOXRED1):c.1289A>G (p.Asn430Ser)	5557 2	FOXRE D1	5084	['GTGGGCCCCACC CGCTAGTTGTCAVC ATGTACTTTGCTACT GGCTTCAGT']	5261	['CCACCCGCTAGT TGTCAVCATGT']	5464- 5466	['CCCACCCGCTAG TTGTCAVCATG', 'CCACCCGCTAGTT GTCAVCATGT', 'CCCGCTAGTTGTC AVCATGTACT']
NM_000071.2(CBS):c.1150A>G (p.Lys384Glu)	875	CBS	5085	['GGTGA TCCCCAT CCCCGAGGACC RAGTTCTCTGAG CGACAGGTGGAT GCT']	5262	['CCCCATCCCGCA GGACC RAGTTTC']	5467- 5470	['CCCCATCCCGC AGGACC RAGTT', 'CCCCATCCCGCA GGACC RAGTTTC', 'CCCATCCCGCAG GACC RAGTTTC', 'CCATCCCGCAGG ACC RAGTTCT']
NM_000552.3(VWF):c.2384A>G (p.Tyr795Cys)	7450	VWF	5086	['GAGTGTACCAAAA CGTGCCAGAACTRT GACCTGGAGTGCAT GAGCATGGGC']	5263	['CCAAAACGTGCC AGAACTRTGAC']	5471	['CCAAAACGTGCC AGAACTRTGAC']
NM_000552.3(VWF):c.1583A>G (p.Asn528Ser)	7450	VWF	5087	['ACCTGCGCCTGT GTGGGAATTACART GGCAACCAGGGCGA CGACTTCCTT']	5264	['CCTGTGTGGGAA TTACARTGGCA']	5472	['CCTGTGTGGGAA TTACARTGGCA']
NM_000308.2(CTSA):c.1238A>G (p.Tyr413Cys)	5476	CTSA	5088	['CTTTAGAAATACC AGATCCTATTATRTA ATGGAGATGTAGAC ATGGCCTGC']	5265	['CCAGATCCTATT ATRTAATGGAG']	5473	['CCAGATCCTATT ATRTAATGGAG']
NM_000277.1(PAH):c.916A>G (p.Ile306Val)	5053	PAH	5089	['TTCTATTTCCCC AATTACAGGAARTT GGCCTTGCCCTCTCTG GGTGCACC']	5266	['CCCCAATTACA GGAARTTGGCC']	5474- 5476	['CCCCAATTACA GGAARTTGGCC', 'CCCCAATTACAG GAARTTGGCCT', 'CCAATTACAGG AARTTGGCCTT']
NM_000512.4(GALNS):c.1460A>G (p.Asn487Ser)	2588	GALNS	5090	['TTGGTCCCCGCGCA GCCCCAGCTCARCG TGTGCAACTGGGCG GTCATGGTA']	5267	['CCGCGCAGCCCC AGCTCARCGTG']	5477	['CCGCGCAGCCCC AGCTCARCGTG']
NM_013319.2(UBIAD1):c.305A>G (p.Asn102Ser)	2991 4	UBIAD 1	5091	['GTGCACGGGGCCG GTAATTTGGTCARC ACTTACTATGACTTT TCCAAGGGC']	5268	['CCGGTAATTTGG TCARCACTTAC']	5478	['CCGGTAATTTGG TCARCACTTAC']
NM_013319.2(UBIAD1):c.305A>G (p.Asn102Ser)	2991	UBIAD	5092	['AGCACCGAGGCCA	5269	['CCATTCTCCATT	5479	['CCATTCTCCATT

BIADI):c.695A>G (p.Asn232Ser)	4	1		TTCTCCATTCCARCAACACCAGGGACATGGAGTCCGAC']		CCARCAACACC']		CCARCAACACC']
NM_000275.2(OCA2):c.1465A>G (p.Asn489Asp)	4948	OCA2	5093	['TGCCACTGCCATCGGGGACCCTCCARATGTCATTATTGTTTCCAACCAAGA']	5270	['CCATCGGGGACCCTCCARATGTC']	5480	['CCATCGGGGACCCTCCARATGTC']
NM_001127255.1(NLRP7):c.2738A>G (p.Asn913Ser)	-1	-	5094	['CTCACAAACCTGGACTTGAGTATCARCAGATAGCTCGTGGATTGTGGATT']	5271	['CCTGGACTTGAGTATCARCCAGA']	5481	['CCTGGACTTGAGTATCARCCAGA']
NM_152783.4(D2HGDH):c.1315A>G (p.Asn439Asp)	7282 94	D2HG DH	5095	['TGCCCTTGTCCTCCAGGAGATGGTRACCTGCACCTCAATGTGACGGCGGA']	5272	['CCTCCAGGAGATGGTRACCTGCA']	5482- 5483	['CCCTCCAGGAGATGGTRACCTGC', 'CCTCCAGGAGATGGTRACCTGCA']
NM_022132.4(MCCC2):c.1309A>G (p.Ile437Val)	6408 7	MCCC 2	5096	['TGTGGCCTGTGCCC AAGTGCCTAAGDTA ACCCTCATCATTGGGGCTCCTA']	5273	['CCCAAGTGCCTAAGDTAACCCCTC']	5484	['CCCAAGTGCCTAAGDTAACCCCTC']
NM_000022.2(ADA):c.219-2A>G	100	ADA	5097	['TTCCCAACCCCTTTCTTCCCTTCCCRGGGGCTGCCGGGAGGCTATCAAAAAG']	5274	['CCCCCTTCTTCCCCTTCCCRGGGG']	5485- 5487	['CCCCCTTCTTCCCCTTCCCRGGGG', 'CCCTTCTTCCCTTCCCRGGGGC', 'CCTTCTTCCCTTCCCRGGGGCT']
NM_017780.3(CHD7):c.3082A>G (p.Ile1028Val)	5563 6	CHD7	5098	['TTTAGTAATTGCCCCATTGTCCACARTCCCAACTGGGAAAGGGAATTCCG']	5275	['CCCCATTGTCCACARTCCCAAC']	5488	['CCCCATTGTCCACARTCCCAAC']
NM_000483.4(APOC2):c.1A>G (p.Met1Val)	-1	-	5099	['TCAATGTTCCAGGTCTCTGGACACTRTGGCACACGACTCCTCCCAGCTCT']	5276	['CCAGGTCTCTGGACACTRTGGGC']	5489	['CCAGGTCTCTGGACACTRTGGGC']
NM_000391.3(TPP1):c.887-10A>G	1200	TPP1	5100	['TGTCCCTCATGCCGGCCTGGATTTTYTTTTTTTTTTTTTTTGGATGGG']	5277	['CCGGCCTGGATTTTYTTTTTTTT']	5490	['CCGGCCTGGATTTTYTTTTTTTT']
NM_017890.4(VPS13B):c.8978A>G (p.Asn2993Ser)	1576 80	VPS13 B	5101	['CTTCTGCCCTGGGCCCTGCTTATCARTGAATCCAAATGGGACCTCTGGCTA']	5278	['CCTGGGCCCTGCTTATCARTGAA']	5491	['CCTGGGCCCTGCTTATCARTGAA']
NM_000226.3(KRT9):c.482A>G (p.Asn161Ser)	3857	KRT9	5102	['GAGAAGAGCACCATGAGGAACTCADTTCTCGGCTGGCCTCT	5279	['CCATGCAGGAAC TCADTTCTCGG']	5492	['CCATGCAGGAAC TCADTTCTCGG']

				TACTTGGAT']				
NM_000529.2(MC2R):c.761A>G (p.Tyr254Cys)	4158	MC2R	5103	['CCAAGTAACCCCTACTGCGCCTGCTRCATGTCTCTCTTCCAGGTGAACGGC']	5280 - 5281	['CCCTACTGCGCC TGCTRCATGTC', 'CCTACTGCGCCTGCTRCATGTCT']	5493-5495	['CCCCTACTGCGCCTGCTRCATGT', 'CCCTACTGCGCCTGCTRCATGTC', 'CCTACTGCGCCTGCTRCATGTCT']
NM_005957.4(MTHFR):c.971A>G (p.Asn324Ser)	4524	MTHFR	5104	['CCAGGCCTCCACTTCTACACCCCTCARCCGCGAGATGGCTACCACAGAGGTG']	5282	['CCTACTTCTACACCCTCARCCGCG']	5496	['CCTACTTCTACACCCTCARCCGCG']
NM_000403.3(GALE):c.101A>G (p.Asn34Ser)	2582	GALE	5105	['GGTACTTGCTGTGGTCATCGATARCTTCCATAATGCCTTCCGTGGTGAG']	5283	['CCTGTGGTCATCGATARCTTCCA']	5497	['CCTGTGGTCATCGATARCTTCCA']
NM_000356.3(TCOF1):c.149A>G (p.Tyr50Cys)	6949	TCOF1	5106	['CAGCCCGTAACCC TTCTGGACATCTRACACTGGCAACAGTAAGTGGTG']	5284 - 5285	['CCCTTCTGGACATCTRTRACACAC', 'CCTTCTGGACATCTRTRACACACT']	5498-5499	['CCCTTCTGGACATCTRTRACACAC', 'CCTTCTGGACATCTRTRACACACT']
NM_012464.4(TLL1):c.1885A>G (p.Ile629Val)	7092	TLL1	5107	['ACTTCTTACCAAAC TTAACGGCACCRTAACCACCCCTGGCTGGCCAAAGGA']	5286	['CCAAACTTAACGGCACCRTAACC']	5500	['CCAAACTTAACGGCACCRTAACC']
NM_000112.3(SLC26A2):c.1273A>G (p.Asn425Asp)	1836	SLC26A2	5108	['GAAATGTATGCCATTGGCTTTTGTRATATCATCCCTTCTTCTTCCACTG']	5287	['CCATTGGCTTTTGTRATATCATC']	5501	['CCATTGGCTTTTGTRATATCATC']
NM_000157.3(GBA):c.680A>G (p.Asn227Ser)	2629	GBA	5109	['ACATCACCCACTTGCTCAAGACCARTGAGCGGTGAATGGGAAGGGGTCA']	5288	['CCTACTGGCTCAAGACCARTGGA']	5502	['CCTACTGGCTCAAGACCARTGGA']
NM_175073.2(APTX):c.602A>G (p.His201Arg)	54840	APTX	5110	['GATAAATACCCAAAGGCCCGTTACCRTTGGCTGGTCTTACGTGGACCTCC']	5289 - 5290	['CCCAAAGGCCCGTTACCRTTGGC', 'CCAAAGGCCCGTTACCRTTGGCT']	5503-5504	['CCCAAAGGCCCGTTACCRTTGGC', 'CCAAAGGCCCGTTACCRTTGGCT']
NM_020638.2(FGF23):c.211A>G (p.Ser71Gly)	8074	FGF23	5111	['TGGCGCACCCCATCAGACCATCTACRGTGAGTAGGGCTTCA GGCTGGGAAG']	5291	['CCCCATCAGACCATCTACRGTGA']	5505-5507	['CCCCATCAGACCATCTACRGTGA', 'CCCATCAGACCACTACRGTGAG', 'CCATCAGACCATCTACRGTGAGT']
NM_021102.3(SPINT2):c.488A>G (p.Tyr163Cys)	10653	SPINT2	5112	['AGGAACTCCTGCAATAACTTCATCTRTGGAGGCTGCCGGGGC']	5292	['CCTGCAATAACTTCATCTRTGGA']	5508	['CCTGCAATAACTTCATCTRTGGA']

				AATAAGAAC']				
NM_004795.3(KL):c.578A>G (p.His193Arg)	9365	KL	5113	['GTGCAGCCCCTGGTCACCCTGTACCRCTGGGACCTGCCCCAGCGCTGCAG']	5293	['CCGTGGTCACCCGTGACCRCTGG']	5509	['CCGTGGTCACCCGTGACCRCTGG']
NM_012193.3(FZD4):c.766A>G (p.Ile256Val)	-1	-	5114	['GTTTTCTACCCTGAGCGCCCCATCRTATTTCTCAGTATGTGCTATAATAT']	5294 - 5295	['CCCTGAGCGCCCCATCRTATTTCT', CCTGAGCGCCCCATCRTATTTCT']	5510- 5511	['CCCTGAGCGCCCCATCRTATTTCT', CCTGAGCGCCCCATCRTATTTCT']
NM_001099274.1(TINF2):c.838A>G (p.Lys280Glu)	2627 7	TINF2	5115	['ATGGGCTCCACTAGGGGAGGCCATDAGGAGCGCCCCACAGTCATGCTGTT']	5296	['CCACTAGGGGAGGCCATDAGGAG']	5512	['CCACTAGGGGAGGCCATDAGGAG']
NM_005682.6(ADGRG1):c.263A>G (p.Tyr88Cys)	9289	ADGR GI	5116	['TCCTTCCCTGACCCAGGGGCCTCTRCCACTTCTGCCTCTACTGGAACCGA']	5297	['CCCAGGGGCCTCTRCCACTTCT']	5513	['CCCAGGGGCCTCTRCCACTTCT']
NM_000369.2(TSHR):c.1856A>G (p.Asp619Gly)	7253	TSHR	5117	['CCGCAGTACAACCAGGGGACAAAGRTACCAAAATTGCCAAGAGGATGGCT']	5298	['CCCAGGGGACAAAGRTACCAAAA']	5514	['CCCAGGGGACAAAGRTACCAAAA']
NM_024009.2(GJB3):c.497A>G (p.Asn166Ser)	2707	GJB3	5118	['ATGCCGCGCCTGGTGAGTGTGCCADCGTGGCCCCCTGCCCAACATCGTG']	5299	['CCTGGTGCAGTGTGCCADCGTGG']	5515	['CCTGGTGCAGTGTGCCADCGTGG']
NM_003722.4(TP63):c.697A>G (p.Lys233Glu)	8626	TP63	5119	['TATCCGCGCCATGCTGTCTACAAARAAAGCTGAGCACGTCACGGAGGTGGT']	5300	['CCATGCCTGTCTACAAARAAGCT']	5516	['CCATGCCTGTCTACAAARAAGCT']
NM_003494.3(DYSF):c.3443-33A>G	8291	DYSF	5120	['CAGCTCTTAACCACCTCCAGCCACTCRCTCTGGCACCTCTGTTTTTCCCTT']	5301	['CCACTCCAGCCACTCRCTCTGGC']	5517	['CCACTCCAGCCACTCRCTCTGGC']
NM_003494.3(DYSF):c.1285-2A>G	8291	DYSF	5121	['AACTTGTCCCCTCCCTGTGTCTTCTRGCTGTGCAGCAAGATCTTGGAGAAG']	5302	['CCCCTCCCTGTGTCTTCTRGCTG']	5518- 5520	['CCCCTCCCTGTGTCTTCTRGCTG', CCCTCCCTGTGTCTTCTRGCTGT', CCTCCCTGTGTCTTCTRGCTGTG']
NM_002408.3(MGAT2):c.785A>G (p.His262Arg)	4247	MGAT 2	5122	['CTTATACTTTTCTTCTAGAAGAGGATCRCTACTTAGCCCCAGACTTTTACCAT']	5303	['CCTAGAAGAGGATCRCTACTTAG']	5521	['CCTAGAAGAGGATCRCTACTTAG']
NM_000492.3(CFTR)	1080	CFTR	5123	['GTGATTATCACCA	5304	['CCAGCACCAGTT	5522	['CCAGCACCAGTT

(TR):c.2738A>G (p.Tyr913Cys)				GCACCAGTTCGTRT TATGTGTTTTACATT TACGTGGGA']		CGTRTTATGTG']		CGTRTTATGTG']
NM_001814.4(CT SC):c.857A>G (p.Gln286Arg)	1075	CTSC	5124	['TCTCAGACCCCAAT CCTAAGCCCTCRGG AGGTTGTGTCTTGTA GCCAGTAT']	5305	['CCCCAATCCTAA GCCCTCRGGAG']	5523- 5525	['CCCCAATCCTAA GCCCTCRGGAG', 'CCCAATCCTAAG CCCTCRGGAGG', 'CCAATCCTAAGC CCTCRGGAGGT']
NM_005144.4(H R):c.-218A>G	5580 6	HR	5125	['TCCGACCCCTCCAA CCTGCGGCCCTRGA GCGCCCCCGCCGCC CCGGGGGAA']	5306	['CCTCCAACCTGC GGCCCTRAGAC']	5526- 5527	['CCTCCAACCTGC GGCCCTRAGAC', 'CCAACCTGCGGC CCTRAGAGCGCC']
NM_018488.2(TB X4):c.1592A>G (p.Gln531Arg)	9496	TBX4	5126	['TCCTTGTCCCGAGA ATCTTCCTTACRGTA CCATTACAGGAATGG GGACTGTG']	5307	['CCCGAGAATCTT CCTTACRGTAC']	5528- 5529	['CCCGAGAATCTT CCTTACRGTAC', 'CCGAGAATCTTCC TTACRGTACC']
NM_001089.2(A BCA3):c.1702A> G (p.Asn568Asp)	21	ABCA3	5127	['ACAGATCACCGTC CTGCTGGGCCACRA CGGTGCCGGGAAGA CCACCACCCT']	5308	['CCGTCCTGCTGG GCCACRACGGT']	5530	['CCGTCCTGCTGG GCCACRACGGT']
NM_000525.3(K CNJ11):c.776A> G (p.His259Arg)	3767	KCNJ1 1	5128	['CTGGTGGCCCCGCT GATCATCTACCRGT TCATTGATGCCAAC AGCCCACTC']	5309 - 5310	['CCCCGCTGATCA TCTACCRGTGC', 'CCCGCTGATCATC TACCRGTGCA']	5531- 5533	['CCCCGCTGATCA TCTACCRGTGC', 'CCCGCTGATCATC TACCRGTGCA', 'CCGCTGATCATCT ACCRGTGCAT']
NM_005587.2(M EF2A):c.788A>G (p.Asn263Ser)	4205	MEF2A	5129	['TCTCCCCCTCCACC AGGTGGTGGTARTC TTGGAATGAACAGT AGGAAACCA']	5311	['CCACCAGGTGGT GGTARTCTTGG']	5534	['CCACCAGGTGGT GGTARTCTTGG']
NM_000098.2(CP T2):c.359A>G (p.Tyr120Cys)	1376	CPT2	5130	['TTTTTAGGACCCTG GTTTGATATGTRCCT ATCTGCTCGAGACT CCGTTGTT']	5312	['CCTGGTTTGATA TGTRCCTATCT']	5535- 5536	['CCCTGGTTTGAT ATGTRCCTATC', 'CCTGGTTTGATAT GTRCCTATCT']
NM_178138.4(L HX3):c.332A>G (p.Tyr111Cys)	8022	LHX3	5131	['GTGCGCCGCGCCC AGGACTTCGTGTRC CACCTGCACTGCTTT GCCTGCGTC']	5313 - 5314	['CCAGGACTTCG TGTRCCACCTG', 'CCAGGACTTCGT GTRCCACCTGC']	5537- 5538	['CCAGGACTTCG TGTRCCACCTG', 'CCAGGACTTCGT GTRCCACCTGC']
NM_005502.3(A BCA1):c.2804A> G (p.Asn935Ser)	19	ABCA1	5132	['CAGATCACCTCCTT CCTGGGCCACARTG GAGCGGGGAAGAC GACCACCATG']	5315	['CCTCCTCCTGG GCCACARTGGA']	5539- 5540	['CCTCCTCCTGG GCCACARTGGA', 'CCTCCTGGGCCA CARTGGAGCG']
m.3260A>G	4567	MT-	5133	['GATGGCAGAGCCC	5316	['CCCGGTAATCGC	5541-	['CCCGGTAATCGC

		TLI		GGTAATCGCATARA ACTTAAACTTTAC AGTCAGAGGT']	- 5317	ATARAACCTTAA', 'CCGGTAATCGCA TARAACCTTAAA']	5542	ATARAACCTTAA', 'CCGGTAATCGCA TARAACCTTAAA']
m.4269A>G	4565	MT-TI	5134	['GCATTCCCCCTCAA ACCTAAGAAATRTG TCTGATAAAAGAGT TACTTTGAT']	5318 - 5319	['CCCTCAAACCTA AGAAATRTGTC', 'CCTCAAACCTAA GAAATRTGTCT']	5543- 5544	['CCCTCAAACCTA AGAAATRTGTC', 'CCTCAAACCTAA GAAATRTGTCT']
m.14495A>G	4541	MT- ND6	5135	['TCCAAAGACAACC ATCATTCCCCCTRA ATAAATTAATAAAA CTATTAACC']	5320	['CCATCATTCCCC CTRAATAAATT']	5545	['CCATCATTCCCC CTRAATAAATT']
NM_002764.3(PR PS1):c.341A>G (p.Asn114Ser)	5631	PRPS1	5136	['CCAATCTCAGCCA AGCTTGTTGCAART ATGCTATCTGTAGC AGGTGCAGAT']	5321	['CCAAGCTTGTTG CAARTATGCTA']	5546	['CCAAGCTTGTTG CAARTATGCTA']
NM_000054.4(A VPR2):c.614A>G (p.Tyr205Cys)	554	AVPR2	5137	['GCGGAGCCCTGGG GCCGTGCACCTRT GTCACCTGGATTGC CCTGATGGTG']	5322	['CCTGGGGCCGTC GCACCTRTGTC']	5547	['CCTGGGGCCGTC GCACCTRTGTC']
NM_000033.3(A BCD1):c.443A>G (p.Asn148Ser)	215	ABCD1	5138	['ATCGCCCTCCCTGC TACCTTCGTCARCA GTGCCATCCGTTAC CTGGAGGGC']	5323 - 5324	['CCCTGCTACCTT CGTCARCAGTG', 'CCTGCTACCTTCG TCARCAGTGC']	5548- 5549	['CCCTGCTACCTT CGTCARCAGTG', 'CCTGCTACCTTCG TCARCAGTGC']
NM_000061.2(BT K):c.1082A>G (p.Tyr361Cys)	695	BTK	5139	['AGCACCATCCCTG AGCTCATTAACTRC CATCAGCACAACCTC TGCAGGTGAG']	5325 - 5326	['CCCTGAGCTCAT TAACTRCCATC', 'CCTGAGCTCATT ACTRCCATCA']	5550- 5551	['CCCTGAGCTCAT TAACTRCCATC', 'CCTGAGCTCATT ACTRCCATCA']
NM_003413.3(ZI C3):c.1213A>G (p.Lys405Glu)	7547	ZIC3	5140	['CTACACGCACCCG AGCTCCCTGCGCRA ACACATGAAGGTAA TTACCTCTTT']	5327	['CCGAGCTCCCTG CGCRAACACAT']	5552- 5553	['CCCAGCTCCCT GCGCRAACACA', 'CCGAGCTCCCTG GCRAACACAT']
NM_005448.2(B MP15):c.704A>G (p.Tyr235Cys)	9210	BMP15	5141	['TTGGACATTGCCTT CTTGTTACTCTRTTT CAATGATACTCATA AAAGCATT']	5328	['CCTTCTTGTTACT CTRTTTCAAT']	5554	['CCTTCTTGTTACT CTRTTTCAAT']
NM_001363.4(D KC1):c.1069A>G (p.Thr357Ala)	1736	DKC1	5142	['ATTAATGACCACA GCGGTCATCTCTRC CTGCGACCATGGTA TAGTAGCCAA']	5329	['CCACAGCGGTCA TCTCTRCCTGC']	5555	['CCACAGCGGTCA TCTCTRCCTGC']
NM_000481.3(A MT):c.125A>G (p.His42Arg)	275	AMT	5143	['CGCAGGACACCGC TCTATGACTTCCRCC TGGCCCACGCGGG AAAATGGTG']	5330	['CCGCTCTATGAC TTCCRCCTGGC']	5556	['CCGCTCTATGAC TTCCRCCTGGC']
NM_003361.3(U	7369	UMOD	5144	['TGCCACGCCCTGG	5331	['CCTGGCCACATG	5557-	['CCCTGGCCACAT

MOD):c.383A>G (p.Asn128Ser)				CCACATGTGTCART GTGGTGGGCAGCTA CTTGTGCGTA']		TGTCARTGTGG']	5558	GTGTCARTGTG', 'CCTGGCCACATGT GTCARTGTGG']
NM_001382.3(DP AGT1):c.509A>G (p.Tyr170Cys)	1798	DPAG T1	5145	['TCTCTCCCCGAGG AATCCTGTACTRTGT CTACATGGGGCTGC TGGCAGTG']	5332	['CCGCAGGAATCC TGTACTRTGTC']	5559	['CCGCAGGAATCC TGTACTRTGTC']
NM_001128177.1 (THRB):c.1324A >G (p.Met442Val)	7068	THRB	5146	['CTGCCATGCCAGC CGCTTCCTGCACRT GAAGGTGGAATGCC CCACAGAACT']	5333	['CCAGCCGCTTCC TGCACRTGAAG']	5560	['CCAGCCGCTTCC TGCACRTGAAG']
NM_000141.4(FG FR2):c.874A>G (p.Lys292Glu)	2263	FGFR2	5147	['TGCCCAGCCCCAC ATCCAGTGGATCRA GCACGTGAAAAGA ACGGCAGTAA']	5334	['CCCACATCCAGT GGATCRAGCAC']	5561- 5563	['CCCCACATCCAG TGGATCRAGCA', 'CCCACATCCAGT GGATCRAGCAC', 'CCACATCCAGTG GATCRAGCACG']
NM_000371.3(TT R):c.401A>G (p.Tyr134Cys)	7276	TTR	5148	['ACCATTGCCGCCCT GCTGAGCCCCTRCT CCTATTCCACCACG GCTGTGCTC']	5335 - 5337	['CCGCCCTGCTGA GCCCCTRCTCC', 'CCCTGCTGAGCCC CTRCTCCTAT', 'CCTGCTGAGCCCC TRCTCCTATT']	5564- 5566	['CCGCCCTGCTGA GCCCCTRCTCC', 'CCCTGCTGAGCCC CTRCTCCTAT', 'CCTGCTGAGCCCC TRCTCCTATT']
NM_000371.3(TT R):c.379A>G (p.Ile127Val)	7276	TTR	5149	['CGACTCCGGCCCC CGCCGCTACACCRT TGCCGCCCTGCTGA GCCCCCTACTC']	5338	['CCCCCGCCGCTA CACRRTTGCCG']	5567- 5569	['CCCCCGCCGCTA CACRRTTGCCG', 'CCCCCGCCGCTAC ACRRTTGCCG', 'CCCCCGCTACA CCRRTTGCCG']
NM_000174.4(GP 9):c.182A>G (p.Asn61Ser)	2815	GP9	5150	['ACCCGCCACCTTCT GCTGGCCAACARCA GCCTTCAGTCCGTG CCCCCGGGA']	5339	['CCTTCTGCTGGC CAACARCAGCC']	5570	['CCTTCTGCTGGC CAACARCAGCC']
NM_000222.2(KI T):c.1924A>G (p.Lys642Glu)	3815	KIT	5151	['ACGGGAAGCCCTC ATGTCTGAACTCRA AGTCCTGAGTTACC TTGGTAATCA']	5340 - 5341	['CCCTCATGTCTG AACTCRAAGTC', 'CCTCATGTCTGAA CTCRAAGTCC']	5571- 5572	['CCCTCATGTCTG AACTCRAAGTC', 'CCTCATGTCTGAA CTCRAAGTCC']
NM_000530.6(M PZ):c.242A>G (p.His81Arg)	4359	MPZ	5152	['TCCCCTCATTCCCTC ATAGATCTTCCRCT ATGCCAAGGGACAA CCCTACATT']	5342	['CCTCATAGATCT TCCRCTATGCC']	5573	['CCTCATAGATCT TCCRCTATGCC']
NM_000233.3(L HCGR):c.1733A> G (p.Asp578Gly)	-1	-	5153	['AAAATGGCAATCC TCATCTTACCGRIT TCACCTGCATGGCA	5343	['CCTCATCTTAC CGRTTTCACCT']	5574	['CCTCATCTTAC CGRTTTCACCT']

				CCTATCTCT']				
NM_000421.3(KRT10):c.1374-2A>G	-1	-	5154	['CCGCCGCGTCCGC CGCTCCGGAACYA AACGGGGTGAGGTC ACATTCGGTT']	5344	['CCGCCGCCTCCG GAACYAAACGG']	5575	['CCGCCGCCTCCG GAACYAAACGG']
NM_000422.2(KRT17):c.274A>G (p.Asn92Asp)	3872	KRT17	5155	['TGAGAAGGCCACC ATGCAGAACCTCVA TGACCGCTGGCCT CCTACCTGGA']	5345	['CCACCATGCAGA ACCTCVATGAC']	5576- 5577	['CCACCATGCAGA ACCTCVATGAC', 'CCATGCAGAACC TCVATGACCGC']
NM_000422.2(KRT17):c.275A>G (p.Asn92Ser)	3872	KRT17	5156	['GAGAAGGCCACCA TGAGAACCTCART GACCGCCTGGCCTC CTACCTGGAC']	5346	['CCACCATGCAGA ACCTCARTGAC']	5578- 5579	['CCACCATGCAGA ACCTCARTGAC', 'CCATGCAGAACC TCARTGACCGC']
NM_000823.3(GHRHR):c.985A>G (p.Lys329Glu)	2692	GHRHR	5157	['TTGTCTTTCCTGCA GGCGTCTCTCCRAG TCGACACTTTTCCTG ATCCCACT']	5347	['CCTGCAGGCGTC TCTCCRAGTCG']	5580	['CCTGCAGGCGTC TCTCCRAGTCG']
NM_000407.4(GPIBB):c.338A>G (p.Tyr113Cys)	-1	-	5158	['GCCGGCCGCCCG AGCGTGCGCCCTDC CGCGACCTGCGTTG CGTGCGCCC']	5348 - 5349	['CCCCGAGCGTGC GCCCTDCCGCG', 'CCCGAGCGTGCG CCCTDCCGCGA']	5581- 5583	['CCCCGAGCGTGC GCCCTDCCGCG', 'CCCGAGCGTGCG CCCTDCCGCGA', 'CCGAGCGTGCGC CCTDCCGCGAC']
NM_001146040.1 (GLRA1):c.920A>G (p.Tyr307Cys)	2741	GLRA1	5159	['CCTCCACCCCACT CTAGGTGTCTVTGT GAAAGCCATTGACA TTTGATG']	5350 - 5351	['CCCCACTCTAGG TGTCTVTGTG', 'CCCACTCTAGGTG TCCTVTGTGA']	5584- 5586	['CCCCACTCTAGG TGTCTVTGTG', 'CCCACTCTAGGTG TCCTVTGTGA', 'CCACTCTAGGTGT CCTVTGTGAA']
NM_182925.4(FLT4):c.3104A>G (p.His1035Arg)	2324	FLT4	5160	['CGCCTCCCCGACC CCAGTGCATCCRCA GAGACCTGGCTGCT CGGAACATT']	5352	['CCGCACCCCACT GCATCCRCAGA']	5587	['CCGCACCCCACT GCATCCRCAGA']
NM_212482.1(FN1):c.2918A>G (p.Tyr973Cys)	2335	FN1	5161	['ACCGGGCTGTCCC CTGGGGTCACCTRT TACTTCAAAGTCTTT GCAGTGAGC']	5353 - 5354	['CCCCTGGGGTCA CCTRTACTTC', 'CCCTGGGGTCAC CCTRTACTTCA']	5588- 5589	['CCCCTGGGGTCA CCTRTACTTC', 'CCCTGGGGTCAC CCTRTACTTCA']
NM_000121.3(EPOR):c.1460A>G (p.Asn487Ser)	2057	EPOR	5162	['GGTTATCCGATG GCCCCTACTCCARC CCTTATGAGAACAG CCTTATCCCA']	5355	['CCGATGGCCCCT ACTCCARCCCT']	5590	['CCGATGGCCCCT ACTCCARCCCT']
NM_001735.2(C5):c.1115A>G (p.Lys372Arg)	727	C5	5163	['CGTCTACCCCTCA CCCAATCTACCYTG ATGGGATATGGAAT	5356 - 5357	['CCCCTACCCAA TCTACCYTGAT', 'CCCTACCCCAATC	5591- 5593	['CCCCTACCCAA TCTACCYTGAT', 'CCCTACCCCAATC

				CCCAGGCTT']		TACCYTGATG']		TACCYTGATG', 'CCTCACCCAATCT ACCYTGATGG']
NM_001844.4(COL2A1):c.4172A>G (p.Tyr1391Cys)	1280	COL2A1	5164	['ACGGAAGGCTCCC AGAACATCACCTRC CACTGCAAGAACAG CATTGCCTAT']	5358 - 5359	['CCAGAACATCA CCTRCCACTGC', 'CCAGAACATCAC CTRCCACTGCA']	5594- 5595	['CCAGAACATCA CCTRCCACTGC', 'CCAGAACATCAC CTRCCACTGCA']
NM_001904.3(CTNNB1):c.121A>G (p.Thr41Ala)	1499	CTNNB1	5165	['CTCTGGAATCCATT CTGGTGCCACTNCC ACAGCTCCTTCTCTG AGTGGTAA']	5360	['CCATTCTGGTGC CACTNCCACAG']	5596	['CCATTCTGGTGC CACTNCCACAG']
NM_000040.1(APOC3):c.280A>G (p.Thr94Ala)	345	APOC3	5166	['GGATTTGGACCCT GAGGTCAGACCARC TTCAGCCGTGGCTG CCTGAGACCT']	5361 - 5362	['CCCTGAGGTCAG ACCARCTTCAG', 'CCTGAGGTCAGA CCARCTTCAGC']	5597- 5598	['CCCTGAGGTCAG ACCARCTTCAG', 'CCTGAGGTCAGA CCARCTTCAGC']
NM_000488.3(SERPINC1):c.655A>G (p.Asn219Asp)	462	SERPINC1	5167	['TGCAGAGCAATCC AGAGCGGCCATCRA CAAATGGGTGTCCA ATAAGACCGA']	5363	['CCAGAGCGGCCA TCRACAAATGG']	5599	['CCAGAGCGGCCA TCRACAAATGG']
NM_001085.4(SERPINA3):c.1240A>G (p.Met414Val)	12	SERPINA3	5168	['TACAGACACCCAG AACATCTTCTTCRTG AGCAAAGTCACCAA TCCCAAGCA']	5364	['CCCAGAACATCT TCTTCRTGAGC']	5600- 5601	['CCCAGAACATCT TCTTCRTGAGC', 'CCAGAACATCTTC TTCRTGAGCA']
NM_001145.4(ANG):c.121A>G (p.Lys41Glu)	-1	-	5169	['CTTCTGACCCAGC ACTATGATGCCRAA CCACAGGGCCGGGA TGACAGATA']	5365	['CCAGCACTATGA TGCCRAACCAC']	5602- 5603	['CCAGCACTATG ATGCCRAACCA', 'CCAGCACTATGA TGCCRAACCAC']
NM_001100.3(ACCTA1):c.350A>G (p.Asn117Ser)	58	ACCTA1	5170	['GAGGCCCCCTCA ATCCCAAGGCCARC CGCGAGAAGATGAC CCAGATCATG']	5366	['CCTCAATCCCAA GGCCARCCGCG']	5604- 5605	['CCTCAATCCCAA AGGCCARCCGC', 'CCTCAATCCCAA GGCCARCCGCG']
NM_014053.3(FLVCR1):c.361A>G (p.Asn121Asp)	2898 2	FLVCR1	5171	['GATCTTCAGCCTGT ACTCGCTGGTCRAC GCCTTTCAGTGGAT CCAGTACAG']	5367	['CCTGTACTCGCT GGTCRACGCCT']	5606	['CCTGTACTCGCT GGTCRACGCCT']
NM_000334.4(SCN4A):c.4078A>G (p.Met1360Val)	6329	SCN4A	5172	['GAAGCAGGCCTTC GACATCACCATCRT GATCCTCATCTGCC CAACATGGT']	5368	['CCTTCGACATCA CCATCRTGATC']	5607	['CCTTCGACATCA CCATCRTGATC']
NM_004519.3(KCNQ3):c.1403A>G (p.Asn468Ser)	3786	KCNQ3	5173	['GAACCAAAGCCTG TTGGCTTAAACART AAAGAGCGTTTCCG CACGGCCTTC']	5369	['CCTGTTGGCTTA AACARTAAAGA']	5608	['CCTGTTGGCTTA AACARTAAAGA']
NM_007375.3(TARD)	2343	TARD	5174	['AATGCCGAACCTA	5370	['CCTAAGCACAAT	5609	['CCTAAGCACAAT

ARDBP):c.800A>G (p.Asn267Ser)	5	BP		AGCACAATAGCART AGACAGTTAGAAAG AAGTGGGAAGA']		AGCARTAGACA']		AGCARTAGACA']
NM_032520.4(GNPTG):c.610-2A>G	8457 2	GNPT G	5175	['TGCTGCCCCTGCAT CCTCCACCTTCRGG GCCATGAGAAGTTG CTGAGGACA']	5371	['CCTGCATCCTCC ACCTTCRGGGC']	5610	['CCTGCATCCTCC ACCTTCRGGGC']
NM_000495.4(COL4A5):c.466-2A>G	1287	COL4A 5	5176	['AGAACTTCCATTG ATGGCTTCTTTTRGG GTGAACCAGGTAGT ATAATTATG']	5372	['CCATTGATGGCT TCTTTTRGGGT']	5611	['CCATTGATGGCT TCTTTTRGGGT']
NM_000495.4(COL4A5):c.1340-2A>G	1287	COL4A 5	5177	['TTGCTATCCTTTCT TTATCTTACTCRGGT GATGAGATATGTGA ACCAGGCC']	5373	['CCTTTCTTTATCT TACTCRGGTG']	5612	['CCTTTCTTTATCT TACTCRGGTG']
NM_000060.3(BTD):c.278A>G (p.Tyr93Cys)	686	BTD	5178	['CTCATGAACCAGA ACCTTGACATCTRT GAACAGCAAGTGAT GACTGCAGCC']	5374	['CCAGAACCTTGA CATCTRTGAAC']	5613	['CCAGAACCTTGA CATCTRTGAAC']
NM_000060.3(BTD):c.641A>G (p.Asn214Ser)	686	BTD	5179	['CTTGTTGACCGCTA CCGTAAACACARCC TCTACTTTGAGGCA GCATTCGAT']	5375	['CCGCTACCGTAA ACACARCCTCT']	5614	['CCGCTACCGTAA ACACARCCTCT']
NM_000094.3(COL7A1):c.425A>G (p.Lys142Arg)	1294	COL7A 1	5180	['CAGCTGGCCCGAC CTGGTGTCGCCARG GTGATCCCTACCCC TACCATGCCT']	5376	['CCCGACCTGGTG TCCCCARGGTG']	5615- 5616	['CCCGACCTGGTG TCCCCARGGTG', 'CCGACCTGGGTGC CCCARGGTGA']
NM_005247.2(FGF3):c.146A>G (p.Tyr49Cys)	2248	FGF3	5181	['GGGGCGCCCCGGC GCCGCAAGCTCTRC TGCGCCACGAAGTA CCACCTCCAG']	5377 - 5378	['CCCGGCGCCGCA AGCTCTRCTGC', 'CCGGCGCCGCAA GCTCTRCTGCG']	5617- 5618	['CCCGGCGCCGCA AGCTCTRCTGC', 'CCGGCGCCGCAA GCTCTRCTGCG']
NM_000313.3(PROS1):c.701A>G (p.Tyr234Cys)	5627	PROS1	5182	['TGTGAATGCCCCG AAGGCTACAGATRT AATCTCAAATCAAA GTCTTGTGAA']	5379 - 5380	['CCCGAAGGCTAC AGATRTAATCT', 'CCGAAGGCTACA GATRTAATCTC']	5619- 5621	['CCCGAAGGCTA CAGATRTAATC', 'CCCGAAGGCTAC AGATRTAATCT', 'CCGAAGGCTACA GATRTAATCTC']
NM_004612.3(TGFBR1):c.134A>G (p.Asn45Ser)	7046	TGFBR 1	5183	['TTCTGCCACCTCTG TACAAAAGACARTT TTACTTGTGTGACA GATGGGCTC']	5381	['CCTCTGTACAAA AGACARTTTTA']	5622	['CCTCTGTACAAA AGACARTTTTA']
m.608A>G	4558	MT-TF	5184	['GTAGCTTACCTCCT CAAAGCAATACRCT GAAAATGTTTAGAC	5382	['CCTCCTCAAAGC AATACRCTGAA']	5623- 5624	['CCTCCTCAAAGC AATACRCTGAA', 'CCTCAAAGCAAT

				GGGCTCACAA']				ACRCTGAAAAT']
NM_001376.4(DYNC1H1):c.2909A>G (p.Tyr970Cys)	1778	DYNC1H1	5185	['CTAAGAATAACCAATCAGGTAATCTRCTTGAATCCACCAATTGAAAGAGTGC']	5383	['CCAATCAGGTAACTCTRCTTGAAT']	5625	['CCAATCAGGTAACTCTRCTTGAAT']
NM_000459.4(TEK):c.2690A>G (p.Tyr897Cys)	7010	TEK	5186	['ATGCTCTCTTCCTTCCCTCCAGGCTVCTTGTACCTGGCCATTGAGTACGCG']	5384	['CCTTCCCTCCAGGCTVCTTGTAC']	5626	['CCTTCCCTCCAGGCTVCTTGTAC']
NM_014191.3(SCN8A):c.5302A>G (p.Asn1768Asp)	6334	SCN8A	5187	['CATGTACATTGCCATCATCCTGGAGRAC TTCAGTGTAGCCACAGAGGAAAG']	5385	['CCATCATCCTGGAGRACCTTCAGT']	5627	['CCATCATCCTGGAGRACCTTCAGT']
NM_002552.4(ORC4):c.521A>G (p.Tyr174Cys)	5000	ORC4	5188	['CATCATAAAAACCAAACACTTCTCTRTAATCTTTTTGACATTCTCAGTCT']	5386	['CAAACACTTCTCTRTAATCTTT']	5628	['CAAACACTTCTCTRTAATCTTT']
NM_004813.2(PEX16):c.992A>G (p.Tyr331Cys)	9409	PEX16	5189	['TACTTGCCCACCTGGCAGAAAATCTRCTTCTACAGTTGGGGCTGACAGACC']	5387 - 5388	['CCACCTGGCAGAAAATCTRCTTC', 'CCTGGCAGAAAACTCTRCTTCTAC']	5629- 5630	['CCACCTGGCAGAAAATCTRCTTC', 'CCTGGCAGAAAACTCTRCTTCTAC']
NM_016952.4(CDON):c.2368A>G (p.Thr790Ala)	5093 7	CDON	5190	['GTTTTTGTTCCTCAAAAGGTTTCARCATACAAATTTAGGGTCATTGCCAT']	5389	['CCCTCAAAGGTTTCARCATACAA']	5631	['CCCTCAAAGGTTTCARCATACAA']
NM_016464.4(TMEM138):c.287A>G (p.His96Arg)	5152 4	TMEM138	5191	['TACTTTGCCCTCAGCATCTCCCTTCRTGTCTGGGTCATGGTAAGAGTGGCA']	5390 - 5391	['CCCTCAGCATCTCCCTTCRTGTC', 'CCTCAGCATCTCCCTTCRTGTCT']	5632- 5633	['CCCTCAGCATCTCCCTTCRTGTC', 'CCTCAGCATCTCCCTTCRTGTCT']
NM_005022.3(PFN1):c.350A>G (p.Glu117Gly)	5216	PFN1	5192	['GTTGATCAAACCA CCGTGGACACCTYCTTTGCCATCAGCAGGACTAGCGC']	5392	['CACCGTGGACACCTYCTTTGCC']	5634	['CACCGTGGACACCTYCTTTGCC']
NM_022787.3(NMNAT1):c.817A>G (p.Asn273Asp)	6480 2	NMNAT1	5193	['GGTCATCCTGGCCCTTTGCAGAGARACACTGCAGAAGCTAAGACATAGGA']	5393	['CCCCTTTGCAGAGARACACTGCA']	5635	['CCCCTTTGCAGAGARACACTGCA']
NM_005340.6(HINT1):c.152A>G (p.His51Arg)	3094	HINT1	5194	['GACATTTCCCTCAAGCACCAACACRTTTCTGGTGATACCAAGAAACAT']	5394 - 5396	['CCCCTCAAGCACCAACACRTTTT', 'CCCTCAAGCACCAACACRTTTTTC', 'CCTCAAGCACCAACACRTTTTCT']	5636- 5638	['CCCCTCAAGCACCAACACRTTTT', 'CCCTCAAGCACCAACACRTTTTTC', 'CCTCAAGCACCAACACRTTTTCT']
NM_005211.3(CSF1R)	1436	CSF1R	5195	['GACTAACCTGCA	5397	['CCTGCAGTGCTT	5639	['CCTGCAGTGCTT

FIR):c.2320-2A>G				GTGCTTTCCTCRGT GCATCCACCGGGAC GTGGCAGCG']		TCCCTCRGTGC']		TCCCTCRGTGC']
NM_001039958.1 (MESP2):c.271A >G (p.Lys91Glu)	1458 73	MESP2	5196	['GCGGCAGAGCGCC AGCGAGCGGGAGRA ACTGCGCATGCGCA CGCTGGCCCG']	5398	['CCAGCGAGCGGG AGRAACTGCGC']	5640	['CCAGCGAGCGGG AGRAACTGCGC']
NM_001099274.1 (TINF2):c.850A> G (p.Thr284Ala)	2627 7	TINF2	5197	['TAGGGGAGGCCAT AAGGAGCGCCCCRC AGTCATGCTGTTTCC CTTTAGGAA']	5399	['CCATAAGGAGCG CCCCRCAGTCA']	5641	['CCATAAGGAGCG CCCCRCAGTCA']
NM_003863.3(DP M2):c.68A>G (p.Tyr23Cys)	8818	DPM2	5198	['GCCGTTAGCCTGAT CATTTACCTRCTA CACCGCCTGGGTGA TTCTCTTG']	5400	['CCTGATCATCTT CACCTRCTACA']	5642	['CCTGATCATCTT CACCTRCTACA']
NM_000530.6(M PZ):c.347A>G (p.Asn116Ser)	4359	MPZ	5199	['AAGGATGGCTCCA TTGTCATACACARC CTAGACTACAGTGA CAATGGCACG']	5401	['CCATTGTCATAC ACARCCTAGAC']	5643	['CCATTGTCATAC ACARCCTAGAC']
NM_000138.4(FB N1):c.3058A>G (p.Thr1020Ala)	2200	FBN1	5200	['ACCCGGATTTGCC ACAAAAGAAATTRC AAATGGAAAGCCTT TCTTCAAAGG']	5402	['CCACAAAAGAA ATTRCAAATGGA']	5644	['CCACAAAAGAA ATTRCAAATGGA']
NM_000169.2(G LA):c.1153A>G (p.Thr385Ala)	-1	-	5201	['GGCCTGTAATCCTG CCTGCTTCATRCAC AGCTCCTCCCTGTG AAAAGGAA']	5403	['CCTGCCTGCTTC ATRCACAGCT']	5645	['CCTGCCTGCTTC ATRCACAGCT']
NM_000257.3(M YH7):c.2206A>G (p.Ile736Val)	4625	MYH7	5202	['AGCGGCCATCCCT GAGGGACAGTTCRT TGATAGCAGGAAGG GGGCAGAGAA']	5404	['CCCTGAGGGACA GTTTCRTTGATA']	5646- 5647	['CCCTGAGGGACA GTTTCRTTGATA', 'CCTGAGGGACAG TTCRTTGATAG']
NM_018972.2(G DAP1):c.368A>G (p.His123Arg)	5433 2	GDAP1	5203	['AGCATGTATTACCC ACGGGTACAACRTT ACCGAGAGCTGCTT GACTCCTTG']	5405	['CCCACGGGTACA ACRTTACCGAG']	5648	['CCCACGGGTACA ACRTTACCGAG']
NM_001946.3(D USP6):c.566A>G (p.Asn189Ser)	1848	DUSP6	5204	['ACTACCATCCGAG TCTGTTGCACTAYTG GGGTCTCGGTCAAG GTCAGACTC']	5406	['CCGAGTCTGTTG CACTAYTGGGG']	5649	['CCGAGTCTGTTG CACTAYTGGGG']
NM_003867.3(FG F17):c.560A>G (p.Asn187Ser)	8822	FGF17	5205	['TACCAAGGCCAGC TGCCCTTCCCCARCC ACGCCGAGAAGCAG AAGCAGTTC']	5407	['CCAGCTGCCCTT CCCCARCCACG']	5650	['CCAGCTGCCCTT CCCCARCCACG']
NM_015560.2(OP	4976	OPAI	5206	['TTTTTATTTTCT	5408	['CCTGAGTAGACC	5651	['CCTGAGTAGACC

A1):c.1146A>G (p.Ile382Met)				GAGTAGACCATRTC CTTAAATGTAAAAG GCCCTGGAC']		ATRTCCTTAAA']		ATRTCCTTAAA']
NM_002972.3(SB F1):c.1249A>G (p.Met417Val)	6305	SBF1	5207	['AAGGCCATGCCCT CCAGCACCTTTCAYC AGGAAATCGTCCTC TACCAGCCCA']	5409	['CCCTCCAGCACC TTCAYCAGGAA']	5652- 5653	['CCCTCCAGCACC TTCAYCAGGAA', 'CCTCCAGCACCTT CAYCAGGAAA']
NM_006876.2(B4 GAT1):c.1168A> G (p.Asn390Asp)	1104 1	B4GAT 1	5208	['GTTCCATCCCCAAA AGGAGGCTGAARAT CAGCACATAAGAT CCTATATCG']	5410	['CCAAAAGGAGG CTGAARATCAGC']	5654- 5656	['CCCCAAAAGGAGG GCTGAARATCA', 'CCAAAAGGAGG CTGAARATCAG', 'CCAAAAGGAGGC TGAARATCAGC']
NM_000218.2(K CNQ1):c.332A>G (p.Tyr111Cys)	3784	KCNQ 1	5209	['CGCACCCACGTCC AGGGCCGCGTCTRC AACTTCCTCGAGCG TCCCACCGGC']	5411	['CCAGGGCCGCGT CTRCAACTTCC']	5657	['CCAGGGCCGCGT CTRCAACTTCC']
NM_000492.3(CF TR):c.1A>G (p.Met1Val)	1080	CFTR	5210	['CAGGGACCCAGC GCCCCGAGAGACRT GCAGAGGTGCCTC TGAAAAGGC']	5412	['CCAGCGCCCGAG AGACCRTGCAG']	5658- 5659	['CCAGCGCCCGGA GAGACCRTGCAG', 'CCAGCGCCCGAG AGACCRTGCAG']
NM_007294.3(B RCA1):c.122A>G (p.His41Arg)	672	BRCA1	5211	['GAACCTGTCTCCAC AAAGTGTGACCRCA TATTTTGCAAGTAA GTTTGAATG']	5413	['CCACAAAGTGTG ACRCATATTT']	5660	['CCACAAAGTGTG ACRCATATTT']
NM_007294.3(B RCA1):c.4485- 2A>G	672	BRCA1	5212	['GTTTTCTCCTTCCA TTTATCTTTCTRGGT CATCCCCTTCTAAAT GCCCATC']	5414	['CCTTCCATTTATC TTTCTRGGTC']	5661- 5662	['CCTTCCATTTATC TTTCTRGGTC', 'CCATTTATCTTTC TRGGTCATCC']
NM_014795.3(ZE B2):c.3134A>G (p.His1045Arg)	9839	ZEB2	5213	['AAACACAAGCACC ACCTTATCGAGCRC TCAAGGCTTCACTC GGGCGAGAAG']	5415	['CCACCTTATCGA GCRCTCAAGGC']	5663	['CCACCTTATCGA GCRCTCAAGGC']
NM_001287.5(CL CN7):c.296A>G (p.Tyr99Cys)	1186	CLCN7	5214	['TGTCCCGCCTGCA GAGCTTGACTRTG ACAACAGTGAGAAC CAGCTGTTC']	5416	['CCTGCAGAGCTT GGACTRTGACA']	5664	['CCTGCAGAGCTT GGACTRTGACA']
NM_080605.3(B3 GALT6):c.1A>G (p.Met1Val)	1267 92	B3GAL T6	5215	['CGCCACGCCCGCC GCAGCAGCTTTCAYG GCGCCCCGCGCCGGG CCGGCGGCC']	5417	['CCCGCCGCAGCA GCTTTCAYGGCG']	5665- 5667	['CCCGCCGCAGCA GCTTTCAYGGCG', 'CCGCCGCAGCAG CTTTCAYGGCGC', 'CCGCAGCAGCTT CAYGGCGCCCC']
NM_000207.2(IN	-1	-	5216	['TCCTGCACCGAGA	5418	['CCGAGAGAGATG	5668	['CCGAGAGAGATG

S):c.*59A>G				GAGATGGAATAARG CCCTTGAACCAGCC CTGCTGTGCC']		GAATAARGCCC']		GAATAARGCCC']
NM_000784.3(CYP27A1):c.1061A>G (p.Asp354Gly)	1593	CYP27 A1	5217	['TGGGCCCTGTACC ACCTCTCAAAGGRC CCTGAGATCCAGGA GGCCTTGAC']	5419	['CCACCTCTCAAA GGRCCCTGAGA']	5669	['CCACCTCTCAAA GGRCCCTGAGA']
NM_000540.2(RYR1):c.14572A>G (p.Asn4858Asp)	6261	RYR1	5218	['CTACCTGTACACCG TGGTGGCCTTCRAC TTCTCCGCAAGTTC TACAACAA']	5420	['CCGTGGTGGCCT TTRACTTCTTC']	5670	['CCGTGGTGGCCT TTRACTTCTTC']
NM_000238.3(KCNH2):c.1478A>G (p.Tyr493Cys)	3757	KCNH 2	5219	['CACCCCGCCGCA TCGCCGTCCACTNC TTCAAGGGCTGGTT CCTCATCGAC']	5421	['CCGCATCGCCGT CCACTNCTTCA']	5671	['CCGCATCGCCGT CCACTNCTTCA']
NM_000335.4(SCN5A):c.688A>G (p.Ile230Val)	6331	SCN5A	5220	['CCGAGTCTCCGG GCCCTGAAAACRT ATCAGTCATTTAG GTGAAAATCA']	5422	['CCGGGCCCTGAA AACTRTATCAG']	5672	['CCGGGCCCTGAA AACTRTATCAG']
NM_000169.2(GLA):c.548-2A>G	-1	-	5221	['TATTTTACCCATTG TTTTCTCATACRGGT TATAAGCACATGTC CTTGCCCC']	5423	['CCCATTGTTTTCT CATACRGGTT']	5673- 5674	['CCCATTGTTTTCT CATACRGGTT', 'CCATTGTTTTCTC ATACRGGTTA']
NM_000146.3(FTL):c.1A>G (p.Met1Val)	2512	FTL	5222	['GTTAGCTCCTTCTT GCCAACCAACCRGT AGCTCCCAGATTCG TCAGAATTA']	5424	['CCTTCTTGCCAA CCAACCRGTAG']	5675	['CCTTCTTGCCAA CCAACCRGTAG']
NM_000531.5(OTC):c.1034A>G (p.Tyr345Cys)	5009	OTC	5223	['GTCATGGTGTCCCT GCTGACAGATTRCT CACCTCAGCTCCAG AAGCCTAAA']	5425 - 5426	['CCCTGCTGACAG ATTRCTCACCT', 'CCTGCTGACAGA TTRCTCACCTC']	5676- 5677	['CCCTGCTGACAG ATTRCTCACCT', 'CCTGCTGACAGA TTRCTCACCTC']
NM_000531.5(OTC):c.350A>G (p.His117Arg)	5009	OTC	5224	['TGTTTTCTTACCAC ACAAGATATTCDDT TGGGTGTGAATGAA AGTCTCACG']	5427	['CCACACAAGATA TTCDDTTGGGT']	5678	['CCACACAAGATA TTCDDTTGGGT']
NM_000531.5(OTC):c.524A>G (p.Asp175Gly)	5009	OTC	5225	['TACCATCCTATCCA GATCCTGGCTGDDT ACCTCACGCTCCAG GTTGGTTTA']	5428	['CCAGATCCTGGC TGDDTACCTCA']	5679	['CCAGATCCTGGC TGDDTACCTCA']
NM_000531.5(OTC):c.527A>G (p.Tyr176Cys)	5009	OTC	5226	['CATCCTATCCAGAT CCTGGCTGATTRCCT CACGCTCCAGGTTG GTTTATTT']	5429	['CCAGATCCTGGC TGATTRCCTCA']	5680	['CCAGATCCTGGC TGATTRCCTCA']
NM_000531.5(OTC):c.527A>G (p.Tyr176Cys)	5009	OTC	5227	['TCTCCTTCATCCCG	5430	['CCGTGCCTTTTA	5681-	['CCCGTGCCTTTT

TC):c.542A>G (p.Glu181Gly)				TGCCTTTTAGGRAC ACTATAGCTCTCTG AAAGGTCTT']		GGRACACTATA']	5682	AGGRACACTAT', 'CCGTGCCTTTTAG GRACACTATA']
NM_024301.4(FK RP):c.1A>G (p.Met1Val)	7914 7	FKRP	5228	['CCAGCTAGCCCCA GACTTCGGCCCCRT GCGGCTCACCCGCT GCCAGGCTGC']	5431	['CCCAGACTTCG GCCCCRTGCGG']	5683- 5685	['CCCAGACTTCG GCCCCRTGCGG', 'CCAGACTTCGG CCCCRTGCGGC', 'CCAGACTTCGGC CCCRGCGGCT']
NM_000321.2(R B1):c.1927A>G (p.Lys643Glu)	5925	RB1	5229	['AGCCTTCCAGACC CAGAAGCCATTGRA ATCTACCTCTCTTTC ACTGTTTTA']	5432	['CCAGAAGCCAT TGRAATCTACC']	5686	['CCAGAAGCCAT TGRAATCTACC']
NM_015713.4(R RM2B):c.581A> G (p.Glu194Gly)	5048 4	RRM2 B	5230	['AAAAGATCCTGAG AAGAAAACCTCTYC TACAGCAGCAAAGG CCACCACTCT']	5433	['CTGAGAAGAAA ACTCCTYCTAC']	5687	['CTGAGAAGAAA ACTCCTYCTAC']
NM_000219.5(K CNE1):c.242A>G (p.Tyr81Cys)	3753	KCNE1	5231	['CACTCGAACGACC CATTCAACGTCTDC ATCGAGTCCGATGC CTGGCAAGAG']	5434	['CCATTCAACGT CTDCATCGAGT']	5688	['CCATTCAACGT CTDCATCGAGT']
NM_003108.3(SO X11):c.347A>G (p.Tyr116Cys)	6664	SOX11	5232	['AAGCACATGGCCG ACTACCCCGACTRC AAGTACCGGCCCG GAAAAAGCCC']	5435	['CCGACTACCCCG ACTRCAAGTAC']	5689	['CCGACTACCCCG ACTRCAAGTAC']
NM_002764.3(PR PS1):c.343A>G (p.Met115Val)	5631	PRPS1	5233	['AATCTCAGCCAAG CTTGTTGCAAARTG CTATCTGTAGCAGG TGCAGATCA']	5436	['CCAAGCTTGTTG CAAARTGCTA']	5690	['CCAAGCTTGTTG CAAARTGCTA']
NM_000546.5(TP 53):c.1101-2A>G	7157	TP53	5234	['TCTCCTCCCTGCTT CTGTCTCTACRGCC ACCTGAAGTCCAAA AAGGGTCA']	5437	['CCTGCTTCTGTCT CCTACRGCCA']	5691	['CCTGCTTCTGTCT CCTACRGCCA']
NM_000166.5(GJ B1):c.580A>G (p.Met194Val)	2705	GJB1	5235	['CGAGAAAACCGTC TTCACCGTCTTCRTG CTAGCTGCCTCTGG CATCTGCAT']	5438	['CCGTCTTCACCG TCTTCRTGCTA']	5692	['CCGTCTTCACCG TCTTCRTGCTA']
NM_003159.2(C DKL5):c.449A>G (p.Lys150Arg)	6792	CDKL5	5236	['TTAATCAGCCACA ATGATGTCCTAARA CTGTGTGACTTTGGT AAGTTAAAA']	5439	['CCACAATGATGT CCTAARACTGT']	5693	['CCACAATGATGT CCTAARACTGT']
NM_000053.3(A TP7B):c.122A>G (p.Asn41Ser)	540	ATP7B	5237	['ATCCAGACCACCTT CATAGCCAACAYTG TCAAAAAGCAAACCT	5440	['CCACCTTCATAG CCAACAYTGTC']	5694- 5695	['CCACCTTCATAG CCAACAYTGTC', 'CCTTCATAGCCAA

				CTTCTTCAT']				CAYTGTCAAA']
NM_006306.3(SMC1A):c.3254A>G (p.Tyr1085Cys)	8243	SMC1 A	5238	['GTGGCTACCAACA TTGATGAGATCTRT AAGGCCCTGTCCCG CAATAGCAGT']	5441	['CCAACATTGATG AGATCTRTAAG']	5696	['CCAACATTGATG AGATCTRTAAG']
NM_005154.4(USP8):c.2150A>G (p.Tyr717Cys)	9101	USP8	5239	['GAACCTTCCAAAC TGAAGCGTCTCCTDC TCCTCCCAGATAT AACCCAGGCT']	5442	['CCAAACTGAAGC GCTCCTDCTCC']	5697	['CCAAACTGAAGC GCTCCTDCTCC']
NM_000117.2(EMDMD):c.266-2A>G	2010	EMD	5240	['TCTGCTACCGCTGC CCCCCTTCCCARGG CTACAATGACGACT ACTATGAAG']	5443	['CCGCTGCCCCCC TCCCARGGCT']	5698	['CCGCTGCCCCCC TCCCARGGCT']
NM_207352.3(CYP4V2):c.1393A>G (p.Arg465Gly)	2854 40	CYP4V 2	5241	['CTACGTGCCCTTCT CTGCTGGCCCCRGG AACTGTATAGGTTT GTATCCATC']	5444	['CCCTTCTCTGCT GGCCCCRGGAA']	5699- 5700	['CCCTTCTCTGCT GGCCCCRGGAA', 'CCTTCTCTGCTGG CCCCRGGAA']
NM_000546.5(TP53):c.709A>G (p.Met237Val)	7157	TP53	5242	['CTGTACCACCATCC ACTACAACACTACRTG TGTAACAGTTCCTG CATGGGCGG']	5445	['CCATCCACTACA ACTACRTGTGT']	5701	['CCATCCACTACA ACTACRTGTGT']
NM_016069.9(PAM16):c.226A>G (p.Asn76Asp)	-1	-	5243	['CTCACCCGTCCCCT CTCCTCTGCAGRAC TATGAACACTTATTT AAGGTGAA']	5446	['CCTCTCCTCTGC AGRACTATGAA']	5702- 5704	['CCCCTCTCCTCT GCAGRACACTATG', 'CCCTCTCCTCTGC AGRACTATGA', 'CCTCTCCTCTGCA GRACACTATGAA']
NM_006785.3(MALT1):c.1019-2A>G	1089 2	MALT 1	5244	['AACACCCCTTTTCT TTTTTTTTCAARGCG AAGGACAAGGTTGC CCTTTTGA']	5447	['CCTTTCTTTTTTT TTCAARGCGA']	5705	['CCTTTCTTTTTTT TTCAARGCGA']
NM_004771.3(MMP20):c.611A>G (p.His204Arg)	9313	MMP2 0	5245	['GGAGAAGGCCTGG GAGGAGATACACRT TTCGACAATGCTGA GAAGTGGACT']	5448	['CCTGGGAGGAGA TACACRTTTCG']	5706	['CCTGGGAGGAGA TACACRTTTCG']
NM_003159.2(CDKL5):c.458A>G (p.Asp153Gly)	6792	CDKL5	5246	['CACAAATGATGTCT AAAACACTGTGTGRCT TTGGTAAGTTAAAA AGAAATTAA']	5449	['CCTAAAACACTGTG TGRCTTTGGTA']	5707	['CCTAAAACACTGTG TGRCTTTGGTA']
NM_001204830.1(LIPT1):c.535A>G (p.Thr179Ala)	-1	-	5247	['CCGGACTACTGCCT ATCACCATTGCRCTT TATTATGTAGTACTG ATGGGAC']	5450	['CCTATCACCATT GCRCTTTATTA']	5708	['CCTATCACCATT GCRCTTTATTA']
NM_000921.4(PDE3A)	5139	PDE3A	5248	['AGTTTCTTCCACTT	5451	['CCACTTGGACCA	5709	['CCACTTGGACCA

E3A):c.1333A>G (p.Thr445Ala)				GGACCACCACCRCC TCGGCCACAGGTCT ACCCACCTT']		CCACCRCCTCG']		CCACCRCCTCG']
NM_000182.4(H ADHA):c.919- 2A>G	3030	HADH A	5249	['TTGCTCAATTCCAG TCTTTACCACCYAA AAAACATATAAAGC ACTTGCTCA']	5452	['CCAGTCTTTACC ACCYAAAAAAC']	5710	['CCAGTCTTTACC ACCYAAAAAAC']
NM_000169.2(G LA):c.620A>G (p.Tyr207Cys)	-1	-	5250	['GTGTA CTCTGTGA GTGGCCTCTTTRTAT GTGGCCCTTTCAAA AGGTGAGA']	5453	['CCTGTGAGTGGC CTCTTTRTATG']	5711	['CCTGTGAGTGGC CTCTTTRTATG']
NM_000238.3(K CNH2):c.2582A> G (p.Asn861Ser)	3757	KCNH 2	5251	['TGGTCCAGCCTGG AGATCACCTTCANC CTGCGAGATGTGAG TTGGCTGCCC']	5454	['CCTGGAGATCAC CTTCANCCTGC']	5712	['CCTGGAGATCAC CTTCANCCTGC']
NM_000218.2(K CNQ1):c.605A>G (p.Asp202Gly)	3784	KCNQ 1	5252	['GCTCCCCCTCTCCT GCACTCCACAGRCC TCATCGTGGTCGTG GCCTCCATG']	5455	['CCTGCACTCCAC AGRCCTCATCG']	5713	['CCTGCACTCCAC AGRCCTCATCG']
NM_012203.1(G RHPR):c.934A>G (p.Asn312Asp)	9380	GRHP R	5253	['CACCATGTCCTTGT TGGCAGCTAACRAC TTGCTGGCTGGCCT GAGAGGGGA']	5456	['CCTTGTGGCAG CTAACRACTTG']	5714	['CCTTGTGGCAG CTAACRACTTG']
NM_021007.2(SC N2A):c.387- 2A>G	6326	SCN2A	5254	['ACTTTGTCTTCCTT GACGATATTCTRCTT TATTCAATATGCTCA TTATGTG']	5457	['CCTTGACGATAT TCTRCTTTATT']	5715	['CCTTGACGATAT TCTRCTTTATT']
NM_002693.2(PO LG):c.2840A>G (p.Lys947Arg)	5428	POLG	5255	['GTGGGCATCAGCC GTGAGCATGCCARA ATCTTCAACTACGG CCGCATCTAT']	5458	['CCGTGAGCATGC CARAATCTTCA']	5716	['CCGTGAGCATGC CARAATCTTCA']
NM_020533.2(M COLN1):c.1406A >G (p.Asn469Ser)	5719	MCOL N1	5256	['TCTGAGTGCCTGTT CTCGCTCATCARTG GGGACGACATGTTT GTGACGTT']	5459	['CCTGTTCTCGCT CATCARTGGGG']	5717	['CCTGTTCTCGCT CATCARTGGGG']
NM_000069.2(C ACNA1S):c.3526 -2A>G	779	CACN AIS	5257	['TCGCTTTCCCATCC TTTTCTTCCCRGGG CTACTTTGGAGACC CCTGGAAT']	5460	['CCCATCCTTTTCC TCCCRGGGC']	5718- 5719	['CCCATCCTTTTCC TCCCRGGGC', 'CCATCCTTTTCTT TCCCRGGGCT']
NM_017662.4(TR PM6):c.3173A>G (p.Tyr1058Cys)	1408 03	TRPM6	5258	['CAAGCTGTCTACCT CTTCGTGCAATRTAT CATCATGGTGAACC TGTTGATT']	5461	['CCTCTTCGTGCA ATRTATCATCA']	5720	['CCTCTTCGTGCA ATRTATCATCA']
NM_006642.3(SD	1080	SDCC	5259	['AATAAACCCCTCTG	5462	['CCTCTGCTTTTGC	5721	['CCTCTGCTTTTGC

CCAG8):c.221-2A>G	6	AG8		CTTTTGCTCTATRGT TAATCAGCTCAAAG ATTTGTTGC']		TCTATRGTTA']		TCTATRGTTA']
NM_003560.2(PLA2G6):c.1349-2A>G	8398	PLA2G6	5260	['CAGCATGCCCTGCTCTGTGCCTCACRGA ACTACAGGATCTCATGCACATCT']	5463	['CCCTGCTCTGTGCCTCACRGAAC']	5722-5723	['CCCTGCTCTGTGCCTCACRGAAC', 'CCTGCTCTGTGCC TCACRGAAC']

EXAMPLE 6: Next Generation C to T Editors

[00351] Other families of cytidine deaminases as alternatives to base editor 3 (BE3) constructs were examined. The different C to T editors were developed to have a narrow or different editing window, alternate sequence specificity to expand targetable substrates, and to have higher activity.

[00352] Using the methods described in Example 4, the pmCDA1 (cytidine deaminase 1 from *Petromyza marinus*) activity at the HeK-3 site is evaluated (Figure 42). The pmCDA1-nCas9-UGI-NLS (nCas9 indicates the Cas9 nickase described herein) construct is active on some sites (e.g., the C bases on the complementary strand at position 9, 5, 4, and 3) that are not accessible with rAPOBEC1 (BE3).

[00353] The pmCDA1 activity at the HeK-2 site is given in Figure 43. The pmCDA1-XTEN-nCas9-UGI-NLS construct is active on sites adjacent to “G,” while rAPOBEC1 analog (BE3 construct) has low activity on “C”s that are adjacent to “G”s, e.g., the C base at position 11 on the complementary strand.

[00354] The percent of total sequencing reads with target C converted to T (Figure 44), C converted to A (Figure 45), and C converted to G (Figure 46) are shown for CDA and APOBEC1 (the BE3 construct).

[00355] The huAPOBEC3G activity at the HeK-2 site is shown in Figure 47. Two constructs were used: huAPOBEC3G-XTEN-nCas9-UGI-NLS and huAPOBEC3G*(D316R_D317R)-XTEN-nCas9-UGI-NLS. The huAPOBEC3G-XTEN-nCas9-UGI-NLS construct has different sequence specificity than rAPOBEC1 (BE3), as shown in Figure 47, the editing window appears narrow, as indicated by APOBEC3G’s decreased activity at position 4 compared to APOBEC1. Mutations made in huAPOBEC3G (D316R and D317R) increased ssDNA binding and resulted in an observable effect on expanding the sites which were edited (compare APOBEC3G with APOBEC3G_RR in Figure 47). Mutations were chosen based on APOBEC3G crystal structure, *see*: Holden *et al.*, Crystal structure of the anti-viral APOBEC3G catalytic domain and functional

implication. *Nature*. (2008); 121-4, the entire contents of which are incorporated herein by reference.

EXAMPLE 7: pmCDA1/huAPOBEC3G/rAPOBEC1 work in E. coli

[00356] LacZ selection optimization for the A to I conversion was performed using a bacterial strain with lacZ encoded on the F plasmid. A critical glutamic acid residue was mutated (e.g., GAG to GGG, Glu to Gly mutation) so that G to A by a cytidine deaminase would restore lacZ activity (Figure 48). Strain CC102 was selected for the selection assay. APOBEC1 and CDA constructs were used in a selection assay to optimize G to A conversion.

[00357] To evaluate the the effect of copy number of the plasmids encoding the deaminase constructs on lacZ reversion frequency, the CDA and APOBEC1 deaminases were cloned into 4 plasmids with different replication origins (hence different copy numbers), SC101, CloDF3, RSF1030, and PUC (copy number: PUC>RSF1030>CloDF3>SC101) and placed under an inducible promoter. The plasmids were individually transformed into E. coli cells harboring F plasmid containing the mutated LacZ gene. The expression of the deaminases were induced and LacZ activity was detected for each construct (Figure 49). As shown in Figure 49, CDA exhibited significantly higher activity than APOBEC1 in all instances, regardless of the plasmid copy number the deaminases were cloned in. Further, In terms of the copy number, the deaminase activity was positively correlated with the copy number of the plasmid they are cloned in, i.e., PUC> CloDF3>SC101.

[00358] LacZ reversions were confirmed by sequencing of the genomic DNA at the lacZ locus. To obtain the genomic DNA containing the corrected LacZ gene, cells were grown media containg X-gal, where cells having LacZ activity form blue colonies. Blue colonies were selected and grown in minimal media containing lactose. The cells were spun down, washed, and re-plated on minimal media plates (lactose). The blue colony at the highest dilution was then selected, and its genomic DNA was sequenced at the lacZ locus (Figure 50).

[00359] A chloramphenicol reversion assay was designed to test the activity of different cytidine deaminases (e.g., CDA, and APOBEC1). A plasmid harboring a mutant CAT1 gene which confers chloramphenicol resistance to bacteria is constructed with RSF1030 as the replication origin. The mutant CAT1 gene encodings a CAT1 protein that has a H195R (CAC to CGC) mutation, rendering the protein inactive (Figure 51). Deamination of the C base-paired to the G base in the CGC codon would convert the codon back to a CAC codon,

restoring the activity of the protein. As shown in Figure 52, CDA outperforms rAPOBEC in *E. coli* in restoring the activity of the chloramphenicol resistance gene. The minimum inhibitory concentration (MIC) of chlor in S1030 with the selection plasmid (pNMG_ch_5) was approximately 1 µg/mL. Both rAPOBEC-XTEN-dCas9-UGI and CDA-XTEN-dCas9-UGI induced DNA correction on the selection plasmid (Figure 53).

[00360] Next, the huAPOBEC3G-XTEN-dCas9-UGI protein was tested in the same assay. Interestingly, huAPOBEC3G-XTEN-dCas9-UGI exhibited different sequence specificity than the rAPOBEC1-XTEN-dCas9-UGI fusion protein. Only position 8 was edited with APOBEC3G-XTEN-dCas9-UGI fusion, as compared to the rAPOBEC1-XTEN-dCas9-UGI fusion (in which positions 3, 6, and 8 were edited) (Figure 54).

EXAMPLE 8: C to T Base Editors with Less Off Target Editing

[00361] Current base editing technologies allow for the sequence-specific conversion of a C:G base pair into a T:A base pair in genomic DNA. This is done via the direct catalytic conversion of cytosine to uracil by a cytidine deaminase enzyme and thus, unlike traditional genome editing technologies, does not introduce double-stranded DNA breaks (DSBs) into the DNA as a first step. See, Komor, A.C., Kim, Y.B., Packer, M.S., Zuris, J.A., and Liu, D.R. (2016), “Programmable editing of a target base in genomic DNA without double-stranded DNA cleavage.” *Nature* 533, 420-424; the entire contents of which are incorporated by reference herein. Instead, catalytically dead SpCas9 (dCas9) or a SpCas9 nickase (dCas9(A840H)) is tethered to a cytidine deaminase enzyme such as rAPOBEC1, pmCDA1, or hAPOBEC3G. The genomic locus of interest is encoded by an sgRNA, and DNA binding and local denaturation is facilitated by the dCas9 portion of the fusion. However, just as wt dCas9 and wt Cas9 exhibit off-target DNA binding and cleavage, current base editors also exhibit C to T editing at Cas9 off-target loci, which limits their therapeutic usefulness.

[00362] It has been reported that the introduction of just three to four mutations into SpCas9 that neutralize nonspecific electrostatic interactions between the protein and the sugar-phosphate backbone of its target DNA, increases the DNA binding specificity of SpCas9. See, Kleinstiver, B.P., Pattanayak, V., Prew, M.S., Tsai, S.Q., Nguyen, N.T., Zheng, Z., and Joung, J.K. (2016) “High-fidelity CRISPR–Cas9 nucleases with no detectable genome-wide off-target effects.” *Nature* 529, 490-495; and Slaymaker, I.M., Gao, L., Zetsche, B., Scott, D.A., Yan, W.X., and Zhang, F. (2015) “Rationally engineered Cas9 nucleases with improved specificity. *Science* 351, 84-88; the entire contents of each are hereby incorporated by reference herein. Four reported neutralizing mutations were therefore

incorporated into the initially reported base editor BE3 (SEQ ID NO: 285), and found that off-target C to T editing of this enzyme is also drastically reduced (Figure 55), with no decrease in on-target editing (Figure 56).

[00363] As shown in Figure 55, HEK293T cells were transfected with plasmids expressing BE3 or HF-BE3 and a sgRNA matching the EMX1 sequence using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target locus, plus the top ten known Cas9 off-target loci for the EMX1 sgRNA, as previously determined by Joung and coworkers using the GUIDE-seq method. See Tsai, S.Q., Zheng, Z., Nguyen, N.T., Liebers, M., Topkar, V.V., Thapar, V., Wyvekens, N., Khayter, C., Iafrate, A.J., Le, L.P., *et al.* (2015) "GUIDE-seq enables genome-wide profiling of off-target cleavage by CRISPR-Cas nucleases." *Nat Biotech* 33, 187-197; the entire contents of which are incorporated by reference herein. EMX1 off-target 5 locus did not amplify and is not shown. Sequences of the on-target and off-target protospacers and protospacer adjacent motifs (PAMs) are displayed (Figure 55). Cellular C to T conversion percentages, defined as the percentage of total DNA sequencing reads with T at each position of an original C within the protospacer, are shown for BE3 and HF-BE3.

[00364] In Figure 56, HEK293T cells were transfected with plasmids expressing BE3 or HF-BE3 and sgRNAs matching the genomic loci indicated using Lipofectamine 2000. Three days after transfection, genomic DNA was extracted, amplified by PCR, and analyzed by high-throughput DNA sequencing at the on-target loci. The percentage of total DNA sequencing reads with all four bases at the target Cs within each protospacer are shown for treatment with BE3 or HF-BE3 (Figure 56). Frequencies of indel formation are shown as well.

[00365] Primary Protein Sequence of HF-BE3 (SEQ ID NO: 285):

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELRKETCLLYEINWGGRHSIWRHTSQNTNKHVEVNFIEKF
TTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYHHADPRNRQGLRDLISSGVTI
QIMTEQESGYCWRNFVNYSNEAHWPYRPHLWVRLYVLELYCILGLPPCLNLRKQPQLTFFTIALQ
SCHYQRLPPHILWATGLKSGSETPGTSESATPESDKKYSIGLAIGTNSVGWAVITDEYKVPKFKVLG
NTDRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRLEESFL
VEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDLN
PDNSDVDKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSKSRRLLENLIAQLPGEKKNGLFGNLIAS
LGLTPNFKSNFDLAEDAQLSKDQYDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITK
APLSASMIKRYDEHHQDLTLLKALVRQQLPKEYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKM
DGTEELLVKLNREDLLRQRTFDNGSIPHQIHLGELHAILRRQEDFYFPFLKDNREKIEKILTFRIPYYVGP
LARGNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTAFDKNLPNEKVLPHKSLLYEYFTVYN
ELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASL
GTYHDLKIKDKDFLDNEENEDILEDIVLTLTLFEDREMIEERLKYAHLFDDKVMKQLKRRRYTGWG
ALSRKLINGIRDKQSGKTILDFLKSDFANRNFMALIHDDSLTFKEDIQKAQVSGQGDLSHEHIANLAGS
PAIKKGILQTVKVVDELVKVMGRHKPENIVIAMARENQTTQKGQKNSRERMKRIIEGKELGSQILKEH

PVENTQLQNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDKNRGKS
 DNVPSSEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDKAGFIKRQLVETRAITKHVAQIL
 DSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYK VREINNYHHAHDAYLNAVVG TALIKKYPK
 LESEFVYGDYK VYDVRKMIKSEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIV
 WDKGRDFATVRK VLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVA
 YSVLVVAKVEKGGKSKLKS VKELLGITIMERSSEKPNIDFLEAKGYKEVKKDLI IKLPKYSLFELENGR
 KRMLASAGELQKGNELALPSKYVNFLYLASHYEKLGSPEDNEQKQLFVEQHKHYLDEIIEQISEFSKR
 VILADANLDKVL SAYNKHDKPIREQAENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIH
 QSITGLYETRIDLSQLGGDSGGSTNLSDIIEKETGKQLVIQESILMLPEEVVEEVIGNKPEDILVHTAYDES
 TDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSPKKRKY

EXAMPLE 9: Development of Base Editors that Use Cas9 Variants and Modulation of the Base Editor Processivity to Increase the Target Range and Precision of the Base Editing Technology

[00366] Unlike traditional genome editing platforms, base editing technology allows precise single nucleotide changes in the DNA without inducing double-stranded breaks (DSBs). See, Komor, A. C. *et al. Nature* **533**, 420-424 (2016). The current generation of base editor uses the NGG PAM exclusively. This limits its ability to edit desired bases within the genome, as the base editor needs to be placed at a precise location where the target base is placed within a 4-base region (the ‘deamination window’), approximately 15 bases upstream of the PAM. See, Komor, A. C. *et al. Nature* **533**, 420-424 (2016). Moreover, due to the high processivity of cytidine deaminase, the base editor may convert all cytidines within its deamination window into thymidines, which could induce amino acid changes other than the one desired by the researcher. See, Komor, A. C. *et al. Nature* **533**, 420-424 (2016).

Expanding the scope of base editing through the development of base editors with Cas9 variants

[00367] Cas9 homologs and other RNA-guided DNA binders that have different PAM specificities were incorporated into the base editor architecture. See, Kleinstiver, B. P. *et al. Nature* **523**, 481–485 (2015); Kleinstiver, B. P. *et al. Nature Biotechnology* **33**, 1293–1298 (2015); and Zetsche, B. *et al. Cell* **163**, 759-771 (2015); the entire contents of each are incorporated by reference herein. Furthermore, innovations that have broadened the PAM specificities of various Cas9 proteins were also incorporated to expand the target reach of the base editor even more. See, Kleinstiver, B. P. *et al. Nature* **523**, 481–485 (2015); and Kleinstiver, B. P. *et al. Nature Biotechnology* **33**, 1293–1298 (2015). The current palette of base editors is summarized in Table 4.

Table 4. New base editors made from Cas9 Variants

Species	PAM	Base Editor Name	Reference for Cas9 variant
<i>S. pyogenes</i>	...NGG	BE3	Wild-type
	...NGA	VQR BE3 or EQR BE3	Kleinstiver, B. P. <i>et al.</i>
	...NGCG	VRER BE3	Kleinstiver, B. P. <i>et al.</i>
<i>S. aureus</i>	...NNGRRT	SaBE3	Wild-type
	...NNNRRT	SaKKH BE3	Kleinstiver, B. P. <i>et al.</i>
<i>L. bacterium</i>	TTN...	dCpf1 BE2	Zetsche, B. <i>et al.</i>

Modulating base editor's processivity through site-directed mutagenesis of rAPOBEC1

[00368] It was reasoned that the processivity of the base editor could be modulated by making point mutations in the deaminase enzyme. The incorporation of mutations that slightly reduce the catalytic activity of deaminase in which the base editor could still catalyze on average one round of cytidine deamination but was unlikely to access and catalyze another deamination within the relevant timescale were pursued. In effect, the resulting base editor would have a narrower deamination window.

[00369] rAPOBEC1 mutations probed in this work are listed in Table 5. Some of the mutations resulted in slight apparent impairment of rAPOBEC1 catalysis, which manifested as preferential editing of one cytidine over another when multiple cytidines are found within the deamination window. Combining some of these mutations had an additive effect, allowing the base editor to discriminate substrate cytidines with higher stringency. Some of the double mutants and the triple mutant allowed selective editing of one cytidine among multiple cytidines that are right next to one another (Figure 57).

Table 5. rAPOBEC1 Point Mutations Investigated

rAPOBEC1 mutation studied in this work	Corresponding mutation in APOBEC3G	Reference
H121R / H122R	D315R / D316R	Holden, L. G. <i>et al.</i>
R126A	R320A	Chen, K-M. <i>et al.</i>
R126E	R320E	Chen, K-M. <i>et al.</i>
R118A	R313A	Chen, K-M. <i>et al.</i>
W90A	W285A	Chen, K-M. <i>et al.</i>
W90Y	W285Y	
R132E	R326E	

Base Editor PAM Expansion and Processivity Modulation

[00370] The next generation of base editors were designed to expand editable cytidines in the genome by using other RNA-guided DNA binders (Figure 58). Using a NGG PAM only

allows for a single target within the “window” whereas the use of multiple different PAMs allows for Cas9 to be positioned anywhere to effect selective deamination. A variety of new base editors have been created from Cas9 variants (Figure 59 and Table 4). Different PAM sites (NGA, Figure 60; NGCG, Figure 61; NNGRRT, Figure 62; and NNHRRT, Figure 63) were explored. Selective deamination was successfully achieved through kinetic modulation of cytidine deaminase point mutagenesis (Figure 65 and Table 5).

[00371] The effect of various mutations on the deamination window was then investigated in cell culture using spacers with multiple cytidines (Figures 66 and 67).

[00372] Further, the effect of various mutations on different genomic sites with limited numbers of cytidines was examined (Figures 68 to 71). It was found that approximately one cytidine will be edited within the deamination window in the spacer, while the rest of the cytidines will be left intact. Overall, the preference for editing is as follows: $C_6 > C_5 \gg C_7 \approx C_4$.

Base Editing Using Cpf1

[00373] Cpf1, a Cas9 homolog, can be obtained as AsCpf1, LbCpf1, or from any other species. Schematics of fusion constructs, including BE2 and BE3 equivalents, are shown in Figure 73. The BE2 equivalent uses catalytically inactive Cpf2 enzyme (dCpf1) instead of Cas9, while the BE3 equivalent includes the Cpf1 mutant, which nicks the target strand. The bottom schematic depicts different fusion architectures to combine the two innovations illustrated above it (Figure 73). The base editing results of HEK293T cell TTTN PAM sites using Cpf1 BE2 were examined with different spacers (Figures 64A to 64C). In some embodiments, Cpf1 may be used in place of a Cas9 domain in any of the base editors provided herein. In some embodiments, the Cpf1 is a protein that is at least 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 98%, 99%, or 99.5% identical to SEQ ID NO 313.

Full Protein Sequence of Cpf1 (SEQ ID NO: 313):

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MSIYQEFVNKYSLSKTLRFELIPQGKTLENIKARGLILDDEKRAKDYKKAKQIIDKYHQFFIEEILSSVCIS
EDLLQNYSDVYFKLKKSDDDNLQKDFKSAKDTIKKQISEYIKDSEKFNLFNQNLIDAKKGQESDLILW
LKQSKDNGIELFKANSDITDIDEALEIISFKGWTTYFKGFHENRKNVYSSNDIPTSIYRIVDDNLPKFLE
NKAKYESLKDKAPEAINYEQIKKDLAEELTFDIDYKTSEVNQRVFSLDEVFEIANFNYYLNQSGITKFNT
IIGGKVFNGENTKRKGINEYINLYSQQINDKTLKKYKMSVLFKQILSDTESKSFVIDKLEDDSDVVTMQL
SFYEQIAAFKTVEEKSIKETLSLLFDDLKAQKLDLSKIYFKNDKSLTDLSQQVFDDYSVIGTAVLEYITQ
QIAPKNLDNPSKKEQELIAKKTEKAKYLSLETIKLAL EEFNKHRDIDKQCRFEEILANFAAIPMIFDEIAQ
NKDNLAQISIKYQNQGKDLLQASAEDDVKAIKDLLDQTNLLHKLKIFHISQSEDKANILDKDEHFYL
VFEECYFELANIVPLYNKIRNYITQKPYSDKFKLNFENSTLANGWDKNKEPDNTAILFIKDDKYLLGV
MNMKNNKIFDDKAIKENKGEYKKIVYKLLPGANKMLPKVFFSAKSIKFYNPSEDILRIRNHSTHTKNG
SPQKGYEKFEFNIEDCRKFIDFYKQSISKHPEWKDFGRFSDTQRYNSIDEFYREVENQYKLTFFENISES
YIDSVVNQGKLYLFQIYNKDFSAYSKGRPNLHTLYWKALFDERNLQDVVYKLNGEAELFYRKQSIPKK
ITHPAKEAIANKNDNPKKESVFEYDLIKDKRFTEDKFFFHCPITINFKSSGANKFNDEINLLLKEKAND
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VHILSIDRGERHLAYYTLVDGKGNIKQDTFNIIGNDRMKTNYHDKLAAIEKDRDSARKDWKKNINKE
 MKEGYLSQVVHEIAKLVEIYNAIVVFEDLNFGFKRGRFKVEKQVYQKLEKMLIEKLNLYL VFKDNEFDK
 TGGVLRAYQLTAPFETFKKMGKQTGIIYYVPAGFTSKICPVTGFVNQLYPKYESVSKSQEFFSKFDKICY
 NLDKGYFEFSFDYKNFGDKAAK GKWTIASFGSRLINFRNSDKNHNWDTREVYPTKELEKLLKDYSIEY
 GHGECIKAAICGESDKKFFAKLTSVLNTILQMRNSKTGTELDYLISPVADVNGNFFDSRQAPKNMPQDA
 DANGAYHIGLKGMLLGRKNNQEGKLNLVIKNEEYFEFVQNRNN

EXAMPLE 10: Increased Fidelity of Base Editing

[00374] Examining the difference between plasmid delivery of BE3 and HF-BE3, it was found that the two edit on-target loci with comparable efficiency (Figures 74 and 75). However, HF-BE3 edited off-target loci much less than BE3, meaning that HF-BE3 has a much higher DNA specificity than BE3 (Figure 76). Deaminase protein lipofection to HEK cells demonstrated that protein delivery of BE3 results in comparable on-target activity, but much better specificity, than plasmid DNA delivery of BE3. Using improved transfection procedures and better plasmids (n=2), the experiment used the following conditions: protein delivery was 125 nM Cas9:sgRNA complex, plasmid delivery was 750ng BE3/HF-BE3 plasmid + 250ng sgRNA plasmid, and lipofection was with 1.5µL of Lipofectamine 2000 per well. EMX-1 off target site 2 and FANCF off-target site 1 showed the most off-target editing with BE3, compared to all of the off-targets assayed (Figures 77 and 78), while HEK-3 showed no significant editing at off-targets for any of the delivery methods (Figure 79). HEK-4 shows some C-to-G editing on at the on-target site, while its off-target sites 1, 3, and 4 showed the most off-target editing of all the assayed sites (Figure 80).

Delivery of BE3 Protein via Micro-injection to Zebrafish

[00375] TYR guide RNAs were tested in an *in vitro* assay for sgRNA activity (Figures 81 and 82). The % HTS reads shows how many C residues were converted to T residues during a 2h incubation with purified BE3 protein and PCR of the resulting product. Experiments used an 80-mer synthetic DNA substrate with the target deamination site in 60bp of its genomic context. This is not the same as % edited DNA strands because only one strand was nicked, so the product is not amplified by PCR. The proportion of HTS reads edited is equal to $x/(2-x)$, where x is the actual proportion of THS reads edited. For 60% editing, the actual proportion of bases edited is 75%. “Off target” is represents BE3 incubated with the same DNA substrate, while bound to an off-target sgRNA. It was found sgRNAs sgRH_13, sgHR_17, and possibly sgHR_16 appeared to be promising targets for *in vivo* injection experiments.

[00376] The delivery of BE3 protein in was tested *in vivo* in zebrafish. Zebrafish embryos (n=16-24) were injected with either scrambled sgRNA, sgHR_13, sgHR_16, or sgHR_17 and purified BE3. Three embryos from each condition were analyzed independently (single embryo) and for each condition, all of the injected embryos were pooled and sequenced as a pool. The results are shown in Figures 83 to 85.

EXAMPLE 11: Uses of Base Editors to Treat Disease

[00377] Base editors or complexes provided herein (*e.g.*, BE3) may be used to modify nucleic acids. For example, base editors may be used to change a cytosine to a thymine in a nucleic acid (*e.g.*, DNA). Such changes may be made to, *inter alia*, alter the amino acid sequence of a protein, to destroy or create a start codon, to create a stop codon, to disrupt splicing donors, to disrupt splicing acceptors or edit regulatory sequences. Examples of possible nucleotide changes are shown in Figure 86.

[00378] Base editors or complexes provided herein (*e.g.*, BE3) may be used to edit an isoform of Apolipoprotein E in a subject. For example, an Apolipoprotein E isoform may be edited to yield an isoform associated with a lower risk of developing Alzheimer's disease. Apolipoprotein E has four isoforms that differ at amino acids 112 and 158. APOE4 is the largest and most common genetic risk factor for late-onset Alzheimer's disease. Arginine residue 158 of APOE4, encoded by the nucleic acid sequence CGC, may be changed to a cysteine by using a base editor (*e.g.*, BE3) to change the CGC nucleic acid sequence to TGC, which encodes cysteine at residue 158. This change yields an APOE3r isoform, which is associated with lower Alzheimer's disease risk. See Figure 87.

[00379] It was tested whether base editor BE3 could be used to edit APOE4 to APOE3r in mouse astrocytes (Figure 88). APOE 4 mouse astrocytes were nucleofected with Cas9 + template or BE3, targeting the nucleic acid encoding Arginine 158 of APOE4. The Cas9 + template yielded only 0.3% editing with 26% indels, while BE3 yielded 75% editing with 5% indels. Two additional base-edited cytosines are silent and do not yield changes to the amino acid sequence (Figure 88).

[00380] Base editors or complexes provided herein may be used to treat prion protein diseases such as Creutzfeldt-Jakob disease and fatal familial insomnia, for example, by introducing mutations into a PRNP gene. Reverting PRNP mutations may not yield therapeutic results, and indels in PRNP may be pathogenic. Accordingly, it was tested whether PRNP could be mutated using base editors (*e.g.*, BE3) to introduce a premature stop codon in the PRNP gene. BE3, associated with its guide RNA, was introduced into HEK cells

or glioblastoma cells and was capable of editing the PRNP gene to change the encoded arginine at residue 37 to a stop codon. BE3 yielded 41% editing (Figure 89).

[00381] Additional genes that may be edited include the following: *APOE* editing of Arg 112 and Arg 158 to treat increased Alzheimer's risk; *APP* editing of Ala 673 to decrease Alzheimer's risk; *PRNP* editing of Arg 37 to treat fatal familial insomnia and other prion protein diseases; *DMD* editing of the exons 23 and 51 splice sites to treat Duchenne muscular dystrophy; *FTO* editing of intron 1 to treat obesity risk; *PDS* editing of exon 8 to treat Pendred syndrome (genetic deafness); *TMCI* editing of exon 8 to treat congenital hearing loss; *CYBB* editing of various patient-relevant mutations to treat chronic granulomatous disease. Additional diseases that may be treated using the base editors provided herein are shown in Table 6, below.

[00382] UGI also plays a key role. Knocking out UDG (which UGI inhibits) was shown to dramatically improve the cleanliness and efficiency of C to T base editing (Figure 90). Furthermore, base editors with nickase and without UGI were shown to produce a mixture of outcomes, with very high indel rates (Figure 91).

EXAMPLE 12: Expanding the Targeting Scope of Base Editing

[00383] Base editing is a new approach to genome editing that uses a fusion protein containing a catalytically defective *Streptococcus pyogenes* Cas9, a cytidine deaminase, and an inhibitor of base excision repair to induce programmable, single-nucleotide C→T (or G→A) changes in DNA without generating double-strand DNA breaks, without requiring a donor DNA template, and without inducing an excess of stochastic insertions and deletions¹. The development of five new C→T (or G→A) base editors that use natural and engineered Cas9 variants with different protospacer-adjacent motif (PAM) specificities to expand the number of sites that can be targeted by base editing by 2.5-fold are described herein. Additionally, new base editors containing mutated cytidine deaminase domains that narrow the width of the apparent editing window from approximately 5 nucleotides to 1 or 2 nucleotides were engineered, enabling the discrimination of neighboring C nucleotides that would previously be edited with comparable efficiency. Together, these developments substantially increase the targeting scope of base editing.

[00384] CRISPR-Cas9 nucleases have been widely used to mediate targeted genome editing². In most genome editing applications, Cas9 forms a complex with a single guide RNA (sgRNA) and induces a double-stranded DNA break (DSB) at the target site specified by the sgRNA sequence. Cells primarily respond to this DSB through the non-homologous

end-joining (NHEJ) repair pathway, which results in stochastic insertions or deletions (indels) that can cause frameshift mutations that disrupt the gene. In the presence of a donor DNA template with a high degree of homology to the sequences flanking the DSB, gene correction can be achieved through an alternative pathway known as homology directed repair (HDR).^{3,4} Unfortunately, under most non-perturbative conditions HDR is inefficient, dependent on cell state and cell type, and dominated by a larger frequency of indels.^{3,4} As most of the known genetic variations associated with human disease are point mutations⁵, methods that can more efficiently and cleanly make precise point mutations are needed.

[00385] Base editing, which enables targeted replacement of a C:G base pair with a T:A base pair in a programmable manner without inducing DSBs¹, has been recently described. Base editing uses a fusion protein between a catalytically inactivated (dCas9) or nickase form of *Streptococcus pyogenes* Cas9 (SpCas9), a cytidine deaminase such as APOBEC1, and an inhibitor of base excision repair such as uracil glycosylase inhibitor (UGI) to convert cytidines into uridines within a five-nucleotide window specified by the sgRNA.¹ The third-generation base editor, BE3, converts C:G base pairs to T:A base pairs, including disease-relevant point mutations, in a variety of cell lines with higher efficiency and lower indel frequency than what can be achieved using other genome editing methods¹. Subsequent studies have validated the deaminase-dCas9 fusion approach in a variety of settings^{6,7}.

[00386] Efficient editing by BE3 requires the presence of an NGG PAM that places the target C within a five-nucleotide window near the PAM-distal end of the protospacer (positions 4-8, counting the PAM as positions 21-23)¹. This PAM requirement substantially limits the number of sites in the human genome that can be efficiently targeted by BE3, as many sites of interest lack an NGG 13- to 17- nucleotides downstream of the target C. Moreover, the high activity and processivity of BE3 results in conversion of all Cs within the editing window to Ts, which can potentially introduce undesired changes to the target locus. Herein, new C:G to T:A base editors that address both of these limitations are described.

[00387] It was thought that any Cas9 homolog that binds DNA and forms an “R-loop” complex⁸ containing a single-stranded DNA bubble could in principle be converted into a base editor. These new base editors would expand the number of targetable loci by allowing non-NGG PAM sites to be edited. The Cas9 homolog from *Staphylococcus aureus* (SaCas9) is considerably smaller than SpCas9 (1053 vs. 1368 residues), can mediate efficient genome editing in mammalian cells, and requires an NNGRRT PAM⁹. SpCas9 was replaced with SaCas9 in BE3 to generate SaBE3 and transfected HEK293T cells with plasmids encoding SaBE3 and sgRNAs targeting six human genomic loci (Figures 92A and 92B). After 3 d, the

genomic loci were subjected to high-throughput DNA sequencing (HTS) to quantify base editing efficiency. SaBE3 enabled C to T base editing of target Cs at a variety of genomic sites in human cells, with very high conversion efficiencies (approximately 50-75% of total DNA sequences converted from C to T, without enrichment for transfected cells) arising from targeting Cs at positions 6-11. The efficiency of SaBE3 on NNGRRT-containing target sites in general exceeded that of BE3 on NGG-containing target sites¹. Perhaps due to its higher average efficiency, SaBE3 can also result in detectable base editing at target Cs at positions outside of the canonical BE3 activity window (Figure 92C). In comparison, BE3 showed significantly reduced editing under the same conditions (0-11%), in accordance with the known SpCas9 PAM preference (Figure 106A)¹⁰. These data show that SaBE3 can facilitate very efficient base editing at sites not accessible to BE3.

[00388] The targeting range of base editors was further expanded by applying recently engineered Cas9 variants that expand or alter PAM specificities. Joung and coworkers recently reported three SpCas9 mutants that accept NGA (VQR-Cas9), NGAG (EQR-Cas9), or NGCG(VRER-Cas9) PAM sequences¹¹. In addition, Joung and coworkers engineered a SaCas9 variant containing three mutations (SaKKH-Cas9) that relax its PAM requirement to NNNRRT¹². The SpCas9 portion of BE3 was replaced with these four Cas9 variants to produce VQR-BE3, EQR-BE3, VRER-BE3, and SaKKH-BE3, which target NNNRRT, NGA, NGAG, and NGCG PAMs respectively. HEK293T cells were transfected with plasmids encoding these constructs and sgRNAs targeting six genomic loci for each new base editor, and measured C to T base conversions using HTS.

[00389] SaKKH-BE3 edited sites with NNNRRT PAMs with efficiencies up to 62% of treated, non-enriched cells (Figure 92D). As expected, SaBE3 was unable to efficiently edit targets containing PAMs that were NNNHRRT (where H = A, C, or T) (Figure 92D). VQR-BE3, EQR-BE3, and VRER-BE3 exhibited more modest, but still substantial base editing efficiencies of up to 50% of treated, non-enriched cells at genomic loci with the expected PAM requirements with an editing window similar to that of BE3 (Figures 92E and 92F). Base editing efficiencies of VQR-BE3, EQR-BE3, and VRER-BE3 in general closely paralleled the reported PAM requirements of the corresponding Cas9 nucleases; for example, EQR-BE3 was unable to efficiently edit targets containing NGAH PAM sequences (Figure 92F). In contrast, BE3 was unable to edit sites with NGA or NGCG PAMs efficiently (0-3%), likely due to its PAM restrictions (Figure 106B).

[00390] Collectively, the properties of SaBE3, SaKKH-BE3, VQR-BE3, EQR-BE3, and VRER-BE3 establish that base editors exhibit a modularity that facilitates their ability to exploit Cas9 homologs and engineered variants.

[00391] Next, base editors with altered activity window widths were developed. All Cs within the activity window of BE3 can be efficiently converted to Ts¹. The ability to modulate the width of this window would be useful in cases in which it is important to edit only a subset of Cs present in the BE3 activity window.

[00392] The length of the linker between APOBEC1 and dCas9 was previously observed to modulate the number of bases that are accessible by APOBEC1 *in vitro*¹. In HEK293T cells, however, varying the linker length did not significantly modulate the width of the editing window, suggesting that in the complex cellular milieu, the relative orientation and flexibility of dCas9 and the cytidine deaminase are not strongly determined by linker length (Figure 96). Next, it was thought that truncating the 5' end of the sgRNA might narrow the base editing window by reducing the length of single-stranded DNA accessible to the deaminase upon formation of the RNA-DNA heteroduplex. HEK293T cells were co-transfected with plasmids encoding BE3 and sgRNAs of different spacer lengths targeting a locus with multiple Cs in the editing window. No consistent changes in the width of base editing when using truncated sgRNAs with 17- to 19-base spacers were observed (Figures 95A to 95C). Truncating the sgRNA spacer to fewer than 17 bases resulted in large losses in activity (Figure 95A).

[00393] As an alternative approach, it was thought that mutations to the deaminase domain might narrow the width of the editing window through multiple possible mechanisms. First, some mutations may alter substrate binding, the conformation of bound DNA, or substrate accessibility to the active site in ways that reduce tolerance for non-optimal presentation of a C to the deaminase active site. Second, because the high activity of APOBEC1 likely contributes to the deamination of multiple Cs per DNA binding event,^{1,13,14} mutations that reduce the catalytic efficiency of the deaminase domain of a base editor might prevent it from catalyzing successive rounds of deamination before dissociating from the DNA. Once any C:G to T:A editing event has taken place, the sgRNA no longer perfectly matches the target DNA sequence and re-binding of the base editor to the target locus should be less favorable. Both strategies were tested in an effort to discover new base editors that distinguish among multiple cytidines within the original editing window.

[00394] Given the absence of an available APOBEC1 structure, several mutations previously reported to modulate the catalytic activity of APOBEC3G, a cytidine deaminase from the same family that shares 42% sequence similarity of its active site-containing domain

to that of APOBEC1, were identified¹⁵. Corresponding APOBEC1 mutations were incorporated into BE3 and evaluated their effect on base editing efficiency and editing window width in HEK293T cells at two C-rich genomic sites containing Cs at positions 3, 4, 5, 6, 8, 9, 10, 12, 13, and 14 (site A); or containing Cs at positions 5, 6, 7, 8, 9, 10, 11, and 13 (site B).

[00395] The APOBEC1 mutations R118A and W90A each led to dramatic loss of base editing efficiency (Figure 97C). R132E led to a general decrease in editing efficiency but did not change the substantially narrow the shape of the editing window (Figure 97C). In contrast, several mutations that narrowed the width of the editing window while maintaining substantial editing efficiency were found (Figures 93A and 97C). The “editing window width” was defined to represent the artificially calculated window width within which editing efficiency exceeds the half-maximal value for that target. The editing window width of BE3 for the two C-rich genomic sites tested was 5.0 (site A) and 6.1 (site B) nucleotides.

[00396] R126 in APOBEC1 is predicted to interact with the phosphate backbone of ssDNA¹³. Previous studies have shown that introducing the corresponding mutation into APOBEC3G decreased catalysis by at least 5-fold¹⁴. Interestingly, when introduced into APOBEC1 in BE3, R126A and R126E increased or maintained activity relative to BE3 at the most strongly edited positions (C5, C6, and C7), while decreasing editing activity at other positions (Figures 93A and 97C). Each of these two mutations therefore narrowed the width of the editing window at site A and site B to 4.4 and 3.4 nucleotides (R126A), or to 4.2 and 3.1 nucleotides (R126E), respectively (Figures 93A and 97C).

[00397] W90 in APOBEC1 (corresponding to W285 in APOBEC3G) is predicted to form a hydrophobic pocket in the APOBEC3G active site and assist in substrate binding¹³. Mutating this residue to Ala abrogated APOBEC3G’s catalytic activity¹³. In BE3, W90A almost completely abrogated base editing efficiency (Figure 97C). In contrast, it was found that W90Y only modestly decreased base editing activity while narrowing the editing window width at site A and site B to 3.8 and 4.9 nucleotides, respectively (Figure 93A). These results demonstrate that mutations to the cytidine deaminase domain can narrow the activity window width of the corresponding base editors.

[00398] W90Y, R126E, and R132E, the three mutations that narrowed the editing window without drastically reducing base editing activity, were combined into doubly and triply mutated base editors. The double mutant W90Y+R126E resulted in a base editor (YE1-BE3) with BE3-like maximal editing efficiencies, but substantially narrowed editing window width (width at site A and site B = 2.9 and 3.0 nucleotides, respectively (Figure 93A). The

W90Y+R132E base editor (YE2-BE3) exhibited modestly lower editing efficiencies (averaging 1.4-fold lower maximal editing yields across the five sites tested compared with BE3), and also substantially narrowed editing window width (width at site A and site B = 2.7 and 2.8 nucleotides, respectively) (Figure 97C). The R126E+R132E double mutant (EE-BE3) showed similar maximal editing efficiencies and editing window width as YE2-BE3 (Figure 97C). The triple mutant W90Y+R126E+R132E (YEE-BE3) exhibited 2.0-fold lower average maximal editing yields but very little editing beyond the C6 position and an editing window width of 2.1 and 1.4 nucleotides for site A and site B, respectively (Figure 97C). These data taken together indicate that mutations in the cytidine deaminase domain can strongly affect editing window widths, in some cases with minimal or only modest effects on editing efficiency.

[00399] The base editing outcomes of BE3, YE1-BE3, YE2-BE3, EE-BE3, and YEE-BE3 were further compared in HEK293T cells targeting four well-studied human genomic sites that contain multiple Cs within the BE3 activity window¹. These target loci contained target Cs at positions 4 and 5 (HEK site 3), positions 4 and 6 (HEK site 2), positions 5 and 6 (EMX1), or positions 6, 7, 8, and 11 (FANCF). BE3 exhibited little (< 1.2-fold) preference for editing any Cs within the position 4-8 activity window. In contrast, YE1-BE3, exhibited a 1.3-fold preference for editing C5 over C4 (HEK site 3), 2.6-fold preference for C6 over C4 (HEK site 2), 2.0-fold preference for C5 over C6 (EMX1), and 1.5-fold preference for C6 over C7 (FANCF) (Figure 93B). YE2-BE3 and EE-BE3 exhibited somewhat greater positional specificity (narrower activity window) than YE1-BE3, averaging 2.4-fold preference for editing C5 over C4 (HEK site 3), 9.5-fold preference for C6 over C4 (HEK site 2), 2.9-fold preference for C5 over C6 (EMX1), and 2.6-fold preference for C7 over C6 (FANCF) (Figure 93B). YEE-BE3 showed the greatest positional selectivity, with a 2.9-fold preference for editing C5 over C4 (HEK site 3), 29.7-fold preference for C6 over C4 (HEK site 2), 7.9-fold preference for C5 over C6 (EMX1), and 7.9-fold preference for C7 over C6 (FANCF) (Figure 93B). The findings establish that mutant base editors can discriminate between adjacent Cs, even when both nucleotides are within the BE3 editing window.

[00400] The product distributions of these four mutants and BE3 were further analyzed by HTS to evaluate their apparent processivity. BE3 generated predominantly T4-T5 (HEK site 3), T4-T6 (HEK site 2), and T5-T6 (EMX1) products in treated HEK293T cells, resulting in, on average, 7.4-fold more products containing two Ts, than products containing a single T. In contrast, YE1-BE3, YE2-BE3, EE-BE3, and YEE-BE3 showed substantially higher preferences for singly edited C4-T5, C4-T6, and T5-C6 products (Figure 93C). YE1-BE3

yielded products with an average single-T to double-T product ratio of 1.4. YE2-BE3 and EE-BE3 yielded products with an average single-T to double-T product ratio of 4.3 and 5.1, respectively (Figure 93C). Consistent with the above results, the YEE-BE3 triple mutant favored single-T products by an average of 14.3-fold across the three genomic loci. (Figure 93C). For the target site in which only one C is within the target window (HEK site 4, at position C5), all four mutants exhibited comparable editing efficiencies as BE3 (Figure 98). These findings indicate that these BE3 mutants have decreased apparent processivity and can favor the conversion of only a single C at target sites containing multiple Cs within the BE3 editing window. These data also suggest a positional preference of $C5 > C6 > C7 \approx C4$ for these mutant base editors, although this preference could differ depending on the target sequence.

[00401] The window-modulating mutations in APOBEC1 were applied to VQR-BE3, allowing selective base editing of substrates at sites targeted by NGA PAM (Figure 107A). However, when these mutations were applied to SaKKH-BE3, a linear decrease in base editing efficiency was observed without the improvement in substrate selectivity, suggesting a different kinetic equilibrium and substrate accessibility of this base editor than those of BE3 and its variants (Figure 107B).

[00402] The five base editors with altered PAM specificities described in this study together increase the number of disease-associated mutations in the ClinVar database that can in principle be corrected by base editing by 2.5-fold (Figures 94A and 94B). Similarly, the development of base editors with narrowed editing windows approximately doubles the fraction of ClinVar entries with a properly positioned NGG PAM that can be corrected by base editing without comparable modification of a non-target C (from 31% for BE3 to 59% for YEE-BE3) (Figures 94A and 94B).

[00403] In summary, the targeting scope of base editing was substantially expanded by developing base editors that use Cas9 variants with different PAM specificities, and by developing a collection of deaminase mutants with varying editing window widths. In theory, base editing should be possible using other programmable DNA-binding proteins (such as Cpf1¹⁶) that create a bubble of single-stranded DNA that can serve as a substrate for a single-strand-specific nucleotide deaminase enzyme.

Materials and Methods

[00404] *Cloning.* PCR was performed using Q5 Hot Start High-Fidelity DNA Polymerase (New England Biolabs). Plasmids for BE and sgRNA were constructed using USER cloning

(New England Biolabs), obtained from previously reported plasmids¹. DNA vector amplification was carried out using NEB 10beta competent cells (New England Biolabs).

[00405] *Cell culture.* HEK293T (ATCC CRL-3216) were cultured in Dulbecco's Modified Eagle's Medium plus GlutaMax (ThermoFisher) supplemented with 10% (v/v) fetal bovine serum (FBS), at 37 °C with 5% CO₂. Immortalized rat astrocytes containing the ApoE4 isoform of the *APOE* gene (Taconic Biosciences) were maintained in Dulbecco's Modified Eagle's Medium plus GlutaMax (ThermoFisher Scientific) supplemented with 10% (v/v) fetal bovine serum (FBS) and 200 µg/mL Geneticin (ThermoFisher Scientific).

[00406] *Transfections.* HEK293T cells were seeded on 48-well collagen-coated BioCoat plates (Corning) and transfected at approximately 85% confluency. 750 ng of BE and 250 ng of sgRNA expression plasmids were transfected using 1.5 µl of Lipofectamine 2000 (ThermoFisher Scientific) per well according to the manufacturer's protocol.

[00407] *High-throughput DNA sequencing of genomic DNA samples.* Transfected cells were harvested after 3 d and the genomic DNA was isolated using the Agencourt DNAdvance Genomic DNA Isolation Kit (Beckman Coulter) according to the manufacturer's instructions. Genomic regions of interest were amplified by PCR with flanking HTS primer pairs listed in the Supplementary Sequences. PCR amplification was carried out with Phusion hot-start II DNA polymerase (ThermoFisher) according to the manufacturer's instructions. PCR products were purified using RapidTips (Diffinity Genomics). Secondary PCR was performed to attach sequencing adaptors. The products were gel-purified and quantified using the KAPA Library Quantification Kit-Illumina (KAPA Biosystems). Samples were sequenced on an Illumina MiSeq as previously described¹.

[00408] *Data analysis.* Nucleotide frequencies were assessed using a previously described MATLAB script¹. Briefly, the reads were aligned to the reference sequence via the Smith-Waterman algorithm. Base calls with Q-scores below 30 were replaced with a placeholder nucleotide (N). This quality threshold results in nucleotide frequencies with an expected theoretical error rate of 1 in 1000.

[00409] Analyses of base editing processivity were performed using a custom python script. This program trims sequencing reads to the 20 nucleotide protospacer sequence as determined by a perfect match for the 7 nucleotide sequences that should flank the target site. These targets were then consolidated and sorted by abundance to assess the frequency of base editing products.

[00410] Bioinformatic analysis of the ClinVar database of human disease-associated mutations was performed in a manner similar to that previously described but with small

adjustments¹. These adjustments enable the identification of targets with PAMs of customizable length and sequence. In addition, this improved script includes a priority ranking of target C positions (C5 > C6 > C7 > C8 ≈ C4), thus enabling the identification of target sites in which the on-target C is either the only cytosine within the window or is placed at a position with higher predicted editing efficiency than any off-target C within the editing window.

References for Example 12

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EXAMPLE 13:

[00411] Using improved transfection procedures and better plasmids, biological replicates (n=3) were used to install the four HF mutations into the Cas9 portion of BE3. The mutations do not significantly effect on-targeting editing with plasmid delivery (Figure 99). At the tested concentration, BE3 protein delivery works; however, the on-target editing is lower than for plasmid delivery (Figure 100). Protein delivery of BE3 with the HF mutations installed reduces on-targeting editing efficiency but still yields some edited cells (Figure 101).

[00412] Both lipofection and installing HF mutations were shown to decrease off-target deamination events. For the four sites shown in Figure 102, the off-target site (OT) with the highest GUIDE-Seq reads and deamination events were assayed (Komor *et al.*, *Nature*, 2016). The specificity ratio was calculated by dividing the off-target editing by the on-target editing at the closest corresponding C. In cases where off-target editing was not detectable, the ratio was set to 100. Thus, a higher specificity ratio indicates a more specific construct. BE3 plasmid delivery showed much higher off-target/on-target editing than protein delivery of BE3, plasmid delivery of HF-BE3, or protein delivery of HF-BE3 (Figures 102 and 105).

[00413] Purified proteins HF-BE3 and BE3 were analyzed *in vitro* for their capabilities to convert C to T residues at different positions in the spacer with the most permissive motif. Both BE3 and HF-BE3 proteins were found to have the same “window” for base editing (Figures 103 and 104).

[00414] A list of the disease targets is given in Table 9. The base to be edited in Table 9 is indicated in bold and underlined.

Table 9. Base Editor Disease Targets

GENE	DISEASE	SPACER	PAM	EDITOR	DEFECT	CELL
RB1	RETINOBLASTOMA	AAT <u>C</u> TAGTAAATAAA TTGATGT	AAAA GT	SAKKH- BE3	SPLICING IMPAIRMENT	J82
PTEN	CANCER	GACCA <u>A</u> CGGCTAAGT GAAGA	TGA	VQR- BE3	W111R	MC116
PIK3C A	CANCER	TC <u>C</u> TTTCTTCACGGTT GCCT	ACTG GT	SAKKH- BE3	K111R	CRL- 5853
PIK3C A	CANCER	CTC <u>C</u> TGCTCAGTGATT TCAG	AGA	VQR- BE3	Q546R	CRL- 2505
TP53	CANCER	TGT <u>C</u> CACACATGTAGTT GTAG	TGG	YEE-BE3	N239D	SNU47 5
HRAS	CANCER	CCTCC <u>C</u> GGCCGGCGG TATCC	AGG	YEE-BE3	Q61R	MC/C AR

Table 6. Exemplary diseases that may be treated using base editors. The protospacer and PAM sequences are shown in the sgRNA (PAM) column. The PAM sequence is shown in parentheses and with the base to be edited indicated by underlining.

Disease target	gene symbol	Base changed	sgRNA (PAM)	Base editor
Prion disease	PRNP	R37*	GGCAGCCGATACCCGGGGCA(GGG) GGGCAGCCGATACCCGGGGC(AGG)	BE3
Pendred syndrome	Slc26a4	c.919-2A>G	TTATTGTCGAAATAAAAGA(AGA) ATTGTCGAAATAAAAGAAG(AGG) TTGTCCGAAATAAAAGAAGA(GGA) GTCCGAAATAAAAGAAGAGGAAAA(AAT) GTCCGAAATAAAAGAAGAGGAAAA(ATT)	BE3 (VQR SaCas9)
Congenital deafness	Tmc1	c.545A>G	CAGGAAGCACGAGGCCACTG(AGG) AACAGGAAGCACGAGGCCAC(TGA) AGGAAGCACGAGGCCACTGA(GGA)	BE3 YE-BE3 YEE-BE3
Acquired deafness	SNHL	S33F	TTGGATTCTGGATCCATTC(TGG)	BE3
Alzheimer's Disease	APP	A673T	TCTGCATCCATCTTCACTTC(AGA)	BE3 VQR
Niemann-Pick Disease Type C	NPC1	I1061T	CTTACAGCCAGTAATGTAC(CGA)	BE3 VQR

[00415] Additional exemplary genes in the human genome that may be targeted by the base editors or complexes of this disclosure are provided herein in Tables 7 and 8. Table 7 includes gene mutations that may be corrected by changing a cytosine (C) to a thymine (T), for example, using a BE3 nucleobase editor. Table 8 includes gene mutations that may be corrected by changing a guanine (G) to an adenine (A), for example, using a BE3 nucleobase editor.

Table 7. Human gene mutations that may be corrected by changing a cytosine (C) to a thymine (T). The gene name, gene symbol, and dbSNP database reference number (RS#) are indicated. Also indicated are exemplary protospacers with their PAM sequences (gRNAs and gRNAall) and the base to be edited, e.g., a C, indicated by a “Y”. The “gRNAs” sequences, from top to bottom, correspond to SEQ ID NOs: 1914-2091. The “gRNA all” sequences, from top to bottom, correspond to SEQ ID NOs: 2192-2540, 3144-3433.

Name	RS# (dbSNP)	GeneSymbol	gRNAs	gRNAall	Phenotypes
NM_000138.4(FBN1): c.3220T>C (p.Cys1074Arg)	137854465	FBN1	[]	[]	[]
NM_001927.3(DES):c.1154T>C (p.Leu385Pro)	57955682	DES	[]	[]	['Myofibrillar myopathy 1', 'not provided']
NM_025152.2(NUBPL):c.815-27T>C	118161496	NUBPL	[]	['TGGTTCYAATGG ATGTCTGCTGG', 'GGTTCYAATGGA TGCTGCTGGG']	['Mitochondrial complex I deficiency', 'not provided']
NM_003000.2(SDHB):c.574T>C (p.Cys192Arg)	786202732	SDHB	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_004586.2(RPS6KA3):c.803T>C (p.Phe268Ser)	122454131	RPS6KA3	[]	[]	['Coffin-Lowry syndrome']

NM_005609.2(PYGM):c.425_528del	764313717	PYGM	[]	['TGGCTGYCAGG GACCCAGCAAGG', 'CTGYCAGGGACC CAGCAAGGAGG']	[]
NM_000124.3(ERCC6):c.2830-2A>G	373227647	ERCC6	[]	[]	['Cockayne syndrome, type B']
NM_000059.3(BRCA2):c.316+2T>C	81002805	BRCA2	['CTTAGGYAAGTAATGCAATATGG']	['CTTAGGYAAGTAATGCAATATGG']	['Familial cancer of breast', 'Breast-ovarian cancer, familial 2', 'Hereditary cancer-predisposing syndrome']
NM_003242.5(TGFBR2):c.923T>C (p.Leu308Pro)	28934568	TGFBR2	[]	['AGTTCCYGACGGCTGAGGAGCGG']	['Loeys-Dietz syndrome 2']
NM_000410.3(HFE):c.314T>C (p.Ile105Thr)	28934596	HFE	[]	[]	['Hemochromatosis type 1']
NM_000308.2(CTSA):c.247T>C (p.Trp83Arg)	28934603	CTSA	[]	[]	['Combined deficiency of sialidase AND beta galactosidase']
NM_033290.3(MID1):c.1877T>C (p.Leu626Pro)	28934611	MID1	[]	[]	['Opitz-Frias syndrome']
NM_000329.2(RPE65):c.1102T>C (p.Tyr368His)	62653011	RPE65	[]	[]	['Leber congenital amaurosis 2', 'Retinitis pigmentosa 20', 'not provided']
NM_007313.2(ABL1):c.814T>C (p.Tyr272His)	121913461	ABL1	[]	['CCAGYACGGGGAGGTGTACGAGG', 'CAGYACGGGGAGGTGTACGAGGG']	[]
NM_000546.5(TP53):c.398T>C (p.Met133Thr)	28934873	TP53	[]	[]	['Li-Fraumeni syndrome 1']
NM_000490.4(AVP):c.200T>C (p.Val67Ala)	28934878	AVP	[]	[]	['Neurohypophyseal diabetes insipidus']
NM_021961.5(TEAD1):c.1261T>C (p.Tyr?His)	11567847	TEAD1	['TCATATTYACAGGCTTGTAAGG']	['TCATATTYACAGGCTTGTAAGG']	[]
NM_002609.3(PDGFRB):c.1973T>C (p.Leu658Pro)	397509381	PDGFRB	[]	[]	['Basal ganglia calcification, idiopathic, 4']
NM_005236.2(ERCC4):c.689T>C (p.Leu230Pro)	397509402	ERCC4	[]	[]	['Fanconi anemia, complementation group Q']
NM_005236.2(ERCC4):c.706T>C (p.Cys236Arg)	397509403	ERCC4	[]	[]	['XERODERMA PIGMENTOSUM, TYPE F/COCKAYNE SYNDROME']
NM_173551.4(ANKS6):c.1322A>G (p.Gln441Arg)	377750405	ANKS6	[]	['AGGGCYGTCGGACCTTCGAGTGG', 'GGGCYGTCGGACCTTCGAGTGGG',	['Nephronophthisis 16']

				'GGCYGTCGGACC TTCGAGTGGGG']	
NM_000142.4(FGFR3):c.1612A>G (p.Ile538Val)	80053154	FGFR3	[]	[]	['Hypochondroplasia']
NM_000441.1(SLC26A4):c.707T>C (p.Leu236Pro)	80338848	SLC26A4	[]	[]	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_000518.4(HBB):c.337T>C (p.Cys113Arg)	35849199	HBB	[]	[]	[]
NM_000104.3(CYP1B1):c.2T>C (p.Met1Thr)	72549389	CYP1B1	[]	[]	['Irido-corneo-trabecular dysgenesis']
NM_000169.2(GLA):c.484T>C (p.Trp162Arg)	28935196	-	[]	[]	['Fabry disease']
NM_001927.3(DES):c.1034T>C (p.Leu345Pro)	57639980	DES	[]	['ATTCCCYGATGAGGCAGATGCGG', 'TCCCYGATGAGGCAGATGCGGG']	['Myofibrillar myopathy 1', 'not provided']
NM_006517.4(SLC16A2):c.1190T>C (p.Leu397Pro)	122455132	SLC16A2	[]	[]	['Allan-Herndon-Dudley syndrome']
NM_020320.3(RARS2):c.35A>G (p.Gln12Arg)	147391618	RARS2	[]	['ATACCYGGCAA GCAATAGCGCGG']	['Pontocerebellar hypoplasia type 6']
NM_000239.2(LYZ):c.221T>C (p.Ile74Thr)	121913547	LYZ	[]	[]	['Familial visceral amyloidosis, Ostertag type']
NM_002977.3(SCN9A):c.2215A>G (p.Ile739Val)	182650126	-	[]	['GTAAYTGCAAG ATCTACAAAAGG']	['Small fiber neuropathy', 'not provided']
NM_004700.3(KCNQ4):c.842T>C (p.Leu281Ser)	80358278	KCNQ4	[]	['ACATYGACAAC CATCGGCTATGG']	['DFNA 2 Nonsyndromic Hearing Loss']
NM_000169.2(GLA):c.806T>C (p.Val269Ala)	28935488	-	['CAGTT AGYGA TTGGC AACTT TGG']	['CAGTTAGYGATT GGCAACTTTGG']	['Fabry disease']
NM_000228.2(LAMB3):c.565-2A>G	370148688	LAMB3	[]	[]	['Junctional epidermolysis bullosa gravis of Herlitz']
NM_052867.2(NALCN):c.1526T>C (p.Leu509Ser)	786203987	NALCN	[]	[]	['CONGENITAL CONTRACTURES OF THE LIMBS AND FACE, HYPOTONIA, AND DEVELOPMENTAL DELAY']
NM_001031.4(RPS28):c.1A>G (p.Met1Val)	786203997	RPS28	['CCAY GATGG CGGCG CGGCG GCGG']	['TGTCCAYGATGG CGGCGGCGG', 'CCAYGATGGCGG CGCGGCGGCGG']	['Diamond-Blackfan anemia with microtia and cleft palate']
NM_005957.4(MTHF)	786204012	MTHFR	[]	['GACCYGCTGCCG']	['Homocysteinemia']

R):c.388T>C (p.Cys130Arg)				TCAGCGCCTGG']	due to MTHFR deficiency']
NM_005957.4(MTHFR):c.1530+2T>C	786204027	MTHFR	['GAAGGYGTGGTGTAGG GAGGCACGG', 'AAGGYGTGGT AGGGA GGCACGGG', 'AGGYGTGGTAGG GAGGCACGGGG']	['GAAGGYGTGGT AGGGAGGCACGG', 'AAGGYGTGGTAG GGAGGCACGGG', 'AGGYGTGGTAGG GAGGCACGGGG']	['Homocysteinemia due to MTHFR deficiency']
NM_005957.4(MTHFR):c.1793T>C (p.Leu598Pro)	786204034	MTHFR	[]	[]	['Homocysteinemia due to MTHFR deficiency']
NM_005957.4(MTHFR):c.1883T>C (p.Leu628Pro)	786204037	MTHFR	[]	['TCCCACYGGACA ACTGCCTCTGG']	['Homocysteinemia due to MTHFR deficiency']
NM_000264.3(PTCH1):c.3168+2T>C	786204056	PTCH1	['ATCAT TGYGA GTGTA TTATA AGG', 'TCATT GYGAG TGTAT TATAA GGG', 'CATTG YGAGT GTATT ATAAG GGG']	['ATCATTGYGAGT GTATTATAAGG', 'TCATTGYGAGTGT TATTATAAGGG', 'CATTGYGAGTGT ATTATAAGGGG']	['Gorlin syndrome']
NM_000182.4(HADHA):c.919-2A>G	200017313	HADHA	[]	[]	['Mitochondrial trifunctional protein deficiency', 'Long-chain 3-hydroxyacyl-CoA dehydrogenase deficiency', 'not provided']
NM_000030.2(AGXT):c.806T>C (p.Leu269Pro)	180177271	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_006121.3(KRT1):c.1436T>C (p.Ile479Thr)	57837128	KRT1	[]	[]	['Ichthyosis, cyclic, with epidermolytic hyperkeratosis', 'not provided']
NM_000521.3(HEXB):c.185C>T (p.Ser62Leu)	820878	HEXB	[]	[]	['Sandhoff disease, infantile type']
NM_000140.3(FECH):c.1137+3A>G	202147607	FECH	[]	['GTAGAYACCTTA GAGAACAATGG']	['Erythropoietic protoporphyria']
NM_015046.5(SETX):c.1166T>C	29001584	SETX	[]	[]	['Amyotrophic lateral sclerosis type

(p.Leu389Ser)					4']
NM_020365.4(EIF2B3):c.1037T>C (p.Ile346Thr)	119474039	EIF2B3	['CCAG AYTGT CAGCA AACAC CTGG']	['CCAGAYTGTCAG CAAACACCTGG']	['Leukoencephalopathy with vanishing white matter']
NM_139058.2(ARX):c.98T>C (p.Leu33Pro)	28936077	ARX	[]	[]	['Mental retardation, with or without seizures, ARX-related, X-linked']
NM_005183.3(CACNA1F):c.2267T>C (p.Ile756Thr)	122456136	CACNA1F	[]	['TGCCAYTGCTGT GGACAACCTGG']	[]
NM_007374.2(SIX6):c.110T>C (p.Leu37Pro)	786204851	SIX6	[]	['GTCGCGCCCGT GGCCCCTGCGG']	['Cataract, microphthalmia and nystagmus']
NM_000339.2(SLC12A3):c.1261T>C (p.Cys421Arg)	28936387	SLC12A3	[]	[]	['Familial hypokalemia-hypomagnesemia']
NM_003865.2(HESX1):c.77T>C (p.Ile26Thr)	28936416	HESX1	[]	[]	['Pituitary hormone deficiency, combined 5']
NM_022114.3(PRDM16):c.2660T>C (p.Leu887Pro)	202115331	PRDM16	[]	[]	['Dilated cardiomyopathy 1LL']
NM_001159287.1(TPI1):c.832T>C (p.Phe278Leu)	121964847	TPI1	[]	[]	['Triosephosphate isomerase deficiency']
NM_001692.3(ATP6V1B1):c.242T>C (p.Leu81Pro)	121964880	ATP6V1B1	[]	[]	[]
NM_000490.4(AVP):c.61T>C (p.Tyr21His)	121964893	AVP	[]	[]	['Neurohypophyseal diabetes insipidus']
NM_000027.3(AGA):c.916T>C (p.Cys306Arg)	121964906	AGA	['GTTAT AYGTC CCAAT GTGAC TGG']	['GTTATAYGTGCC AATGTGACTGG']	['Aspartylglycosaminuria']
NM_000138.4(FBN1):c.1468+2T>C	794728167	FBN1	[]	['ATTGGYACGTGA TCCATCCTAGG']	['Thoracic aortic aneurysms and aortic dissections']
NM_000027.3(AGA):c.214T>C (p.Ser72Pro)	121964909	AGA	[]	['GACGGCYCTGTA GGCTTTGGAGG']	['Aspartylglycosaminuria']
NM_004453.3(ETFDH):c.1001T>C (p.Leu334Pro)	377686388	ETFDH	[]	[]	['Glutaric aciduria, type 2']
NM_001385.2(DPYS):c.1078T>C (p.Trp360Arg)	121964924	DPYS	['CGTA ATAYG GGAAA AAGGC GTGG', 'AATAY GGGAA AAAGG CGTGG TGG', 'ATAYG GGAAA AAGGC	['CGTAATAYGGG AAAAAGGCGTGG', 'AATAYGGGAAAA AGGCGTGGTGG', 'ATAYGGGAAAA GGCGTGGTGGG']	['Dihydropyrimidase deficiency']

			GTGGT GGG']		
NM_004453.3(ETFDH):c.2T>C (p.Met1Thr)	121964953	ETFDH	[]	[]	['Glutaric acidemia IIC']
NM_000071.2(CBS):c.1616T>C (p.Leu539Ser)	121964968	CBS	[]	[]	['Homocystinuria, pyridoxine-responsive']
NM_000170.2(GLDC):c.2T>C (p.Met1Thr)	121964978	GLDC	[]	['CGGCCAYGCAGTCCTGTGCCAGG', 'GGCCAYGCAGTCCTGTGCCAGGG']	['Non-ketotic hyperglycinemia']
NM_000108.4(DLD):c.1178T>C (p.Ile393Thr)	121964991	DLD	[]	[]	['Maple syrup urine disease, type 3']
NM_014425.3(INVS):c.1478T>C (p.Leu493Ser)	121964995	INVS	[]	[]	['Infantile nephronophthisis']
NM_000398.6(CYB5R3):c.382T>C (p.Ser128Pro)	121965006	CYB5R3	[]	[]	['Methemoglobinemia type 2']
NM_000398.6(CYB5R3):c.446T>C (p.Leu149Pro)	121965008	CYB5R3	[]	['CTGTCYGGTCTAC CAGGGCAAAGG']	['METHEMOGLOBINEMIA, TYPE I']
NM_000398.6(CYB5R3):c.610T>C (p.Cys204Arg)	121965011	CYB5R3	[]	[]	['Methemoglobinemia type 2']
NM_000398.6(CYB5R3):c.218T>C (p.Leu73Pro)	121965013	CYB5R3	[]	[]	['METHEMOGLOBINEMIA, TYPE I']
NM_001103.3(ACTN2):c.683T>C (p.Met228Thr)	786205144	ACTN2	['CCTAA AAAYG TTGGA TGCTG AAGG']	['CCTAAAAYGTTG GATGCTGAAGG']	['Dilated cardiomyopathy 1AA']
NM_000548.3(TSC2):c.3106T>C (p.Ser1036Pro)	45517281	TSC2	[]	[]	['Tuberous sclerosis syndrome', 'Tuberous sclerosis 2']
NM_000203.4(IDUA):c.1469T>C (p.Leu490Pro)	121965027	IDUA	[]	[]	['Mucopolysaccharidosis, MPS-I-H/S', 'Hurler syndrome', 'not provided']
NM_001122764.1(PPOX):c.35T>C (p.Ile12Thr)	28936677	PPOX	[]	[]	['Variegate porphyria']
NM_000525.3(KCNJ11):c.440T>C (p.Leu147Pro)	28936678	KCNJ11	[]	[]	['Islet cell hyperplasia']
NM_001025107.2(ADAR):c.1883T>C (p.Leu628Pro)	28936680	ADAR	[]	[]	['Symmetrical dyschromatosis of extremities']
NM_001025107.2(ADAR):c.2609T>C (p.Phe870Ser)	28936681	ADAR	[]	[]	['Symmetrical dyschromatosis of extremities']
NM_000557.4(GDF5):c.1322T>C (p.Leu441Pro)	28936683	-	[]	[]	['Brachydactyly type A2', 'Fibular hypoplasia and complex brachydactyly']

NM_000274.3(OAT):c.163T>C (p.Tyr55His)	121965037	OAT	[]	[]	['Ornithine aminotransferase deficiency']
NM_000274.3(OAT):c.1205T>C (p.Leu402Pro)	121965043	OAT	[]	[]	['Ornithine aminotransferase deficiency']
NM_000223.3(KRT12):c.386T>C (p.Met129Thr)	28936695	KRT12	[]	[]	['Meesman corneal dystrophy', 'not provided']
NM_000128.3(F11):c.901T>C (p.Phe301Leu)	121965064	F11	[]	['TGATYTCCTGGGAGAAGAAGCTGG']	['Hereditary factor XI deficiency disease']
NM_000128.3(F11):c.166T>C (p.Cys56Arg)	121965069	F11	[]	[]	['Hereditary factor XI deficiency disease']
NM_000235.3(LIPA):c.599T>C (p.Leu200Pro)	121965086	LIPA	[]	[]	['Lysosomal acid lipase deficiency']
NM_001199.3(BMP1):c.*241T>C	786205217	BMP1	[]	[]	['Osteogenesis imperfecta type 13']
NM_004974.3(KCNA2):c.788T>C (p.Ile263Thr)	786205231	KCNA2	[]	[]	['EPILEPTIC ENCEPHALOPATHY, EARLY INFANTILE, 32']
NM_000548.3(TSC2):c.5150T>C (p.Leu1717Pro)	45517398	TSC2	[]	['GCCCYGCACGC AAATGTGAGTGG', 'CCCYGCACGCAA ATGTGAGTGGG']	['Tuberous sclerosis syndrome', 'not provided']
NM_000212.2(ITGB3):c.176T>C (p.Leu59Pro)	5918	ITGB3	[]	[]	['Myocardial infarction', 'Posttransfusion purpura', 'Thrombocytopenia, neonatal alloimmune', 'Fracture, hip, susceptibility to']
m.9191T>C	386829069	MT-ATP6	[]	[]	['Leigh disease']
NM_000419.3(ITGA2B):c.1787T>C (p.Ile596Thr)	76811038	ITGA2B	[]	[]	['Glanzmann thrombasthenia']
NM_002294.2(LAMP2):c.864+2T>C	730880485	LAMP2	[]	[]	['Cardiomyopathy', 'Danon disease']
NM_000138.4(FBN1):c.7111T>C (p.Trp2371Arg)	794728264	FBN1	[]	[]	['Thoracic aortic aneurysms and aortic dissections']
NM_000531.5(OTC):c.143T>C (p.Phe48Ser)	72554315	OTC	[]	[]	['not provided']
NM_178454.4(DRAM2):c.79T>C (p.Tyr27His)	786205662	DRAM2	[]	[]	['Retinal dystrophy']
NM_000138.4(FBN1):c.7531T>C (p.Cys2511Arg)	794728272	FBN1	[]	[]	['Thoracic aortic aneurysms and aortic dissections']
NM_016218.2(POLK):c.609T>C (p.Asn203=)	786205684	POLK	[]	[]	['Malignant tumor of prostate']
NM_016218.2(POLK):c.*66T>C	786205688	POLK	[]	[]	['Malignant tumor of prostate']
NM_000354.5(SERP1)	28937312	SERPINA7	[]	[]	[]

NA7):c.740T>C (p.Leu247Pro)					
NM_000531.5(OTC):c.284T>C (p.Leu95Ser)	72554346	OTC	['ACAA GATYG TCTAC AGAAA CAGG']	['ACAAGATYGTCT ACAGAAACAGG']	['not provided']
NM_015662.2(IFT172):c.770T>C (p.Leu257Pro)	786205857	IFT172	[]	['TTGTGCYAGGAA GTTATGACAGG']	['RETINITIS PIGMENTOSA 71']
NM_000531.5(OTC):c.386+2T>C	72554359	OTC	[]	[]	['not provided']
NM_001135669.1(XPR1):c.434T>C (p.Leu145Pro)	786205901	XPR1	[]	[]	['BASAL GANGLIA CALCIFICATION, IDIOPATHIC, 6']
NM_001135669.1(XPR1):c.419T>C (p.Leu140Pro)	786205903	XPR1	[]	[]	['BASAL GANGLIA CALCIFICATION, IDIOPATHIC, 6']
NM_001135669.1(XPR1):c.653T>C (p.Leu218Ser)	786205904	XPR1	[]	['GCGTTYACGTGT CCCCCCTTTGG', 'CGTTYACGTGTC CCCCCCTTTGGG']	['BASAL GANGLIA CALCIFICATION, IDIOPATHIC, 6']
NM_181457.3(PAX3): c.268T>C (p.Tyr90His)	104893654	PAX3	[]	[]	['Klein-Waardenberg syndrome']
NM_001987.4(ETV6): c.1046T>C (p.Leu349Pro)	786205155	ETV6	[]	[]	['Thrombocytopenia', 'LEUKEMIA, ACUTE LYMPHOBLASTIC : ALL']
NM_000055.2(BCHE): c.1004T>C (p.Leu335Pro)	104893684	BCHE	[]	[]	['Deficiency of butyrylcholine esterase']
NM_000388.3(CASR): c.382T>C (p.Phe128Leu)	104893696	CASR	[]	[]	['Hypocalcemia, autosomal dominant 1']
NM_000388.3(CASR): c.1835T>C (p.Phe612Ser)	104893698	CASR	[]	[]	['Hypocalcemia, autosomal dominant 1']
NM_000388.3(CASR): c.2641T>C (p.Phe881Leu)	104893704	CASR	[]	['ACGCTYTCAAGG TGGCTGCCCGG', 'CGCTYTCAAGGT GGCTGCCCGGG']	['Hypercalciuric hypercalcemia']
NM_000388.3(CASR): c.374T>C (p.Leu125Pro)	104893708	CASR	[]	[]	['Hypocalcemia, autosomal dominant 1', 'Hypocalcemia, autosomal dominant 1, with bartter syndrome']
NM_000388.3(CASR): c.2362T>C (p.Phe788Leu)	104893711	CASR	[]	[]	['Hypocalcemia, autosomal dominant 1']
NM_000388.3(CASR): c.38T>C (p.Leu13Pro)	104893717	CASR	[]	[]	['Hypercalciuric hypercalcemia, familial, type 1']
NM_006580.3(CLDN16):c.500T>C	104893725	CLDN16	[]	[]	['Primary hypomagnesemia']

(p.Leu167Pro)					
NM_006580.3(CLDN16):c.434T>C (p.Leu145Pro)	104893731	CLDN16	[]	[]	['Primary hypomagnesemia']
NM_000041.3(APOE):c.388T>C (p.Cys130Arg)	429358	APOE	[]	[]	['Familial type 3 hyperlipoproteinemia']
NM_198159.2(MITF):c.1051T>C (p.Ser351Pro)	104893744	MITF	[]	[]	['Waardenburg syndrome type 2A']
NM_198159.2(MITF):c.1195T>C (p.Ser399Pro)	104893747	MITF	[]	['ACTTYCCCTTAT TCCATCCACGG', 'CTTYCCCTTATTC CATCCACGGG']	['Waardenburg syndrome type 2A']
NM_001122757.2(POU1F1):c.655T>C (p.Trp219Arg)	104893758	POU1F1	[]	[]	['Pituitary hormone deficiency, combined 1']
NM_000539.3(RHO):c.133T>C (p.Phe45Leu)	104893770	RHO	[]	['CATGYTTCTGCT GATCGTGCTGG', 'ATGYTTCTGCTG ATCGTGCTGGG']	['Retinitis pigmentosa 4']
NM_003106.3(SOX2):c.290T>C (p.Leu97Pro)	104893802	-	[]	[]	['Microphthalmia syndromic 3']
NM_024009.2(GJB3):c.101T>C (p.Leu34Pro)	28937583	GJB3	[]	[]	['Erythrokeratoderma variabilis']
NM_003907.2(EIF2B5):c.1882T>C (p.Trp628Arg)	28937596	EIF2B5	[]	['AGGCCYGGAGC CCTGTTTTAGG']	['Leukoencephalopathy with vanishing white matter']
NM_000551.3(VHL):c.188T>C (p.Leu63Pro)	104893827	VHL	[]	[]	['Pheochromocytoma']
NM_000320.2(QDPR):c.106T>C (p.Trp36Arg)	104893865	QDPR	[]	[]	['Dihydropteridine reductase deficiency']
NM_001151.3(SLC25A4):c.293T>C (p.Leu98Pro)	104893876	SLC25A4	[]	['GCAGCYCTTCTT AGGGGGTGTGG']	['Autosomal dominant progressive external ophthalmoplegia with mitochondrial DNA deletions 2']
NM_006005.3(WFS1):c.2486T>C (p.Leu829Pro)	104893883	WFS1	[]	['ACCATCCYGGA GGGCCGCCTGGG']	['WFS1-Related Disorders']
NM_001018077.1(NR3C1):c.1712T>C (p.Val571Ala)	104893911	NR3C1	['AAGY GATTG CAGCA GTGAA ATGG']	['AAGYGATTGCA GCAGTGAAATGG']	['Pseudohermaphroditism, female, with hypokalemia, due to glucocorticoid resistance']
NM_001018077.1(NR3C1):c.2318T>C (p.Leu773Pro)	104893912	NR3C1	[]	[]	['Glucocorticoid resistance, generalized']
NM_003122.4(SPINK1):c.2T>C (p.Met1Thr)	104893938	SPINK1	[]	[]	['Hereditary pancreatitis']
NM_000165.4(GJA1):c.52T>C (p.Ser18Pro)	104893962	GJA1	[]	['CTACYCAACTGC TGGAGGGAAGG']	['Oculodentodigital dysplasia']
NM_000416.2(IFNGR1):c.260T>C (p.Ile87Thr)	104893973	IFNGR1	['TGTA ATAYT TCTGA	['TGTAATAYTTCT GATCATGTTGG']	['Disseminated atypical mycobacterial

			TCATG TTGG']		infection', 'Mycobacterium tuberculosis, susceptibility to']
NM_000434.3(NEU1): c.718T>C (p.Trp240Arg)	104893978	NEU1	[]	['GCCTCCYGGCGC TACGGAAGTGG', 'CCTCCYGGCGCT ACGGAAGTGGG', 'CTCCYGGCGCTA CGGAAGTGGGG']	['Sialidosis, type II']
NM_153704.5(TM67):c.755T>C (p.Met252Thr)	202149403	TM67	[]	[]	['Joubert syndrome 6']
NM_000162.3(GCK):c.391T>C (p.Ser131Pro)	104894010	GCK	[]	[]	['Maturity-onset diabetes of the young, type 2']
NM_004577.3(PSPH): c.155T>C (p.Met52Thr)	104894036	PSPH	[]	[]	['Deficiency of phosphoserine phosphatase']
NM_000193.3(SHH):c.995T>C (p.Val332Ala)	104894052	SHH	[]	[]	['Single upper central incisor']
NM_000282.3(PCCA): c.491T>C (p.Ile164Thr)	202247815	PCCA	[]	[]	['Propionic acidemia']
NM_002546.3(TNFRSF11B):c.349T>C (p.Phe117Leu)	104894092	TNFRSF11B	[]	['TAGAGYTCTGCT TGAAACATAGG']	['Hyperphosphatase mia with bone disease']
NM_000532.4(PCCB): c.1556T>C (p.Leu519Pro)	202247822	PCCB	[]	[]	['Propionic acidemia']
NM_006412.3(AGPAT2):c.683T>C (p.Leu228Pro)	104894100	AGPAT2	[]	[]	['Congenital generalized lipodystrophy type 1']
NM_000238.3(KCNH2):c.2366T>C (p.Ile789Thr)	794728388	KCNH2	[]	[]	['Cardiac arrhythmia']
NM_001243133.1(NLRP3):c.1058T>C (p.Leu353Pro)	28937896	NLRP3	[]	[]	['Familial cold urticaria']
NM_021020.3(LZTS1):c.85T>C (p.Ser29Pro)	28937897	LZTS1	[]	[]	[]
NM_000102.3(CYP17A1):c.316T>C (p.Ser106Pro)	104894135	CYP17A1	[]	['CATCGCGYCCAA CAACCGTAAGG', 'ATCGCGYCCAAC AACCGTAAGGG']	['Complete combined 17-alpha- hydroxylase/17,20- lyase deficiency']
NM_000102.3(CYP17A1):c.1216T>C (p.Trp406Arg)	104894143	CYP17A1	[]	[]	['Complete combined 17-alpha- hydroxylase/17,20- lyase deficiency']
NM_000102.3(CYP17A1):c.1358T>C (p.Phe453Ser)	104894151	CYP17A1	[]	['AGCTCTYCCTCA TCATGGCCTGG']	['Combined partial 17-alpha- hydroxylase/17,20- lyase deficiency']
NM_005097.3(LGI1):c.136T>C (p.Cys46Arg)	104894166	LGI1	[]	[]	['Epilepsy, lateral temporal lobe, autosomal dominant']

NM_005097.3(LGI1):c.695T>C (p.Leu232Pro)	104894167	LGI1	[]	[]	['Epilepsy, lateral temporal lobe, autosomal dominant']
NM_000281.3(PCBD1):c.244T>C (p.Cys82Arg)	104894177	PCBD1	[]	[]	['Hyperphenylalaninemia, BH4-deficient, D']
NM_003476.4(CSRP3):c.131T>C (p.Leu44Pro)	104894205	CSRP3	[]	[]	['Familial hypertrophic cardiomyopathy 12', 'not specified']
NM_000315.2(PTH):c.67T>C (p.Ser23Pro)	104894272	PTH	[]	[]	['Hypoparathyroidism familial isolated']
NM_005055.4(RAPSN):c.848T>C (p.Leu283Pro)	104894293	RAPSN	[]	[]	['Myasthenic syndrome, congenital, associated with acetylcholine receptor deficiency', 'MYASTHENIC SYNDROME, CONGENITAL, 11, ASSOCIATED WITH ACETYLCHOLINE RECEPTOR DEFICIENCY']
NM_000518.4(HBB):c.344T>C (p.Leu115Pro)	36015961	HBB	[]	['TGTGTGCGGCC CATCACTTTGG']	['Beta thalassemia intermedia']
NM_005055.4(RAPSN):c.41T>C (p.Leu14Pro)	104894300	RAPSN	[]	[]	['MYASTHENIC SYNDROME, CONGENITAL, 11, ASSOCIATED WITH ACETYLCHOLINE RECEPTOR DEFICIENCY']
NM_000531.5(OTC):c.2T>C (p.Met1Thr)	72552295	OTC	['AGAA GAYGC TGTTT AATCT GAGG']	['AGAAGAYGCTG TTTAATCTGAGG']	['not provided']
NM_020661.2(AICDA):c.238T>C (p.Trp80Arg)	104894320	AICDA	[]	[]	['Immunodeficiency with hyper IgM type 2']
NM_020661.2(AICDA):c.317T>C (p.Leu106Pro)	104894321	AICDA	[]	[]	['Immunodeficiency with hyper IgM type 2']
NM_020661.2(AICDA):c.452T>C (p.Phe151Ser)	104894327	AICDA	[]	[]	['Immunodeficiency with hyper IgM type 2']
NM_000486.5(AQP2):c.646T>C (p.Ser216Pro)	104894329	-	[]	[]	[]
NM_020638.2(FGF23):c.287T>C (p.Met96Thr)	104894343	FGF23	[]	[]	['Tumoral calcinosis, familial, hyperphosphatemic']
NM_021044.2(DHH):c.2T>C (p.Met1Thr)	104894346	DHH	[]	[]	['46,XY gonadal dysgenesis, partial,

					with minifascicular neuropathy']
NM_000217.2(KCNA1):c.1223T>C (p.Val408Ala)	104894352	KCNA1	[]	[]	['Episodic ataxia type 1']
NM_000432.3(MYL2):c.52T>C (p.Phe18Leu)	104894370	MYL2	[]	[]	['Familial hypertrophic cardiomyopathy 10']
NM_080911.2(UNG):c.752T>C (p.Phe251Ser)	104894380	UNG	[]	[]	['Immunodeficiency with hyper IgM type 5']
NM_000192.3(TBX5):c.161T>C (p.Ile54Thr)	104894384	TBX5	[]	[]	['Holt-Oram syndrome']
NM_175929.2(FGF14):c.449T>C (p.Phe150Ser)	104894393	FGF14	[]	[]	['Spinocerebellar ataxia 27']
NM_007262.4(PARK7):c.497T>C (p.Leu166Pro)	28938172	PARK7	[]	[]	['Parkinson disease 7']
NM_004004.5(GJB2):c.229T>C (p.Trp77Arg)	104894397	GJB2	[]	[]	['Deafness, autosomal recessive 1A', 'not provided']
NM_001130089.1(KARS):c.517T>C (p.Tyr173His)	397514745	KARS	['TTCYATGATCTTGATCTTCGA GGAGA GGG']	['CTTCYATGATCTTCGAGGAGAGG', 'TTCYATGATCTTCGAGGAGAGGG']	['Deafness, autosomal recessive 89']
NM_000161.2(GCH1):c.662T>C (p.Met221Thr)	104894434	GCH1	[]	[]	['Dystonia, dopa-responsive, with or without hyperphenylalaninemia, autosomal recessive']
NM_032409.2(PINK1):c.1040T>C (p.Leu347Pro)	28940285	-	[]	[]	['Parkinson disease 6, autosomal recessive early-onset']
NM_006177.3(NRL):c.479T>C (p.Leu160Pro)	104894463	NRL	[]	[]	['Retinal degeneration, autosomal recessive, clumped pigment type']
NM_152443.2(RDH12):c.523T>C (p.Ser175Pro)	104894472	RDH12	[]	['TCCYCGGTGGCTCACCACATTGG']	['Leber congenital amaurosis 13']
NM_002435.2(MPI):c.413T>C (p.Met138Thr)	104894495	MPI	[]	[]	['Congenital disorder of glycosylation type 1B']
NM_001159702.2(FHL1):c.457T>C (p.Cys153Arg)	122458144	FHL1	[]	[]	['Myopathy, reducing body, X-linked, childhood-onset']
NM_183235.2(RAB27A):c.389T>C (p.Leu130Pro)	104894498	RAB27A	[]	[]	['Griscelli syndrome type 2']
NM_001018005.1(TPM1):c.284T>C (p.Val95Ala)	104894504	TPM1	[]	[]	['Familial hypertrophic cardiomyopathy 3', 'Cardiomyopathy']

NM_000485.2(APRT): c.329T>C (p.Leu110Pro)	104894508	APRT	[]	[]	['Adenine phosphoribosyltransferase deficiency']
NM_000303.2(PMM2) :c.131T>C (p.Val44Ala)	104894534	PMM2	[]	[]	['Carbohydrate-deficient glycoprotein syndrome type I']
NM_024006.5(VKORC1):c.134T>C (p.Val45Ala)	104894540	VKORC1	[]	[]	['Warfarin response']
NM_001614.3(ACTG1) :c.1109T>C (p.Val370Ala)	104894547	ACTG1	[]	[]	['Deafness, autosomal dominant 20']
NM_001128085.1(ASPA):c.454T>C (p.Cys152Arg)	104894548	-	[]	[]	['Spongy degeneration of central nervous system']
NM_004870.3(MPDU1):c.356T>C (p.Leu119Pro)	104894587	MPDU1	[]	['TTCCYGGTCATG CACTACAGAGG']	['Congenital disorder of glycosylation type 1F']
NM_004870.3(MPDU1):c.2T>C (p.Met1Thr)	104894588	MPDU1	[]	['AATAYGGCGGC CGAGGCGGACGG']	['Congenital disorder of glycosylation type 1F']
NM_004870.3(MPDU1):c.221T>C (p.Leu74Ser)	104894589	MPDU1	[]	[]	['Congenital disorder of glycosylation type 1F']
NM_153006.2(NAGS) :c.1289T>C (p.Leu430Pro)	104894605	-	[]	[]	['Hyperammonemia, type III']
NM_000304.3(PMP22) :c.47T>C (p.Leu16Pro)	104894617	PMP22	[]	[]	['Charcot-Marie-Tooth disease, type IA']
NM_000304.3(PMP22) :c.82T>C (p.Trp28Arg)	104894626	PMP22	[]	['TAGCAAYGGAT CGTGGGCAATGG']	['Charcot-Marie-Tooth disease, type IE']
NM_018129.3(PNPO): c.784T>C (p.Ter262Gln)	104894631	PNPO	[]	['ACCTYA ACTCTG GGACCTGCTGG']	['Pyridoxal 5'-phosphate-dependent epilepsy']
NM_173477.4(USH1G) :c.143T>C (p.Leu48Pro)	104894651	USH1G	[]	[]	['Usher syndrome, type 1G']
NM_000371.3(TTR):c.191T>C (p.Phe64Ser)	104894665	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis', 'AMYLOIDOSIS, LEPTOMENINGEAL, TRANSTHYRETIN-RELATED']
NM_024301.4(FKRP): c.899T>C (p.Val300Ala)	104894691	FKRP	[]	[]	['Limb-girdle muscular dystrophy-dystroglycanopathy, type C5']
NM_032551.4(KISS1R):c.305T>C (p.Leu102Pro)	104894703	KISS1R	[]	['GCCCTGCGYTAC CCGTGCCCGG', 'TGCGTACCCGC TGCCCGGCTGG']	[]
NM_000660.5(TGFB1) :c.673T>C	104894719	TGFB1	[]	[]	['Diaphyseal dysplasia']

(p.Cys225Arg)					
NM_000229.1(LCAT): c.524-22T>C	794726664	LCAT	[]	[]	['Fish-eye disease']
NM_003332.3(TYRO BP):c.2T>C (p.Met1Thr)	104894732	TYROBP	[]	[]	['Polycystic lipomembranous osteodysplasia with sclerosing leukoencephalopathy']
NM_000074.2(CD40L G):c.464T>C (p.Leu155Pro)	104894769	CD40LG	[]	[]	['Immunodeficiency with hyper IgM type 1']
NM_000495.4(COL4A 5):c.438+2T>C	281874738	COL4A5	['TCCAGYAAGTTATAAAATTTGG']	['CTCCAGYAAGTTATAAAATTTGG', 'TCCAGYAAGTTATAAAATTTGGG']	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A 5):c.4690T>C (p.Cys1564Arg)	281874745	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_178152.2(DCX):c. .373T>C (p.Tyr125His)	104894781	DCX	[]	[]	['Lissencephaly, X-linked', 'Subcortical laminar heterotopia, X-linked']
NM_006579.2(EBP):c. 53T>C (p.Leu18Pro)	104894795	EBP	[]	[]	['MEND SYNDROME']
NM_001097642.2(GJ B1):c.397T>C (p.Trp133Arg)	104894813	GJB1	[]	[]	['X-linked hereditary motor and sensory neuropathy']
NM_001165963.1(SC N1A):c.2690T>C (p.Leu897Ser)	794726761	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_000166.5(GJB1): c.407T>C (p.Val136Ala)	104894826	GJB1	[]	['ATGYCATCAGCGTGGTGTCCGG']	['Dejerine-Sottas disease', 'X-linked hereditary motor and sensory neuropathy']
NM_001165963.1(SC N1A):c.769T>C (p.Cys257Arg)	794726771	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy', 'not provided']
NM_001165963.1(SC N1A):c.1033T>C (p.Cys345Arg)	794726782	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_001122606.1(LA MP2):c.961T>C (p.Trp321Arg)	104894859	LAMP2	[]	['CAGCTACYGGGATGCCCCCTGG', 'AGCTACYGGGATGCCCCCTGGG']	['Danon disease']
m.10237T>C	193302927	MT-ND3	[]	[]	['Leber optic atrophy']
NM_033290.3(MID1): c.884T>C (p.Leu295Pro)	104894866	MID1	[]	[]	['Opitz-Frias syndrome']
m.10663T>C	193302933	MT-ND4L	[]	[]	['Leber optic atrophy']
NM_001165963.1(SC N1A):c.4055T>C (p.Leu1352Pro)	794726821	-	['TTCYGGTTGTTCTTATTCTGG']	['TTCYGGTTTGTCTTATTCTGG']	['Severe myoclonic epilepsy in infancy']
NM_000475.4(NR0B1)	104894907	NR0B1	[]	[]	['Congenital adrenal

);c.890T>C (p.Leu297Pro)					hypoplasia, X-linked']
NM_022567.2(NYX):c.302T>C (p.Ile101Thr)	104894911	NYX	[]	[]	['Congenital stationary night blindness, type 1A']
NM_000513.2(OPN1MW):c.607T>C (p.Cys203Arg)	104894914	OPN1MW	[]	[]	['Colorblindness, partial, deutan series', 'Cone monochromatism']
NM_006517.4(SLC16A2):c.1313T>C (p.Leu438Pro)	104894931	SLC16A2	[]	['TGAGCYGGTGGGCCCAATGCAGG']	['Allan-Herndon-Dudley syndrome']
NM_000330.3(RS1):c.38T>C (p.Leu13Pro)	104894935	RS1	[]	['TTACTTCYCTTTGGCTATGAAGG']	['Juvenile retinoschisis', 'not provided']
NM_000116.4(TAZ):c.352T>C (p.Cys118Arg)	104894937	TAZ	['AAGYGTGTGCCTGTGTGCCGAGG']	['AAGYGTGTGCCTGTGTGCCGAGG']	['3-Methylglutaconic aciduria type 2']
NM_006517.4(SLC16A2):c.1481T>C (p.Leu494Pro)	104894938	SLC16A2	[]	[]	['Allan-Herndon-Dudley syndrome']
NM_001109878.1(TBX22):c.641T>C (p.Leu214Pro)	104894946	TBX22	[]	[]	['Cleft palate with ankyloglossia']
NM_001011658.3(TRAPPC2):c.248T>C (p.Phe83Ser)	104894948	-	[]	[]	['Spondyloepiphyseal dysplasia tarda']
NM_003140.2(SRY):c.326T>C (p.Phe109Ser)	104894956	SRY	[]	[]	['46,XY sex reversal, type 1']
NM_003140.2(SRY):c.203T>C (p.Ile68Thr)	104894968	SRY	[]	[]	['46,XY sex reversal, type 1']
NM_201269.2(ZNF644):c.1759A>G (p.Ile587Val)	146936371	ZNF644	[]	[]	['Myopia 21, autosomal dominant']
NM_001004434.2(SLC30A2):c.161A>G (p.His54Arg)	587776926	SLC30A2	[]	[]	['Reduced zinc in breast milk']
NM_000492.3(CFTR):c.3469-20T>C	373002889	CFTR	[]	[]	['Cystic fibrosis']
NM_001848.2(COL6A1):c.957+2T>C	794727060	COL6A1	['ACAAAGGYGAGC GCGTG GGCTG CTGG', 'CAAGGYGAGC GTGGG CTGCT GGG']	['ACAAGGYGAGC GTGGGCTGCTGG', 'CAAGGYGAGCGT GGGCTGCTGGG']	['Ullrich congenital muscular dystrophy', 'Bethlem myopathy']
m.4336T>C	41456348	MT-TQ	[]	[]	[]
NM_001065.3(TNFRSF1A):c.175T>C (p.Cys59Arg)	104895217	TNFRSF1A	[]	['TGCYGTACCAAGTGCCACAAAGG']	['TNF receptor-associated periodic fever syndrome (TRAPS)']
NM_003072.3(SMARCA4):c.3032T>C	281875229	SMARCA4	[]	[]	['Mental retardation, autosomal dominant']

(p.Met1011Thr)					16', 'not provided']
NM_019885.3(CYP26B1):c.436T>C (p.Ser146Pro)	281875232	CYP26B1	[]	[]	['Radiohumeral fusions with other skeletal and craniofacial anomalies', 'not provided']
NM_000182.4(HADHA):c.180+3A>G	781222705	HADHA	[]	[]	['Mitochondrial trifunctional protein deficiency', 'Long-chain 3-hydroxyacyl-CoA dehydrogenase deficiency', 'not provided']
NM_000208.2(INSR):c.1124-2A>G	587776819	INSR	[]	[]	['Pineal hyperplasia AND diabetes mellitus syndrome']
NM_006329.3(FBLN5):c.506T>C (p.Ile169Thr)	28939072	FBLN5	['GACAYTGATGAATGTCGCTATGG']	['GACAYTGATGAATGTCGCTATGG']	['Age-related macular degeneration 3']
NM_000431.3(MVK):c.803T>C (p.Ile268Thr)	104895304	MVK	['CTCAAYAGATGCTGCCCTGG']	['CTCAAYAGATGCTGCCCTGG']	['Hyperimmunoglobulin D with periodic fever', 'Mevalonic aciduria']
NM_024960.4(PANK2):c.437T>C (p.Met146Thr)	28939088	PANK2	[]	[]	['Hypoprebetalipoproteinemia, acanthocytosis, retinitis pigmentosa, and pallidal degeneration']
NM_005359.5(SMAD4):c.1499T>C (p.Ile500Thr)	281875321	SMAD4	[]	[]	['Myhre syndrome', 'not provided']
NM_003793.3(CTSF):c.692A>G (p.Tyr231Cys)	143889283	CTSF	[]	['CTCCAYACTGAGCTGTGCCACGG']	['Ceroid lipofuscinosis, neuronal, 13']
NM_001159702.2(FHL1):c.310T>C (p.Cys104Arg)	122459147	FHL1	[]	['GGGGYGCTTCAAGGCCATTGTGG']	['Myopathy, reducing body, X-linked, childhood-onset']
NM_001159702.2(FHL1):c.625T>C (p.Cys209Arg)	122459149	FHL1	[]	[]	['Emery-dreifuss muscular dystrophy 6']
NM_006214.3(PHYH):c.135-2A>G	201578674	PHYH	[]	[]	['Refsum disease, adult, 1']
NM_006329.3(FBLN5):c.679T>C (p.Ser227Pro)	28939370	FBLN5	[]	[]	['Autosomal recessive cutis laxa type IA']
NM_004329.2(BMPRI1A):c.1409T>C (p.Met470Thr)	199476089	BMPRI1A	[]	[]	['Juvenile polyposis syndrome']
NM_005154.4(USP8):c.2152T>C (p.Ser718Pro)	672601307	USP8	[]	[]	['Pituitary dependent hypercortisolism']
NM_020184.3(CNNM4)	74552543	CNNM4	[]	['AAGCTCCYGGA']	['Cone-rod

4):c.971T>C (p.Leu324Pro)				CTTTTTTCTGGG']	dystrophy amelogenesis imperfecta']
NM_000734.3(CD247) :c.2T>C (p.Met1Thr)	672601318	CD247	[]	[]	['Immunodeficiency due to defect in cd3- zeta']
NM_016042.3(EXOS C3):c.712T>C (p.Trp238Arg)	672601332	EXOSC3	[]	[]	['Pontocerebellar hypoplasia, type 1b']
NC_012920.1.m.1448 4T>C	199476104	MT-ND6	[]	[]	['Leber optic atrophy', 'Leigh disease']
m.10158T>C	199476117	MT-ND3	[]	['AAAYCCACCCCT TACGAGTGCGG']	['Leigh disease', 'Leigh syndrome due to mitochondrial complex I deficiency', 'Mitochondrial complex I deficiency']
NM_020451.2(SEPN1) :c.872+2T>C	794727808	SEPN1	[]	['TTCCGGYGAGTG GGCCACACTGG']	['Congenital myopathy with fiber type disproportion', 'Eichsfeld type congenital muscular dystrophy']
NM_005022.3(PFN1): c.350A>G (p.Glu117Gly)	140547520	PFN1	[]	['CACCTYCTTTGC CCATCAGCAGG']	['Amyotrophic lateral sclerosis 18']
NM_032551.4(KISS1 R):c.443T>C (p.Leu148Ser)	28939719	KISS1R	[]	[]	[]
NM_000084.4(CLCN5) :c.674T>C (p.Leu225Pro)	273585645	CLCN5	[]	[]	['Dent disease 1']
NM_000030.2(AGXT) :c.605T>A (p.Ile202Asn)	536352238	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_000060.3(BTD):c .212T>C (p.Leu71Pro)	397514333	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c .248T>C (p.Leu83Ser)	397514347	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c .445T>C (p.Phe149Leu)	397514359	BTD	[]	['TCACCGCYTCAA TGACACAGAGG']	['Biotinidase deficiency']
NM_000060.3(BTD):c .743T>C (p.Ile248Thr)	397514382	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c .764T>C (p.Ile255Thr)	397514384	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c .833T>C (p.Leu278Pro)	397514389	BTD	[]	[]	['Biotinidase deficiency']
NM_000061.2(BTK):c .2T>C (p.Met1Thr)	128620186	BTK	['AGCT AYGGC CGCAG TGATT CTGG']	['AGCTAYGGCCG CAGTGATTCTGG']	['X-linked agammaglobulinemi a']
m.15572T>C	207459996	MT-CYB	[]	[]	['Familial colorectal cancer']

NM_000060.3(BTD):c.1096T>C (p.Ser366Pro)	397514399	BTD	[]	[]	['Biotinidase deficiency']
m.15197T>C	207460001	MT-CYB	[]	['CTAYCCGCCATC CCATACATTGG']	['Exercise intolerance']
m.14849T>C	207460004	MT-CYB	[]	[]	[]
NM_000060.3(BTD):c.1214T>C (p.Leu405Pro)	397514406	BTD	[]	['TTCACCCYGGTC CCTGTCTGGGG']	['Biotinidase deficiency']
NM_000060.3(BTD):c.1252T>C (p.Cys418Arg)	397514408	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c.1267T>C (p.Cys423Arg)	397514412	BTD	[]	[]	['Biotinidase deficiency']
NM_001128177.1(THRB):c.1336T>C (p.Cys446Arg)	121918703	THRB	[]	[]	['Thyroid hormone resistance, generalized, autosomal dominant']
NM_000060.3(BTD):c.1459T>C (p.Trp487Arg)	397514422	BTD	[]	[]	['Biotinidase deficiency']
NM_198056.2(SCN5A):c.3963+2T>C	397514447	SCN5A	[]	[]	['Progressive familial heart block type 1A']
NM_020461.3(TUBGCP6):c.2546A>G (p.Glu849Gly)	368449236	TUBGCP6	[]	[]	['Microcephaly with chorioretinopathy, autosomal recessive']
NM_006225.3(PLCD1):c.562T>C (p.Cys188Arg)	397514471	PLCD1	[]	[]	['Leukonychia totalis']
NM_001161581.1(POC1A):c.398T>C (p.Leu133Pro)	397514488	POC1A	['AGCYGTGGG ACAAGAGCAG CCGG']	['AGCYGTGGGAC AAGAGCAGCCGG']	['Short stature, onychodysplasia, facial dysmorphism, and hypotrichosis']
NM_005340.6(HINT1):c.250T>C (p.Cys84Arg)	397514489	HINT1	['CAAGAAAYG TGCTGCTGAT CTGG', 'AAGAAAYGTG CTGCTGATCT GGG']	['CAAGAAAYGTG CTGCTGATCTGG', 'AAGAAAYGTGCT GCTGATCTGGG']	['Gamstorp-Wohlfart syndrome']
NM_000051.3(ATM):c.2T>C (p.Met1Thr)	786203606	ATM	[]	[]	['Ataxia-telangiectasia syndrome', 'Hereditary cancer-predisposing syndrome']
NM_004281.3(BAG3):c.1385T>C (p.Leu462Pro)	397514507	BAG3	[]	[]	['Dilated cardiomyopathy 1HH']
NM_183075.2(CYP2U1):c.784T>C	397514515	CYP2U1	[]	[]	['Spastic paraplegia 56, autosomal']

(p.Cys262Arg)					recessive']
NM_006177.3(NRL):c.287T>C (p.Met96Thr)	397514516	NRL	[]	['GAGGCCAYGGA GCTGCTGCAGGG']	['Retinitis pigmentosa 27']
NM_000344.3(SMN1):c.388T>C (p.Tyr130His)	397514518	SMN1	['CACT GGAYA TGGAA ATAGA GAGG']	['CACTGGAYATG GAAATAGAGAGG']	['Kugelberg-Welander disease']
NM_152692.4(C1GALT1C1):c.577T>C (p.Ser193Pro)	397514537	C1GALT1C1	[]	[]	['Polyagglutinable erythrocyte syndrome']
NM_024531.4(SLC52A2):c.368T>C (p.Leu123Pro)	397514538	SLC52A2	[]	[]	['Brown-Vialetto-Van Laere syndrome 2']
NM_000138.4(FBN1):c.5746T>C (p.Cys1916Arg)	794728238	FBN1	[]	[]	['Thoracic aortic aneurysms and aortic dissections']
NM_000138.4(FBN1):c.6274T>C (p.Trp2092Arg)	794728246	FBN1	[]	[]	['Thoracic aortic aneurysms and aortic dissections']
NM_017802.3(DNAAF5):c.2384T>C (p.Leu795Pro)	397514561	DNAAF5	[]	[]	['Ciliary dyskinesia, primary, 18']
NM_206933.2(USH2A):c.12295-2A>G	151148854	USH2A	[]	[]	['Usher syndrome, type 2A']
NM_000531.5(OTC):c.134T>C (p.Leu45Pro)	72554312	OTC	[]	['CTCACTCYAAAA AACTTTACCGG']	['Ornithine carbamoyltransferase deficiency', 'not provided']
NM_178012.4(TUBB2B):c.350T>C (p.Leu117Pro)	397514569	TUBB2B	[]	['GGTCCYGGATGT GGTGAGGAAGG']	['Polymicrogyria, asymmetric']
NM_000431.3(MVK):c.764T>C (p.Leu255Pro)	397514570	MVK	[]	[]	['Porokeratosis, disseminated superficial actinic 1']
NM_000431.3(MVK):c.122T>C (p.Leu41Pro)	397514571	MVK	[]	['CGGCYTCAACCC CACAGCAATGG', 'GGCYTCAACCCC ACAGCAATGGG']	['Porokeratosis, disseminated superficial actinic 1']
NM_000531.5(OTC):c.167T>C (p.Met56Thr)	72554320	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.188T>C (p.Leu63Pro)	72554324	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.227T>C (p.Leu76Ser)	72554328	OTC	[]	[]	['not provided']
NM_004055.4(CAPN5):c.731T>C (p.Leu244Pro)	397514602	CAPN5	[]	[]	['Vitreoretinopathy, neovascular inflammatory']
NM_133497.3(KCNV2):c.491T>C (p.Phe164Ser)	397514604	KCNV2	[]	[]	['Retinal cone dystrophy 3B']
NM_006567.3(FARS2):c.986T>C (p.Ile329Thr)	397514611	FARS2	[]	[]	['Combined oxidative phosphorylation deficiency 14']
NM_018344.5(SLC29A3):c.607T>C (p.Ser203Pro)	397514626	SLC29A3	['ACTG ATAYC AGGTG AGAGC	['ACTGATAYCAG GTGAGAGCCAGG', 'CTGATAYCAGGT	['Histiocytosis-lymphadenopathy plus syndrome']

			CAGG']	GAGAGCCAGGG']	
NM_000108.4(DLD):c.140T>C (p.Ile47Thr)	397514651	DLD	['ACAGTTAYAGGTTCTGGTCCTGG', 'GTTAYAGGTTCTGGTCCTGGAGG']	['ACAGTTAYAGGTTCTGGTCCTGG', 'GTTAYAGGTTCTGGTCCTGGAGG']	['Maple syrup urine disease, type 3']
NM_020632.2(ATP6V0A4):c.1739T>C (p.Met580Thr)	3807153	ATP6V0A4	[]	[]	['Renal tubular acidosis, distal, autosomal recessive']
NM_000238.3(KCNH2):c.1945+6T>C	794728380	KCNH2	['CTGTGAGYGTGCCAGGGGCGGG', 'TGAGYGTGCCAGGGGCGGGCGG']	['CTGTGAGYGTGCCAGGGGCGGG', 'TGAGYGTGCCAGGGGCGGGCGG']	['Cardiac arrhythmia']
NM_001033053.2(NLRP1):c.230T>C (p.Met77Thr)	397514692	NLRP1	[]	[]	['Corneal intraepithelial dyskeratosis and ectodermal dysplasia']
NM_000238.3(KCNH2):c.2396T>C (p.Leu799Pro)	794728390	KCNH2	[]	['GCCATCCYGGGTATGGGGTGGGG', 'CCATCCYGGGTATGGGGTGGGG', 'CATCCYGGGTATGGGGTGGGG']	['Cardiac arrhythmia']
NM_014845.5(FIG4):c.524T>C (p.Leu175Pro)	397514707	FIG4	[]	[]	['Yunis Varon syndrome']
NM_001199107.1(TBC1D24):c.686T>C (p.Phe229Ser)	397514713	TBC1D24	[]	['GGTCTYTGACGTTTCCTGGTGG']	['Early infantile epileptic encephalopathy 16']
NM_080605.3(B3GALT6):c.193A>G (p.Ser65Gly)	397514719	B3GALT6	[]	['CGCYGGCCACAGCACTGCCAGG']	['Spondyloepimetaphyseal dysplasia with joint laxity']
NM_004183.3(BEST1):c.253T>C (p.Tyr85His)	28940274	BEST1	[]	[]	['Viteliform dystrophy', 'not provided']
NM_005689.2(ABCB6):c.1067T>C (p.Leu356Pro)	397514756	ABCB6	[]	[]	['Dyschromatosis universalis hereditaria 3']
NM_000551.3(VHL):c.488T>C (p.Leu163Pro)	28940297	VHL	[]	[]	[]
NM_000218.2(KCNQ1):c.1025T>C (p.Leu342Pro)	794728522	KCNQ1	[]	[]	['Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.1251+2T>C	794728528	KCNQ1	[]	[]	['Cardiac arrhythmia']

NM_000498.3(CYP11B2):c.1382T>C (p.Leu461Pro)	72554627	-	[]	[]	['Corticosterone methyl oxidase type 1 deficiency']
NM_130799.2(MEN1):c.547T>C (p.Trp183Arg)	794728649	MEN1	[]	[]	['not provided']
NM_213653.3(HFE2):c.238T>C (p.Cys80Arg)	28940586	HFE2	[]	[]	['Hemochromatosis type 2A']
NM_198056.2(SCN5A):c.4299+6T>C	794728934	SCN5A	[]	[]	['not provided']
NM_000548.3(TSC2):c.1946+2T>C	397515247	TSC2	[]	[]	['Tuberous sclerosis syndrome']
NM_000256.3(MYBP3):c.3796T>C (p.Cys1266Arg)	730880608	MYBP3	[]	['GAGYGCCGCCTGGAGGTGCGAGG']	['Cardiomyopathy']
NM_016381.5(TREX1):c.530T>C (p.Val177Ala)	79993407	TREX1	[]	[]	['Aicardi Goutieres syndrome 1']
NM_001382.3(DPAGT1):c.503T>C (p.Leu168Pro)	397515329	DPAGT1	[]	['AATCCYGTACTA TGTCTACATGG', 'ATCCYGTACTAT GTCTACATGGG', 'TCCYGTACTATG TCTACATGGGG']	['Congenital disorder of glycosylation type 1J']
NM_000372.4(TYR):c.265T>C (p.Cys89Arg)	28940877	TYR	[]	[]	['Tyrosinase-negative oculocutaneous albinism', 'not provided']
NM_000375.2(UROS):c.-26-177T>C	397515348	UROS	[]	[]	['Congenital erythropoietic porphyria']
NM_015102.4(NPHP4):c.2972T>C (p.Phe991Ser)	28940891	NPHP4	[]	[]	['Nephronophthisis 4']
NM_020822.2(KCNT1):c.2386T>C (p.Tyr796His)	397515406	KCNT1	[]	[]	['Epilepsy, nocturnal frontal lobe, 5']
NM_000061.2(BTK):c.1516T>C (p.Cys506Arg)	128621200	BTK	[]	[]	['X-linked agammaglobulinemia']
NM_006383.3(CIB2):c.272T>C (p.Phe91Ser)	397515411	CIB2	[]	[]	['Deafness, autosomal recessive 48']
NM_000061.2(BTK):c.1741T>C (p.Trp581Arg)	128621205	BTK	['ACATTYGGGC TTTTG GTAAG TGG']	['ACATTYGGGCTT TTGGTAAGTGG']	['X-linked agammaglobulinemia']
NM_018127.6(ELAC2):c.460T>C (p.Phe154Leu)	397515465	ELAC2	[]	['ATAYTTTCTGGT CCATTGAAAGG']	['Combined oxidative phosphorylation deficiency 17']
NM_199355.2(ADAMTS18):c.605T>C (p.Leu202Pro)	397515468	ADAMTS18	[]	[]	['Microcornea, myopic chorioretinal atrophy, and telecanthus']
NM_023110.2(FGFR1)	397515481	FGFR1	[]	[]	['Hartsfield']

):c.494T>C (p.Leu165Ser)					syndrome']
NM_001059.2(TACR3):c.766T>C (p.Tyr256His)	397515483	TACR3	[]	[]	['Hypogonadotropic hypogonadism 11 with or without anosmia']
m.14325T>C	397515505	MT-ND6	[]	[]	['Leber optic atrophy']
NM_004333.4(BRAF):c.1783T>C (p.Phe595Leu)	794729219	BRAF	[]	[]	['Cardiofaciocutaneous syndrome']
NM_000370.3(TTPA):c.548T>C (p.Leu183Pro)	397515525	TTPA	[]	[]	['Ataxia with vitamin E deficiency']
NM_000375.2(UROS):c.139T>C (p.Ser47Pro)	397515527	UROS	[]	[]	['Congenital erythropoietic porphyria']
NM_001006657.1(WDR35):c.1592T>C (p.Leu531Pro)	397515533	WDR35	[]	[]	['Cranioectodermal dysplasia 2']
NM_004595.4(SMS):c.449T>C (p.Ile150Thr)	397515552	SMS	[]	[]	['Snyder Robinson syndrome']
NM_005211.3(CSF1R):c.2483T>C (p.Phe828Ser)	397515557	CSF1R	[]	['CATCTYTGACTGTGTCTACACGG']	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_000026.2(ADSL):c.1339T>C (p.Ser447Pro)	777821034	ADSL	[]	[]	['not provided']
NM_194248.2(OTOF):c.3413T>C (p.Leu1138Pro)	397515599	OTOF	[]	['AGGTGCGTCTGGGCCTACGG', 'GGTGCYGTCTGGGCCTACGGG']	['Deafness, autosomal recessive 9']
NM_002608.2(PDGFB):c.356T>C (p.Leu119Pro)	397515632	PDGFB	[]	[]	['Idiopathic basal ganglia calcification 5']
NM_000404.2(GLB1):c.152T>C (p.Ile51Thr)	72555390	GLB1	[]	[]	['Gangliosidosis GM1 type 3']
NM_000116.4(TAZ):c.310T>C (p.Phe104Leu)	397515741	TAZ	[]	[]	['3-Methylglutaconic aciduria type 2']
NM_000138.4(FBN1):c.2341T>C (p.Cys781Arg)	397515766	FBN1	[]	['GGACAAYGTAGAAATACTCCTGG']	['Marfan syndrome']
NM_000138.4(FBN1):c.4222T>C (p.Cys1408Arg)	397515802	FBN1	[]	[]	['Marfan syndrome']
NM_000112.3(SLC26A2):c.-26+2T>C	386833492	SLC26A2	['GAGAGGYGA GAAGA GGGAA GCGG']	['GAGAGGYGAGA AGAGGGAAGCGG']	['Diastrophic dysplasia']
NM_000256.3(MYBPC3):c.1351+2T>C	397515897	MYBPC3	['AAAGGYGGG CCTGG GACCTGAGG']	['AAAGGYGGGCC TGGGACCTGAGG']	['Familial hypertrophic cardiomyopathy 4', 'Cardiomyopathy']
NM_000045.3(ARG1):c.32T>C (p.Ile11Thr)	28941474	ARG1	[]	[]	['Arginase deficiency']
NM_004820.3(CYP7B)	587777222	CYP7B1	[]	[]	['Spastic paraplegia',

1):c.889A>G (p.Thr297Ala)					'Spastic paraplegia 5A']
NM_017909.3(RMND1):c.713A>G (p.Asn238Ser)	144972972	RMND1	[]	[]	['Combined oxidative phosphorylation deficiency 11']
NM_000314.6(PTEN):c.545T>C (p.Leu182Ser)	794729664	PTEN	[]	[]	['Macrocephaly/autism syndrome']
NM_000256.3(MYBPC3):c.821+2T>C	397516076	MYBPC3	['CACG YGAGT GGCCA TCCTC AGGG']	['GCACGYGAGTG GCCATCCTCAGG', 'CACGYGAGTGGC CATCCTCAGGG']	['Familial hypertrophic cardiomyopathy 4', 'not specified']
NM_000257.3(MYH7):c.1370T>C (p.Ile457Thr)	397516103	MYH7	[]	[]	['Cardiomyopathy', 'not specified']
NM_000257.3(MYH7):c.2093T>C (p.Val698Ala)	397516130	MYH7	[]	[]	['Familial hypertrophic cardiomyopathy 1', 'not specified']
NM_000257.3(MYH7):c.2546T>C (p.Met849Thr)	397516156	MYH7	['GGAG AYGGC CTCCA TGAAG GAGG']	['GGAGAYGGCCT CCATGAAGGAGG']	['Primary familial hypertrophic cardiomyopathy', 'Cardiomyopathy']
NM_000271.4(NPC1):c.1133T>C (p.Val378Ala)	120074134	NPC1	[]	[]	['Niemann-Pick disease type C1']
NM_000520.4(HEXA):c.538T>C (p.Tyr180His)	28941771	HEXA	[]	[]	[]
NM_024426.4(WT1):c.1351T>C (p.Phe451Leu)	28941777	WT1	[]	[]	['Diffuse mesangial sclerosis']
NM_024426.4(WT1):c.1378T>C (p.Phe460Leu)	28941779	WT1	[]	[]	['Frasier syndrome']
NM_000257.3(MYH7):c.602T>C (p.Ile201Thr)	397516258	MYH7	[]	[]	['Dilated cardiomyopathy 1S', 'Cardiomyopathy']
NM_000257.3(MYH7):c.788T>C (p.Ile263Thr)	397516269	MYH7	[]	[]	['Primary familial hypertrophic cardiomyopathy', 'Familial hypertrophic cardiomyopathy 1', 'Cardiomyopathy']
NM_001429.3(EP300):c.3573T>A (p.Tyr1191Ter)	565779970	EP300	[]	['CTTAYTACAGTT ACCAGAACAGG']	['Rubinstein-Taybi syndrome 2']
NM_080605.3(B3GALT6):c.1A>G (p.Met1Val)	786200938	B3GALT6	[]	['AGCTTCAYGGCG CCCGCGCCGGG', 'TCAYGGCGCCCG CGCCGGGCCGG']	['Spondyloepimetaphyseal dysplasia with joint laxity']
NM_032551.4(KISS1R):c.937T>C (p.Tyr313His)	587777844	KISS1R	[]	[]	[]
NM_000257.3(MYH7)	369437262	MYH7	[]	[]	['Familial

:c.5326A>G (p.Ser1776Gly)					hypertrophic cardiomyopathy 1', 'Cardiomyopathy', 'not specified']
NM_000441.1(SLC26A4):c.164+2T>C	397516420	SLC26A4	[]	[]	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_000441.1(SLC26A4):c.765+2T>C	397516432	SLC26A4	[]	[]	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_000551.3(VHL):c.497T>C (p.Val166Ala)	397516445	VHL	[]	[]	['Von Hippel-Lindau syndrome', 'Hereditary cancer-predisposing syndrome']
NM_000256.3(MYBP3):c.709T>C (p.Tyr237His)	730880624	MYBPC3	[]	[]	['Cardiomyopathy']
NM_000531.5(OTC):c.392T>C (p.Leu131Ser)	72556252	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.394T>C (p.Ser132Pro)	72556253	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.416T>C (p.Leu139Ser)	72556259	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.476T>C (p.Ile159Thr)	72556269	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.490T>C (p.Ser164Pro)	72556273	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.526T>C (p.Tyr176His)	72556282	OTC	['GGCT GATYA CCTCA CGCTC CAGG']	['GGCTGATYACCT CACGCTCCAGG', 'GATYACCTCACG CTCCAGGTTGG']	['not provided']
NM_000531.5(OTC):c.536T>C (p.Leu179Pro)	72556286	OTC	[]	[]	['not provided']
NM_000229.1(LCAT):c.698T>C (p.Leu233Pro)	28942087	LCAT	[]	['ATCTCTCYTGGG GCTCCCTGGGG', 'TCTCYTGGGGCT CCCTGGGGTGG']	['Norum disease']
NM_174936.3(PCSK9):c.646T>C (p.Phe216Leu)	28942112	PCSK9	[]	[]	['Hypercholesterolemia, autosomal dominant, 3']
NM_004572.3(PKP2):c.2386T>C (p.Cys796Arg)	794729098	PKP2	[]	[]	['not provided']
NM_000061.2(BTK):c.1223T>C (p.Leu408Pro)	128621198	BTK	['AGCY GGGGA CTGGA CAATT TGGG']	['GAGCYGGGGAC TGGACAATTTGG', 'AGCYGGGGACTG GACAATTTGGG']	['X-linked agammaglobulinemia']
NM_000061.2(BTK):c.1625T>C (p.Leu542Pro)	128621203	BTK	[]	['TCGGCCYGTCCA GGTGAGTGTGG']	['X-linked agammaglobulinemia with growth hormone']

					deficiency']
NM_006383.3(CIB2):c.368T>C (p.Ile123Thr)	397515412	CIB2	[]	['CTTCAYCTGCAA GGAGGACCTGG']	['Deafness, autosomal recessive 48']
NM_001943.3(DSG2):c.523+2T>C	397516709	DSG2	[]	[]	['Arrhythmogenic right ventricular cardiomyopathy, type 10', 'Cardiomyopathy']
NM_032575.2(GLIS2):c.523T>C (p.Cys175Arg)	587777353	GLIS2	[]	[]	['Nephronophthisis 7']
NM_000492.3(CFTR):c.3230T>C (p.Leu1077Pro)	139304906	CFTR	[]	[]	['Cystic fibrosis']
NM_000492.3(CFTR):c.1853T>C (p.Ile618Thr)	139468767	CFTR	[]	[]	['Cystic fibrosis', 'not provided']
NM_002755.3(MAP2K1):c.388T>C (p.Tyr130His)	397516793	MAP2K1	[]	[]	['Cardiofaciocutaneous syndrome 3']
NM_000525.3(KCNJ11):c.755T>C (p.Val252Ala)	193929352	KCNJ11	[]	[]	['Permanent neonatal diabetes mellitus']
NM_000352.4(ABCC8):c.404T>C (p.Leu135Pro)	193929364	ABCC8	[]	['AAGCYGCTAATT GGTAGGTGAGG']	['Permanent neonatal diabetes mellitus']
NM_000071.2(CBS):c.833T>C (p.Ile278Thr)	5742905	CBS	['ATCAYTGGGGTGGATCCCGAAGG', 'TCAYTGGGGTGGATCCCGAAGGG']	['ATCAYTGGGGTGGATCCCGAAGG', 'TCAYTGGGGTGGATCCCGAAGGG']	['Homocystinuria due to CBS deficiency', 'Homocystinuria, pyridoxine-responsive', 'not provided']
NM_001038.5(SCNN1A):c.1477T>C (p.Trp493Arg)	5742912	SCNN1A	[]	[]	['Bronchiectasis with or without elevated sweat chloride 2', 'not specified']
NM_000030.2(AGXT):c.2T>C (p.Met1Thr)	138584408	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_005633.3(SOS1):c.1649T>C (p.Leu550Pro)	397517153	SOS1	[]	[]	['Noonan syndrome 4', 'Rasopathy']
NM_014714.3(IFT140):c.4078T>C (p.Cys1360Arg)	431905520	IFT140	['GCAGYGTGA GCTGC TCCTG GAGG']	['CAAGCAGYGTGAGCTGCTCCTGG', 'GCAGYGTGAGCTGCTCCTGGAGG']	['Renal dysplasia, retinal pigmentary dystrophy, cerebellar ataxia and skeletal dysplasia']
NM_022168.3(IFIH1):c.1009A>G (p.Arg337Gly)	587777447	IFIH1	[]	[]	['Aicardi-goutieres syndrome 7']
NG_012123.1:g.2493A>G	1024611	CCL2	[]	[]	['Coronary artery disease, modifier of', 'Coronary artery disease,']

					development of, in hiv', 'Mycobacterium tuberculosis, susceptibility to']
m.3394T>C	41460449	MT-ND1	['GGCY ATATA CAACT ACGCA AAGG']	['GGCYATATACA ACTACGCAAAGG']	['Leber optic atrophy']
NM_001127328.2(ACADM):c.997A>G (p.Lys333Glu)	77931234	ACADM	[]	[]	['Medium-chain acyl-coenzyme A dehydrogenase deficiency', 'not provided']
NM_005859.4(PURA):c.299T>C (p.Leu100Pro)	587782995	PURA	[]	[]	['Neonatal hypotonia', 'Intellectual disability', 'Seizures', 'Delayed speech and language development', 'Global developmental delay', 'Mental retardation, autosomal dominant 31']
NM_000368.4(TSC1):c.539T>C (p.Leu180Pro)	118203396	TSC1	[]	[]	['Tuberous sclerosis syndrome', 'Tuberous sclerosis 1']
NM_000256.3(MYBPC3):c.1696T>C (p.Cys566Arg)	730880695	MYBPC3	[]	[]	['Cardiomyopathy']
m.7275T>C	267606884	MT-CO1	[]	[]	['Familial colorectal cancer']
NM_000257.3(MYH7):c.1400T>C (p.Ile467Thr)	730880872	MYH7	[]	['TCGAGAYCTTCG ATGTGAGTTGG', 'CGAGAYCTTCGA TGTGAGTTGGG']	['Cardiomyopathy']
NM_002977.3(SCN9A):c.647T>C (p.Phe216Ser)	80356469	SCN9A	[]	[]	['Primary erythromelgia']
NM_002977.3(SCN9A):c.2543T>C (p.Ile848Thr)	80356474	-	[]	['AAGATCAYTGGT AACTCAGTAGG', 'AGATCAYTGGTA ACTCAGTAGGG', 'GATCAYTGGTAA CTCAGTAGGGG']	['Primary erythromelgia']
NM_001164277.1(SLC37A4):c.352T>C (p.Trp118Arg)	80356489	SLC37A4	[]	['GGGCYGGCCCC CATGTGGGAAGG']	['Glucose-6-phosphate transport defect', 'not provided']
NM_001457.3(FLNB):c.4804T>C (p.Ser1602Pro)	80356501	FLNB	[]	[]	[]
NM_152296.4(ATP1A3):c.2338T>C (p.Phe780Leu)	80356536	ATP1A3	[]	['GCCCYTCCTGCT GTTCATCATGG']	['Dystonia 12']

NM_206933.2(USH2A):c.5857+2T>C	397518022	-	[]	[]	['Usher syndrome, type 2A']
NM_194248.2(OTOF):c.1544T>C (p.Ile515Thr)	80356586	OTOF	[]	[]	['Deafness, autosomal recessive 9', 'Auditory neuropathy, autosomal recessive, 1']
NM_000335.4(SCN5A):c.3745T>C (p.Phe1249Leu)	45589741	SCN5A	[]	[]	['Acquired long QT syndrome']
NM_194248.2(OTOF):c.3032T>C (p.Leu1011Pro)	80356596	OTOF	[]	['GATGCYGGTGTTCGACAACCTGG']	['Deafness, autosomal recessive 9', 'Auditory neuropathy, autosomal recessive, 1']
NM_000525.3(KCNJ11):c.124T>C (p.Cys42Arg)	80356610	KCNJ11	[]	[]	['Permanent neonatal diabetes mellitus', 'Transient neonatal diabetes mellitus 3', 'MATURITY-ONSET DIABETES OF THE YOUNG, TYPE 13']
NM_000257.3(MYH7):c.2723T>C (p.Leu908Pro)	730880900	MYH7	[]	[]	['Cardiomyopathy']
NM_152296.4(ATP1A3):c.1112T>C (p.Leu371Pro)	606231433	ATP1A3	[]	[]	['Alternating hemiplegia of childhood 2']
NM_000083.2(CLCN1):c.857T>C (p.Val286Ala)	80356689	CLCN1	[]	['AGGAGYGCTATTAGCATCGAGG']	['Myotonia congenita']
NM_000083.2(CLCN1):c.920T>C (p.Phe307Ser)	80356701	CLCN1	[]	[]	['Myotonia congenita']
NM_007375.3(TARDBP):c.*83T>C	80356744	TARDBP	[]	[]	['Amyotrophic lateral sclerosis type 10']
NM_152296.4(ATP1A3):c.1250T>C (p.Leu417Pro)	606231449	ATP1A3	[]	[]	['Dystonia 12']
NM_001876.3(CPT1A):c.1451T>C (p.Leu484Pro)	80356793	CPT1A	[]	[]	['Carnitine palmitoyltransferase I deficiency']
NM_000088.3(COL1A1):c.4391T>C (p.Leu1464Pro)	72656353	COL1A1	[]	[]	['Osteogenesis imperfecta type III']
NM_000089.3(COL1A2):c.279+2T>C	72656357	COL1A2	[]	[]	['Ehlers-Danlos syndrome, type 7B']
NM_015046.5(SETX):c.1807A>G (p.Asn603Asp)	116205032	SETX	[]	[]	['Spinocerebellar ataxia autosomal recessive 1']
m.4409T>C	118203884	MT-TM	[]	['AGGYCAGCTAAATAAGCTATCGG']	['Mitochondrial myopathy']
m.5874T>C	118203891	MT-TY	[]	[]	[]
NM_000130.4(F5):c.1160T>C (p.Ile387Thr)	118203911	F5	[]	[]	['Thrombophilia due to activated protein']

					C resistance']
NM_173596.2(SLC39A5):c.911T>C (p.Met304Thr)	587777625	SLC39A5	[]	['AGAACAYGCTG GGGCTTTTGC GG']	['Myopia 24, autosomal dominant']
NM_024120.4(NDUFAF5):c.686T>C (p.Leu229Pro)	118203929	NDUFAF5	[]	[]	['Mitochondrial complex I deficiency']
NM_003159.2(CDKL5):c.602T>C (p.Leu201Pro)	587783087	CDKL5	[]	['ATTCYTGGGGAG CTTAGCGATGG']	['not provided']
NM_000046.3(ARSB):c.349T>C (p.Cys117Arg)	118203939	ARSB	[]	[]	['MUCOPOLYSAC CHARIDOSIS, TYPE VI, SEVERE']
NM_000046.3(ARSB):c.707T>C (p.Leu236Pro)	118203940	ARSB	[]	[]	['MUCOPOLYSAC CHARIDOSIS, TYPE VI, MILD']
NM_013319.2(UBIAD1):c.511T>C (p.Ser171Pro)	118203951	UBIAD1	[]	['TCTGGCYCCTTT CTCTACACAGG', 'GGCYCCTTTCTCT ACACAGGAGG']	['Schnyder crystalline corneal dystrophy']
NM_138387.3(G6PC3):c.554T>C (p.Leu185Pro)	118203969	G6PC3	[]	[]	['Severe congenital neutropenia 4, autosomal recessive']
NM_006364.2(SEC23A):c.1144T>C (p.Phe382Leu)	118204000	SEC23A	[]	[]	['Cranioleptoculoseptal dysplasia']
NM_000429.2(MAT1A):c.914T>C (p.Leu305Pro)	118204004	MAT1A	[]	[]	['Methionine adenosyltransferase deficiency, autosomal recessive']
NM_000018.3(ACADVL):c.1372T>C (p.Phe458Leu)	118204017	ACADVL	[]	['TCGCATCYTCCG GATCTTTGAGG', 'CGCATCYTCCGG ATCTTTGAGGG', 'GCATCYTCCGGA TCTTTGAGGGG']	['Very long chain acyl-CoA dehydrogenase deficiency']
NM_000833.4(GRIN2A):c.2T>C (p.Met1Thr)	397518466	GRIN2A	[]	['CTAYGGGCAGA GTGGGCTATTGG']	['Focal epilepsy with speech disorder with or without mental retardation']
NM_015702.2(MMADHC):c.776T>C (p.Leu259Pro)	118204044	MMADHC	[]	[]	['Homocystinuria, cbID type, variant 1']
NM_018077.2(RBM28):c.1052T>C (p.Leu351Pro)	118204055	RBM28	[]	[]	['Alopecia, neurologic defects, and endocrinopathy syndrome']
NM_000237.2(LPL):c.662T>C (p.Ile221Thr)	118204061	LPL	[]	[]	['Hyperlipoproteinaemia, type I']
NM_000237.2(LPL):c.337T>C (p.Trp113Arg)	118204069	LPL	[]	['GGACYGGCTGTC ACGGGCTCAGG']	['Hyperlipoproteinaemia, type I']
NM_000237.2(LPL):c.755T>C (p.Ile252Thr)	118204080	LPL	[]	['GTGAYTGCAGA GAGAGGACTTGG']	['Hyperlipoproteinaemia, type I']
NM_000155.3(GALT)	111033726	GALT	[]	[]	['Deficiency of

:c.580T>C (p.Phe194Leu)					UDPglucose- hexose-1-phosphate uridylyltransferase']
NM_000190.3(HMBS) :c.739T>C (p.Cys247Arg)	118204111	HMBS	[]	['GCTTCGCGCAT CGCTGAAAGGG']	['Acute intermittent porphyria']
NM_000190.3(HMBS) :c.242T>C (p.Leu81Pro)	118204119	HMBS	[]	[]	['Acute intermittent porphyria']
NM_001363.4(DKC1): c.1193T>C (p.Leu398Pro)	199422253	DKC1	[]	[]	['Dyskeratosis congenita X-linked']
NM_004278.3(PIGL): c.500T>C (p.Leu167Pro)	145303331	PIGL	[]	[]	['Zunich neuroectodermal syndrome']
NM_000531.5(OTC):c .602T>C (p.Leu201Pro)	72558407	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c .779T>C (p.Leu260Ser)	72558441	OTC	['ATGT ATYAA TTACA GACAC TTGG']	['ATGTATYAATTA CAGACACTTGG']	['not provided']
NM_000531.5(OTC):c .803T>C (p.Met268Thr)	72558449	OTC	[]	[]	['not provided']
NM_000256.3(MYBP C3):c.1814A>G (p.Asp605Gly)	372371774	MYBPC3	[]	[]	['Primary dilated cardiomyopathy', 'Cardiomyopathy', 'not specified']
NM_000531.5(OTC):c .947T>C (p.Phe316Ser)	72558471	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c .1005+2T>C	72558484	OTC	['ATCAT GGYAA GCAAG AAACA AGG']	['ATCATGGYAAG CAAGAAACAAGG']	['not provided']
NM_000531.5(OTC):c .1018T>C (p.Ser340Pro)	72558489	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c .1022T>C (p.Leu341Pro)	72558490	OTC	[]	[]	['not provided']
NM_007294.3(BRCA1)):c.5291T>C (p.Leu1764Pro)	80357281	BRCA1	['GGCY AGAAA TCTGTT GCTAT GGG']	['GGGCYAGAAAT CTGTTGCTATGG', 'GGCYAGAAATCT GTTGCTATGGG']	['Familial cancer of breast', 'Breast- ovarian cancer, familial 1']
NM_000035.3(ALDO B):c.442T>C (p.Trp148Arg)	118204430	ALDOB	['GGAA GYGGC GTGCT GTGCT GAGG']	['GGAAGYGGCGT GCTGTGCTGAGG']	['Hereditary fructosuria']
NM_000512.4(GALN S):c.413T>C (p.Val138Ala)	118204436	GALNS	[]	[]	['Mucopolysaccharid osis, MPS-IV-A']
NM_024782.2(NHEJ1)):c.367T>C	118204452	NHEJ1	[]	[]	['Severe combined immunodeficiency']

(p.Cys123Arg)					with microcephaly, growth retardation, and sensitivity to ionizing radiation']
NM_007294.3(BRCA1):c.65T>C (p.Leu22Ser)	80357438	BRCA1	[]	['AAATCTYAGAGTGTCCCATCTGG']	['Familial cancer of breast', 'Breast-ovarian cancer, familial 1', 'Hereditary cancer-predisposing syndrome']
m.12297T>C	121434464	MT-TL2	['GTCYT AGGCC CCAAA AATTT TGG']	['GTCYTAGGCCCC AAAAATTTTGG']	['Cardiomyopathy, mitochondrial']
NM_001040431.2(COA3):c.215A>G (p.Tyr72Cys)	139877390	COA3	[]	['CCAYCTGGGGA GGTAGGTTTCAGG']	[]
m.10010T>C	121434476	MT-TG	[]	[]	['Exercise intolerance']
NM_000860.5(HPGD):c.577T>C (p.Ser193Pro)	121434481	HPGD	[]	[]	['Digital clubbing, isolated congenital']
NM_024915.3(GRHL2):c.1192T>C (p.Tyr398His)	587777737	GRHL2	[]	[]	['Ectodermal dysplasia/short stature syndrome']
NM_032374.4(APOPT1):c.353T>C (p.Phe118Ser)	587777786	APOPT1	[]	[]	['Cytochrome-c oxidase deficiency']
NM_001605.2(AARS):c.2251A>G (p.Arg751Gly)	143370729	AARS	[]	[]	['EPILEPTIC ENCEPHALOPATHY, EARLY INFANTILE, 29']
NM_000257.3(MYH7):c.2479T>C (p.Trp827Arg)	730880744	MYH7	[]	[]	['Cardiomyopathy']
NM_017415.2(KLHL3):c.1160T>C (p.Leu387Pro)	199469630	KLHL3			['Pseudohypoadosteronism, type 2']
NM_017415.2(KLHL3):c.1280T>C (p.Met427Thr)	199469642	KLHL3			['Pseudohypoadosteronism, type 2']
NM_005859.4(PURA):c.563T>C (p.Ile188Thr)	793888527	PURA	[]	['GACCA YTGCGCT GCCCGCGCAGG', 'ACCA YTGCGCTG CCCGCGCAGGG', 'CCA YTGCGCTGC CCCGCGCAGGGG']	['not provided', 'Mental retardation, autosomal dominant 31']
NM_007294.3(BRCA1):c.212+2T>C	80358026	BRCA1	[]	[]	['Familial cancer of breast', 'Breast-ovarian cancer, familial 1']
NM_002878.3(RAD51D):c.1A>G (p.Met1Val)	561425038	-	[]	['CGCCCA YGTTCC CCGCAGGCCGG']	['Hereditary cancer-predisposing syndrome']
NM_018960.4(GNMT):c.149T>C (p.Leu50Pro)	121907888	GNMT	[]	[]	['Glycine N-methyltransferase deficiency']

NM_007294.3(BRCA1):c.5074+2T>C	80358089	BRCA1	[]	[]	['Breast-ovarian cancer, familial 1']
NC_012920.1:m.11984T>C	200911567	MT-ND4	[]	[]	['Leigh disease']
NM_000280.4(PAX6):c.773T>C (p.Phe258Ser)	121907925	PAX6	[]	[]	['Congenital ocular coloboma', 'Coloboma of optic disc']
NM_020117.9(LARS):c.1118A>G (p.Tyr373Cys)	201861847	LARS	[]	[]	['Infantile liver failure syndrome 1']
NM_024105.3(ALG12):c.473T>C (p.Leu158Pro)	121907934	ALG12	[]	['TCCYGCTGGCCC TCGCGGCCTGG']	['Congenital disorder of glycosylation type 1G']
NM_000152.3(GAA):c.953T>C (p.Met318Thr)	121907936	GAA	[]	[]	['Glycogen storage disease type II, infantile']
NM_000520.4(HEXA):c.1453T>C (p.Trp485Arg)	121907968	HEXA	[]	[]	['Tay-Sachs disease']
NM_000520.4(HEXA):c.632T>C (p.Phe211Ser)	121907974	HEXA	[]	[]	['Tay-Sachs disease']
NM_000053.3(ATP7B):c.2123T>C (p.Leu708Pro)	121908000	ATP7B	[]	[]	['Wilson disease']
NM_000375.2(UROS):c.217T>C (p.Cys73Arg)	121908012	UROS	[]	[]	['Congenital erythropoietic porphyria']
NM_153212.2(GJB4):c.409T>C (p.Phe137Leu)	80358207	GJB4	[]	['CCTCATCYTCAA GGCCGCCGTGG']	['Erythrokeratoderma variabilis']
NM_000403.3(GALE):c.548T>C (p.Leu183Pro)	121908045	GALE	[]	[]	['UDPglucose-4-epimerase deficiency']
NM_002353.2(TACSTD2):c.557T>C (p.Leu186Pro)	80358228	TACSTD2	[]	['TCGGCYGCACCC CAAGTTCGTGG']	['Lattice corneal dystrophy Type III']
NM_001563.3(IMP1):c.461T>C (p.Leu154Pro)	713993047	IMP1	[]	[]	['Macular dystrophy, vitelliform, 4']
NM_138691.2(TMC1):c.1543T>C (p.Cys515Arg)	121908076	TMC1	[]	['AGGACCTYGCTGGAAACAATGG', 'ACCTYGCTGGGAACAATGGTGG', 'CCTYGCTGGGAAACAATGGTGGG']	['Deafness, autosomal recessive 7']
NM_000271.4(NPC1):c.3182T>C (p.Ile1061Thr)	80358259	NPC1	[]	[]	['Niemann-Pick disease type C1']
NM_006432.3(NPC2):c.295T>C (p.Cys99Arg)	80358264	NPC2	[]	[]	['Niemann-Pick disease type C2']
NM_017838.3(NHP2):c.415T>C (p.Tyr139His)	121908089	NHP2	[]	['GGAGGCTYACGATGAGTGCCTGG', 'GGCTYACGATGATGCCTGGAGG']	['Dyskeratosis congenita autosomal recessive 1', 'Dyskeratosis congenita, autosomal recessive']

					2']
NM_005857.4(ZMPSTE24):c.1018T>C (p.Trp340Arg)	121908093	ZMPSTE24	[]	[]	['Mandibuloacral dysplasia with type B lipodystrophy', 'not provided']
NM_001195794.1(CLRN1):c.488T>C (p.Leu163Pro)	121908142	CLRN1	[]	[]	['Usher syndrome, type 3']
NM_057176.2(BSND):c.35T>C (p.Ile12Thr)	121908144	BSND	[]	[]	['Sensorineural deafness with mild renal dysfunction']
NM_001243133.1(NLRP3):c.1718T>C (p.Phe573Ser)	121908152	NLRP3	[]	[]	['Familial cold urticaria', 'Chronic infantile neurological, cutaneous and articular syndrome']
NM_001243133.1(NLRP3):c.926T>C (p.Phe309Ser)	121908154	NLRP3	[]	['GGTGCCTYTGAC GAGCACATAGG']	['Familial cold urticaria', 'Chronic infantile neurological, cutaneous and articular syndrome']
NM_153741.1(DPM3):c.254T>C (p.Leu85Ser)	121908155	DPM3	[]	[]	['Congenital disorder of glycosylation type 1O']
NM_001033855.2(DCLRE1C):c.2T>C (p.Met1Thr)	121908158	DCLRE1C	[]	['GGCGCTAYGAG TTCTTTCGAGGG', 'GCGCTAYGAGTT CTTTCGAGGGG']	['Histiocytic medullary reticulosis']
NM_017696.2(MCM9):c.1732+2T>C	587777871	MCM9	[]	[]	['Premature ovarian failure 1', 'Ovarian dysgenesis 4']
NM_031433.3(MFRP):c.545T>C (p.Ile182Thr)	121908190	-	[]	[]	['Nanophthalmos 2']
NM_001127221.1(CACNA1A):c.2141T>C (p.Val714Ala)	121908213	CACNA1A	[]	[]	['Familial hemiplegic migraine type 1']
NM_001127221.1(CACNA1A):c.4469T>C (p.Phe1490Ser)	121908233	CACNA1A	[]	[]	['Episodic ataxia type 2']
NM_133459.3(CCBE1):c.520T>C (p.Cys174Arg)	121908254	CCBE1	[]	[]	['Hennekam lymphangiectasia-lymphedema syndrome']
NM_018129.3(PNPO):c.2T>C (p.Met1Thr)	796052870	PNPO	[]	['CCCCCAYGACGT GCTGGCTGCGG', 'CCCCAYGACGTG CTGGCTGCGGG', 'CCCAYGACGTGC TGGCTGCGGGG']	['not provided']
NM_014845.5(FIG4):c.122T>C (p.Ile41Thr)	121908287	FIG4	[]	[]	['Charcot-Marie-Tooth disease, type 4J', 'not provided']
NM_001005741.2(GBA):c.751T>C (p.Tyr251His)	121908300	GBA	['GCCA GAYAC TTTGT GAAGT	['GCCAGAYACTTT GTGAAGTAAGG', 'CCAGAYACTTTG TGAAGTAAGGG']	['Gaucher disease, type 1']

			AAGG']		
NM_020427.2(SLURP1):c.43T>C (p.Trp15Arg)	121908318	SLURP1	[]	['GCAGCCYGGAG CATGGGCTGTGG']	['Acroerythrokeratoderma']
NM_020427.2(SLURP1):c.229T>C (p.Cys77Arg)	121908319	SLURP1	[]	[]	['Acroerythrokeratoderma']
NM_000787.3(DBH):c.339+2T>C	74853476	DBH	[]	[]	['Dopamine beta hydroxylase deficiency']
NM_017882.2(CLN6):c.200T>C (p.Leu67Pro)	154774633	CLN6	['AGCYGGTAT TCCCT CTCGA GTGG']	['AGCYGGTATTCC CTCTCGAGTGG']	['Adult neuronal ceroid lipofuscinosis', 'not provided']
NM_022124.5(CDH23):c.5663T>C (p.Phe1888Ser)	121908352	CDH23	[]	['CTCACCTYCAAC ATCACTGCGGG']	['Deafness, autosomal recessive 12']
NM_054027.4(ANKH):c.143T>C (p.Met48Thr)	121908407	ANKH	['GTCGAGAYG CTGGC CAGCT ACGG', 'TCGAGAYGCT GGCCA GCTAC GGG']	['GTCGAGAYGCT GGCCAGCTACGG', 'TCGAGAYGCTGG CCAGCTACGGG']	['Chondrocalcinosis 2']
NM_004924.4(ACTN4):c.784T>C (p.Ser262Pro)	121908417	ACTN4	[]	[]	['Focal segmental glomerulosclerosis 1']
NM_014384.2(ACAD8):c.455T>C (p.Met152Thr)	121908418	ACAD8	[]	[]	['Deficiency of isobutyryl-CoA dehydrogenase']
NM_153717.2(EVC):c.919T>C (p.Ser307Pro)	121908426	EVC	[]	[]	['Chondroectodermal dysplasia', 'Curry-Hall syndrome']
NM_001040108.1(MLH3):c.3826T>C (p.Trp1276Arg)	121908439	MLH3	[]	[]	['Hereditary nonpolyposis colorectal cancer type 7']
NM_013339.3(ALG6):c.1432T>C (p.Ser478Pro)	121908444	ALG6	[]	[]	['Congenital disorder of glycosylation type 1C']
NM_003835.3(RGS9):c.895T>C (p.Trp299Arg)	121908449	RGS9	[]	[]	['Prolonged electroretinal response suppression']
NM_022336.3(EDAR):c.259T>C (p.Cys87Arg)	121908451	EDAR	[]	[]	['Autosomal recessive hypohidrotic ectodermal dysplasia syndrome']
NM_014270.4(SLC7A9):c.131T>C (p.Ile44Thr)	121908485	SLC7A9	[]	[]	['Cystinuria']
NM_000030.2(AGXT):c.613T>C (p.Ser205Pro)	121908520	AGXT	[]	['CCTGTACYCGGG CTCCAGAAGG']	['Primary hyperoxaluria, type I']

NM_000030.2(AGXT):c.731T>C (p.Ile244Thr)	121908525	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_000026.2(ADSL):c.1312T>C (p.Ser438Pro)	119450940	ADSL	[]	[]	['Adenylosuccinate lyase deficiency']
NM_000026.2(ADSL):c.674T>C (p.Met225Thr)	119450945	ADSL	['AAGAYGGTGACAGAAAAGGCAGG']	['AAGAYGGTGACAGAAAAGGCAGG']	['Adenylosuccinate lyase deficiency']
NM_014985.3(CEP152):c.2000A>G (p.Lys667Arg)	200879436	CEP152	[]	[]	['Seckel syndrome 5', 'not specified']
NM_002755.3(MAP2K1):c.158T>C (p.Phe53Ser)	121908594	MAP2K1	[]	[]	['Cardiofaciocutaneous syndrome 3', 'Rasopathy']
NM_004820.3(CYP7B1):c.647T>C (p.Phe216Ser)	121908612	CYP7B1	[]	[]	['Spastic paraplegia 5A']
NM_004273.4(CHST3):c.776T>C (p.Leu259Pro)	121908616	CHST3	[]	[]	['Spondyloepiphyseal dysplasia with congenital joint dislocations']
NM_004273.4(CHST3):c.920T>C (p.Leu307Pro)	121908618	CHST3	[]	['CGTGCGYGGCCTCGCGCATGGTGG']	['Spondyloepiphyseal dysplasia with congenital joint dislocations']
NM_004273.4(CHST3):c.857T>C (p.Leu286Pro)	121908620	CHST3	[]	[]	['Spondyloepiphyseal dysplasia with congenital joint dislocations']
NM_000050.4(ASS1):c.535T>C (p.Trp179Arg)	121908646	ASS1	[]	[]	[]
NM_030761.4(WNT4):c.35T>C (p.Leu12Pro)	121908653	WNT4	[]	[]	['Mullerian aplasia and hyperandrogenism']
NM_006432.3(NPC2):c.199T>C (p.Ser67Pro)	11694	NPC2	[]	['TATTCAGYCTAA AAGCAGCAAGG']	['Niemann-Pick disease type C2']
NM_003839.3(TNFRSF11A):c.523T>C (p.Cys175Arg)	121908656	TNFRSF11A	[]	[]	['Osteopetrosis autosomal recessive 7']
NM_000022.2(ADA):c.320T>C (p.Leu107Pro)	121908739	ADA	[]	['CCTGCGYGGCCAACTCCAAAGTGG']	['Severe combined immunodeficiency due to ADA deficiency']
NM_000140.3(FECH):c.315-48T>C	2272783	FECH	[]	[]	['Erythropoietic protoporphyria']
NM_000059.3(BRCA2):c.7529T>C (p.Leu2510Pro)	80358979	BRCA2	[]	[]	['Familial cancer of breast', 'Breast-ovarian cancer, familial 2', 'Fanconi anemia, complementation group D1']
NM_003722.4(TP63):c.1033T>C (p.Cys345Arg)	121908837	TP63	[]	[]	['Ectrodactyly, ectodermal dysplasia, and cleft

					lip/palate syndrome 3']
NM_003722.4(TP63):c.1646T>C (p.Ile549Thr)	121908845	TP63	[]	[]	['Hay-Wells syndrome of ectodermal dysplasia', 'Rapp-Hodgkin ectodermal dysplasia syndrome']
NM_000059.3(BRCA2):c.7958T>C (p.Leu2653Pro)	80359022	BRCA2	[]	['TGCYTCTTCAAC TAAAATACAGG']	['Familial cancer of breast', 'Breast-ovarian cancer, familial 2']
NM_000369.2(TSHR):c.1891T>C (p.Phe631Leu)	121908861	TSHR	[]	[]	['Hyperthyroidism, nonautoimmune', 'Thyroid adenoma, hyperfunctioning']
NM_000369.2(TSHR):c.1358T>C (p.Met453Thr)	121908864	TSHR	[]	[]	['Hyperthyroidism, nonautoimmune']
NM_000369.2(TSHR):c.1526T>C (p.Val509Ala)	121908874	TSHR	[]	[]	['Hyperthyroidism, nonautoimmune']
NM_000369.2(TSHR):c.1798T>C (p.Cys600Arg)	121908884	TSHR	[]	[]	['Hypothyroidism, congenital, nongoitrous, 1']
NM_000369.2(TSHR):c.1400T>C (p.Leu467Pro)	121908885	TSHR	[]	[]	['Hypothyroidism, congenital, nongoitrous, 1']
NM_001457.3(FLNB):c.703T>C (p.Ser235Pro)	121908896	FLNB	[]	[]	['Boomerang dysplasia']
NM_003880.3(WISP3):c.232T>C (p.Cys78Arg)	121908902	WISP3	[]	['AAAATCYGTGCC AAGCAACCAGG', 'AAATCYGTGCCA AGCAACCAGGG', 'AATCYGTGCCAA GCAACCAGGGG']	['Progressive pseudorheumatoid dysplasia']
NM_003880.3(WISP3):c.1000T>C (p.Ser334Pro)	121908903	WISP3	[]	[]	['Progressive pseudorheumatoid dysplasia']
NM_002977.3(SCN9A):c.4382T>C (p.Ile1461Thr)	121908914	-	[]	[]	['Paroxysmal extreme pain disorder']
NM_004086.2(COCH):c.349T>C (p.Trp117Arg)	121908929	-	[]	[]	['Deafness, autosomal dominant 9']
NM_004086.2(COCH):c.1535T>C (p.Met512Thr)	121908934	-	['AGAT AYGGC TTCTA AACCG AAGG']	['AGATAYGGCTTC TAAACCGAAGG']	['Deafness, autosomal dominant 9']
NM_006892.3(DNMT3B):c.808T>C (p.Ser270Pro)	121908947	DNMT3B	[]	['CAAGTTCYCCGA GGTGAGTCCGG', 'AAGTTCYCCGAG GTGAGTCCGGG', 'AGTTCYCCGAGG TGAGTCCGGGG']	['Centromeric instability of chromosomes 1,9 and 16 and immunodeficiency']
NM_000226.3(KRT9):c.503T>C	61157095	KRT9	[]	[]	['Epidermolytic palmoplantar

(p.Leu168Ser)					keratoderma', 'not provided']
NM_000051.3(ATM): c.8584+2T>C	730881326	-	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000492.3(CFTR): c.3857T>C (p.Phe1286Ser)	121909028	CFTR	[]	['AGCCTYTGAGT GATACCACAGG']	['Cystic fibrosis']
NM_000492.3(CFTR): c.3763T>C (p.Ser1255Pro)	121909041	CFTR	[]	[]	['Cystic fibrosis']
NM_001040667.2(HSF4): c.341T>C (p.Leu114Pro)	121909048	HSF4	[]	[]	['Cataract, zonular']
NM_005025.4(SERPINI1): c.145T>C (p.Ser49Pro)	121909051	SERPINI1	[]	[]	['Familial encephalopathy with neuroserpin inclusion bodies']
NM_002700.2(POU4F3): c.668T>C (p.Leu223Pro)	121909057	POU4F3	[]	[]	['Deafness, autosomal dominant 15']
NM_003322.4(TULP1): c.1471T>C (p.Phe491Leu)	121909074	TULP1	[]	[]	['Retinitis pigmentosa 14']
NM_003322.4(TULP1): c.1145T>C (p.Phe382Ser)	121909076	TULP1	[]	[]	['Retinitis pigmentosa', 'Retinitis pigmentosa 14']
NM_003000.2(SDHB): c.487T>C (p.Ser163Pro)	33927012	SDHB	[]	[]	['Pheochromocytoma', 'Hereditary Paraganglioma-Pheochromocytoma Syndromes', 'Hereditary cancer-predisposing syndrome', 'Cowden syndrome 2', 'not specified', 'not provided']
NM_005603.4(ATP8B1): c.863T>C (p.Leu288Ser)	121909099	ATP8B1	[]	[]	['Progressive intrahepatic cholestasis']
NM_005603.4(ATP8B1): c.1982T>C (p.Ile661Thr)	121909100	ATP8B1	[]	[]	['Progressive intrahepatic cholestasis', 'Benign recurrent intrahepatic cholestasis 1']
NM_000483.4(APOC2): c.142T>C (p.Trp48Arg)	120074115	-	[]	[]	['Apolipoprotein C2 deficiency']
NM_000543.4(SMPD1): c.911T>C (p.Leu304Pro)	120074124	SMPD1	['CACY TGTGA GGAAG TTCCT GGGG']	['AGCACYTGTGA GGAAGTTCCTGG', 'GCACYTGTGAGG AAGTTCCTGGG', 'CACYGTGTGAGGA AGTTCCTGGGG']	['Sphingomyelin/cholesterol lipidosi', 'Niemann-Pick disease, type A', 'Niemann-Pick disease, type B', 'not provided']
NM_000085.4(CLCNKB): c.1294T>C	121909135	CLCNKB	[]	['CTTTGTCYATGG TGAGTCTGGGG']	['Bartter syndrome type 3']

(p.Tyr432His)					
NM_001300.5(KLF6): c.346T>C (p.Ser116Pro)	121909139	KLF6	[]	[]	[]
m.12338T>C	201863060	MT-ND5	[]	[]	['Leber optic atrophy']
NM_001300.5(KLF6): c.190T>C (p.Trp64Arg)	121909142	KLF6	['TCTGY GGACC AAAAT CATT TGG']	['TCTGYGGACCAA AATCATTCTGG']	[]
NM_001300.5(KLF6): c.506T>C (p.Leu169Pro)	121909143	KLF6	[]	['GGAGCYGCCCTC GCCAGGGAAGG']	[]
NM_000271.4(NPC1): c.337T>C (p.Cys113Arg)	120074136	NPC1	[]	[]	['Niemann-Pick disease type C1']
NM_000019.3(ACAT1): c.935T>C (p.Ile312Thr)	120074146	ACAT1	['CAAG AAYAG TAGGT AAGGC CAGG']	['CAAGAAYAGTA GGTAAGGCCAGG']	['Deficiency of acetyl-CoA acetyltransferase']
NM_017890.4(VPS13B): c.8459T>C (p.Ile2820Thr)	120074155	VPS13B	[]	[]	['Cohen syndrome']
NM_017653.3(DYM): c.1624T>C (p.Cys542Arg)	120074165	DYM	[]	[]	['Smith McCort dysplasia']
NM_001089.2(ABCA3): c.302T>C (p.Leu101Pro)	121909182	ABCA3	[]	['GCACYTGTGATC AACATGCGAGG']	['Surfactant metabolism dysfunction, pulmonary, 3']
NM_001089.2(ABCA3): c.4658T>C (p.Leu1553Pro)	121909183	ABCA3	[]	[]	['Surfactant metabolism dysfunction, pulmonary, 3']
NM_001089.2(ABCA3): c.977T>C (p.Leu326Pro)	121909185	ABCA3	[]	[]	['Surfactant metabolism dysfunction, pulmonary, 3']
NM_000474.3(TWIST1): c.392T>C (p.Leu131Pro)	121909189	TWIST1	[]	[]	['Saethre-Chotzen syndrome']
NM_000503.5(EYA1): c.1459T>C (p.Ser487Pro)	121909200	EYA1	[]	['CACTCYCGCTCA TTCCTCCCGG']	['Melnick-Fraser syndrome']
NM_000358.2(TGFBI): c.1619T>C (p.Phe540Ser)	121909214	TGFBI	[]	[]	['Lattice corneal dystrophy type 3A']
NM_000426.3(LAMA2): c.8282T>C (p.Ile2761Thr)	115650537	LAMA2	['TTGA YAGGG AGCAA GCACT TCGG', TGAYA GGGAG CAAGC AGTTC GGG']	['TTGAYAGGGAG CAAGCAGTTCGG', TGAYAGGGAGCA AGCAGTTCGGG']	['Merosin deficient congenital muscular dystrophy']

NM_000314.6(PTEN): c.209T>C (p.Leu70Pro)	121909226	PTEN	[]	[]	['Cowden syndrome 1', 'Hereditary cancer-predisposing syndrome']
NM_000314.6(PTEN): c.335T>C (p.Leu112Pro)	121909230	PTEN	[]	[]	['Lhermitte-Duclos disease']
NM_000314.6(PTEN): c.722T>C (p.Phe241Ser)	121909240	PTEN	[]	[]	['Macrocephaly/autism syndrome']
NM_004970.2(IGFALS): c.1618T>C (p.Cys540Arg)	121909247	IGFALS	[]	['GGACYGTGGCTGCCCTCTCAAGG']	['Acid-labile subunit deficiency']
NM_005570.3(LMAN1): c.2T>C (p.Met1Thr)	121909253	LMAN1	[]	['AGAYGGCGGGA TCCAGGCAAAGG']	['Combined deficiency of factor V and factor VIII, 1']
NM_005055.4(RAPSN): c.416T>C (p.Phe139Ser)	121909256	RAPSN	[]	[]	['Pena-Shokeir syndrome type I']
NM_000391.3(TPP1): c.887-10A>G	755445790	TPP1	['TTTTT TTTTT TTTTT TTGAGG']	['TTTTTTTTTTTTTT TTTTTTGAGG']	['Ceroid lipofuscinosis, neuronal, 2', 'not provided']
NM_006302.2(MOGS): c.1954T>C (p.Phe652Leu)	121909292	MOGS	[]	[]	['Congenital disorder of glycosylation type 2B']
NM_005379.3(MYO1A): c.2728T>C (p.Ser910Pro)	121909306	MYO1A	[]	[]	['Deafness, autosomal dominant 48']
NM_178151.2(DCX): c.128T>C (p.Leu43Ser)	587783521	DCX	[]	[]	['Heterotopia']
NM_001127221.1(CACNA1A): c.5126T>C (p.Ile1709Thr)	121909326	CACNA1A	[]	[]	['Spinocerebellar ataxia 6', 'Familial hemiplegic migraine type 1', 'Episodic ataxia type 2']
NM_001451.2(FOXF1): c.1138T>C (p.Ter380Arg)	121909337	FOXF1	['TGATGYGAGGCTGCGCCG CAGG']	['TGATGYGAGGCTGCCGCCG CAGG']	['Alveolar capillary dysplasia with misalignment of pulmonary veins']
NM_000163.4(GHR): c.341T>C (p.Phe114Ser)	121909357	GHR	[]	[]	['Laron-type isolated somatotropin defect']
NM_000163.4(GHR): c.512T>C (p.Ile171Thr)	121909367	GHR	[]	[]	['Laron-type isolated somatotropin defect']
NM_000339.2(SLC12A3): c.1868T>C (p.Leu623Pro)	121909385	SLC12A3	[]	['CAACCYGGCCCTCAGCTACTCGG']	['Familial hypokalemia-hypomagnesemia']
NM_001174089.1(SLC4A11): c.2480T>C (p.Leu827Pro)	121909394	SLC4A11	[]	[]	['Corneal dystrophy and perceptive deafness']
NM_001174089.1(SLC4A11): c.589T>C (p.Ser197Pro)	121909395	SLC4A11	[]	[]	['Corneal dystrophy and perceptive deafness']
NM_000519.3(HBD): c.	34975911	HBD	[]	[]	['delta Thalassemia']

.-127T>C					
NM_002427.3(MMP13):c.224T>C (p.Phe75Ser)	121909497	MMP13	[]	['TTCTYCGGCTTAGAGGTGACTGG']	['Spondyloepimetaphyseal dysplasia, Missouri type']
NM_002427.3(MMP13):c.221T>C (p.Phe74Ser)	121909498	MMP13	[]	[]	[]
NM_002427.3(MMP13):c.272T>C (p.Met91Thr)	121909499	MMP13	['GTCA YGAAA AAGCC AAGAT GCGG', 'TCAYG AAAAA GCCAA GATCGGG']	['GTCA YGAAAAA GCCAAGATGCGG', 'TCAYGAAAAAGC CAAGATGCGGG']	[]
NM_000751.2(CHRND):c.283T>C (p.Phe95Leu)	121909506	CHRND	[]	[]	['Lethal multiple pterygium syndrome']
NM_000751.2(CHRND):c.188T>C (p.Leu63Pro)	121909508	CHRND	[]	['AACCYCATCTCCCTGGTGAGAGG']	['MYASTHENIC SYNDROME, CONGENITAL, 3B, FAST-CHANNEL']
NM_001100.3(ACTA1):c.287T>C (p.Leu96Pro)	121909519	ACTA1	[]	['CGAGCYTCGCGTGGCTCCCAGG']	['Nemaline myopathy 3']
NM_001100.3(ACTA1):c.668T>C (p.Leu223Pro)	121909530	ACTA1	[]	[]	['Congenital myopathy with fiber type disproportion']
NM_000488.3(SERPINC1):c.1141T>C (p.Ser381Pro)	121909565	SERPINC1	[]	[]	['Antithrombin III deficiency']
NM_000488.3(SERPINC1):c.442T>C (p.Ser148Pro)	121909569	SERPINC1	[]	[]	['Antithrombin III deficiency']
NM_000488.3(SERPINC1):c.667T>C (p.Ser223Pro)	121909572	SERPINC1	[]	['TGGGTGYCCAATAAGACCGAAGG']	['Antithrombin III deficiency']
NM_000488.3(SERPINC1):c.379T>C (p.Cys127Arg)	121909573	SERPINC1	[]	[]	['Antithrombin III deficiency']
NM_023110.2(FGFR1):c.899T>C (p.Ile300Thr)	121909633	FGFR1	[]	[]	['Interfrontal craniofaciosynostosis']
NM_023110.2(FGFR1):c.1141T>C (p.Cys381Arg)	121909634	FGFR1	[]	[]	['Osteoglophonic dysplasia']
NM_182925.4(FLT4):c.3131T>C (p.Leu1044Pro)	121909651	FLT4	[]	[]	['Hereditary lymphedema type I']
NM_182925.4(FLT4):c.3257T>C (p.Ile1086Thr)	121909655	FLT4	[]	[]	['Hereditary lymphedema type I']
NM_000145.3(FSHR):c.479T>C (p.Ile160Thr)	121909659	FSHR	[]	[]	['Ovarian dysgenesis 1']
NM_000145.3(FSHR):c.1634T>C	121909664	FSHR	[]	[]	['Ovarian hyperstimulation']

(p.Ile545Thr)					syndrome']
NM_000821.6(GGCX):c.896T>C (p.Phe299Ser)	121909677	GGCX	[]	['TATGTYCTCCTA CGTCATGCTGG']	['Pseudothrombocytopenia-like disorder with multiple coagulation factor deficiency']
NM_001018077.1(NR3C1):c.2209T>C (p.Phe737Leu)	121909727	NR3C1	[]	['CTATTGTCCTCA AACATTTTTGG']	['Glucocorticoid resistance, generalized']
NM_005271.3(GLUD1):c.1501T>C (p.Ser501Pro)	121909732	GLUD1	[]	[]	['Hyperinsulinism-hyperammonemia syndrome']
NM_004614.4(TK2):c.278A>G (p.Asn93Ser)	142291440	TK2	[]	[]	['Mitochondrial DNA depletion syndrome 2']
NM_032977.3(CASP10):c.440T>C (p.Met147Thr)	121909776	CASP10	[]	[]	['Neoplasm of stomach']
NM_000250.1(MPO):c.518A>G (p.Tyr173Cys)	78950939	MPO	['TGCGGYATT TGCTC CGGG']	['GTGCGGYATTTG TCCTGCTCCGG', 'TGCGGYATTTGT CCTGCTCCGGG']	['Myeloperoxidase deficiency']
NM_001139.2(ALOX12B):c.1562A>G (p.Tyr521Cys)	199766569	ALOX12B	[]	[]	['Autosomal recessive congenital ichthyosis 2']
NM_022041.3(GAN):c.1268T>C (p.Ile423Thr)	119485091	GAN	['AAGAAAYC TACGC CATGG GTGG']	['AAGAAAAYCTA CGCCATGGGTGG', 'AAAAYCTACGCC ATGGGTGGAGG']	['Giant axonal neuropathy']
NM_014009.3(FOXP3):c.970T>C (p.Phe324Leu)	122467173	FOXP3	['GACAGAGYT CCTCC ACAAC ATGG']	['GACAGAGYTCCT CCACAACATGG']	['Insulin-dependent diabetes mellitus secretory diarrhea syndrome']
NM_014009.3(FOXP3):c.1099T>C (p.Phe367Leu)	122467175	FOXP3	[]	[]	['Insulin-dependent diabetes mellitus secretory diarrhea syndrome']
NM_004239.3(TRIP11):c.2102A>G (p.Asn701Ser)	139539448	TRIP11	[]	[]	['Achondrogenesis, type IA']
NM_001104.3(ACTN3):c.1729C>T (p.Arg577Ter)	1815739	ACTN3	[]	[]	['Sprinting performance', 'Actn3 deficiency']
NM_002693.2(POLG):c.2636A>G (p.Gln879Arg)	368587966	POLG	[]	[]	['not provided']
NM_000552.3(VWF):c.3178T>C (p.Cys1060Arg)	61748497	VWF	[]	[]	['von Willebrand disease type 2N', 'not provided']
NM_000552.3(VWF):c.3445T>C (p.Cys1149Arg)	61748511	VWF	[]	[]	['von Willebrand disease type 1', 'not provided']
NM_000184.2(HBG2):c.125T>C (p.Phe42Ser)	34878913	HBG2	['CAGAGGTYC TTTGA CAGCT']	['CAGAGGTYCTTT GACAGCTTTGG']	['Cyanosis, transient neonatal']

			TTGG']		
NM_000371.3(TTR):c.190T>C (p.Phe64Leu)	138065384	TTR	[]	[]	['Cardiomyopathy', 'not specified']
NM_000402.4(G6PD):c.1058T>C (p.Leu353Pro)	76723693	G6PD	[]	[]	['Glucose 6 phosphate dehydrogenase deficiency', 'Anemia, nonspherocytic hemolytic, due to G6PD deficiency', 'not provided']
NM_177405.2(CECR1):c.355A>G (p.Thr119Ala)	775440641	CECR1	[]	[]	['Idiopathic livedo reticularis with systemic involvement']
NM_000218.2(KCNQ1):c.401T>C (p.Leu134Pro)	199472685	KCNQ1			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.625T>C (p.Ser209Pro)	199472705	KCNQ1			['Atrial fibrillation, familial, 3', 'Atrial fibrillation']
NM_000218.2(KCNQ1):c.752T>C (p.Leu251Pro)	199472716	KCNQ1			['Congenital long QT syndrome', 'Cardiac arrhythmia', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.824T>C (p.Phe275Ser)	199472729	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.832T>C (p.Tyr278His)	199472731	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.845T>C (p.Leu282Pro)	199472733	KCNQ1			['Congenital long QT syndrome']
NM_000257.3(MYH7):c.730T>C (p.Phe244Leu)	730880849	MYH7	[]	[]	['Cardiomyopathy']
NM_000218.2(KCNQ1):c.908T>C (p.Leu303Pro)	199472740	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.913T>C (p.Trp305Arg)	199472741	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.1045T>C (p.Ser349Pro)	199472764	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.1117T>C (p.Ser373Pro)	199472766	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.1165T>C (p.Ser389Pro)	199472772	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.1174T>C (p.Trp392Arg)	199472774	KCNQ1			['Congenital long QT syndrome']

NM_000218.2(KCNQ1):c.1541T>C (p.Ile514Thr)	199472786	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.1696T>C (p.Ser566Pro)	199472803	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.1805T>C (p.Leu602Pro)	199472818	-			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.608T>C (p.Leu203Pro)	199472823	KCNQ1			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2):c.65T>C (p.Phe22Ser)	199472826	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.86T>C (p.Phe29Ser)	199472831	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.89T>C (p.Ile30Thr)	199472832	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.160T>C (p.Tyr54His)	199472843	KCNH2			['Congenital long QT syndrome']
NM_001165963.1(SCN1A):c.662T>C (p.Leu221Pro)	796052961	SCN1A	[]	[]	['not provided']
NM_000238.3(KCNH2):c.287T>C (p.Ile96Thr)	199472853	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.322T>C (p.Cys108Arg)	199472859	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.872T>C (p.Met291Thr)	199472881	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1238T>C (p.Leu413Pro)	199472893	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1279T>C (p.Tyr427His)	199472898	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1387T>C (p.Phe463Leu)	199472904	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1655T>C (p.Leu552Ser)	199472918	KCNH2	[]	[]	['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000061.2(BTK):c.1955T>C (p.Leu652Pro)	128622212	BTK	[]	[]	['X-linked agammaglobulinemia']
NM_000238.3(KCNH2):c.1691T>C (p.Leu564Pro)	199472924	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1702T>C (p.Trp568Arg)	199472927	KCNH2			['Congenital long QT syndrome']
NM_000138.4(FBN1):	794728333	FBN1	[]	[]	['Thoracic aortic

c.5726T>C (p.Ile1909Thr)					aneurysms and aortic dissections']
NM_015884.3(MBTP S2):c.1424T>C (p.Phe475Ser)	122468179	MBTPS2	[]	[]	['IFAP syndrome with or without BRESHECK syndrome']
NM_000238.3(KCNH 2):c.1985T>C (p.Ile662Thr)	199472980	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH 2):c.2033T>C (p.Leu678Pro)	199472981	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH 2):c.2078T>C (p.Leu693Pro)	199472983	KCNH2			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH 2):c.2309T>C (p.Val770Ala)	199472994	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH 2):c.3146T>C (p.Leu1049Pro)	199473026	KCNH2			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.278T>C (p.Phe93Ser)	199473052	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.544T>C (p.Cys182Arg)	199473066	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.689T>C (p.Ile230Thr)	199473073	SCN5A			['Cardiac conduction defect, nonspecific']
NM_000335.4(SCN5A):c.1187T>C (p.Val396Ala)	199473103	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.1190T>C (p.Ile397Thr)	199473105	SCN5A			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.2018T>C (p.Leu673Pro)	199473141	SCN5A			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.2516T>C (p.Leu839Pro)	199473164	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.2783T>C (p.Leu928Pro)	199473178	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.2804T>C (p.Leu935Pro)	199473179	SCN5A			['Brugada syndrome']
NM_198056.2(SCN5A):c.3010T>C (p.Cys1004Arg)	199473183	SCN5A	[]	[]	['Congenital long QT syndrome', 'not specified', 'not provided']
NM_000335.4(SCN5A):c.3679T>C (p.Tyr1227His)	199473205	SCN5A	[]	[]	['Brugada syndrome']
NM_000335.4(SCN5A):c.3713T>C (p.Leu1238Pro)	199473210	SCN5A			['Brugada syndrome']
NM_000492.3(CFTR):	139573311	CFTR	[]	['TTCACYTCTAAT	['Cystic fibrosis']

c.1400T>C (p.Leu467Pro)				GGTGATTATGG', 'TCACYTCTAATG GTGATTATGGG']	
NM_000335.4(SCN5A)):c.3929T>C (p.Leu1310Pro)	199473219	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A)):c.4027T>C (p.Phe1343Leu)	199473228	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A)):c.4028T>C (p.Phe1343Ser)	199473229	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A)):c.4034T>C (p.Leu1345Pro)	199473231	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A)):c.4340T>C (p.Ile1447Thr)	199473251	SCN5A			['Brugada syndrome']
NM_198056.2(SCN5A)):c.4493T>C (p.Met1498Thr)	199473263	SCN5A			['Congenital long QT syndrome', 'not provided']
NM_000335.4(SCN5A)):c.4742T>C (p.Leu1581Pro)	199473275	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A)):c.4778T>C (p.Phe1593Ser)	199473277	SCN5A			['Congenital long QT syndrome']
NM_000335.4(SCN5A)):c.5179T>C (p.Cys1727Arg)	199473302	SCN5A			['Brugada syndrome']
NM_000219.5(KCNE1)):c.158T>C (p.Phe53Ser)	199473355	KCNE1			['Congenital long QT syndrome']
NM_000219.5(KCNE1)):c.259T>C (p.Trp87Arg)	199473361	KCNE1			['Congenital long QT syndrome']
NM_000891.2(KCNJ2)):c.301T>C (p.Cys101Arg)	199473374	KCNJ2			['Ventricular tachycardia']
NM_000218.2(KCNQ1)):c.560T>C (p.Leu187Pro)	199473399	KCNQ1	[]	[]	['Congenital long QT syndrome', 'Cardiac arrhythmia', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1)):c.572T>C (p.Leu191Pro)	199473401	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1)):c.1052T>C (p.Phe351Ser)	199473402	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1)):c.1058T>C (p.Leu353Pro)	199473403	KCNQ1			['Congenital long QT syndrome', 'Cardiac arrhythmia', 'Long QT syndrome, LQT1 subtype']
NM_000238.3(KCNH2)):c.202T>C	199473417	KCNH2			['Congenital long QT syndrome',

(p.Phe68Leu)					'Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.341T>C (p.Leu114Pro)	199473448	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.716T>C (p.Leu239Pro)	199473458	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.742T>C (p.Trp248Arg)	199473459	KCNQ1			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.797T>C (p.Leu266Pro)	199473460	KCNQ1			['Long QT syndrome', 'Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.829T>C (p.Ser277Pro)	199473461	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.910T>C (p.Trp304Arg)	199473466	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.1550T>C (p.Ile517Thr)	199473478	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.1661T>C (p.Val554Ala)	199473481	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.2T>C (p.Met1Thr)	199473485	KCNQ1			['Congenital long QT syndrome', 'KCNQ1-related Jervell and Lange-Nielsen syndrome']
NM_000238.3(KCNH2):c.260T>C (p.Leu87Pro)	199473495	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1700T>C (p.Ile567Thr)	199473519	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1705T>C (p.Tyr569His)	199473520	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1816T>C (p.Ser606Pro)	199473523	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1889T>C (p.Val630Ala)	199473526	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1945T>C (p.Ser649Pro)	199473530	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.2452T>C (p.Ser818Pro)	199473537	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.2573T>C	199473539	KCNH2			['Congenital long QT syndrome']

(p.Ile858Thr)					
NM_000335.4(SCN5A):c.407T>C (p.Leu136Pro)	199473557	SCN5A			['Brugada syndrome']
NM_020166.4(MCCC1):c.1310T>C (p.Leu437Pro)	119103215	MCCC1	[]	[]	['3 Methylcrotonyl-CoA carboxylase 1 deficiency']
NM_198056.2(SCN5A):c.944T>C (p.Leu315Pro)	199473564	SCN5A			['Brugada syndrome', 'not provided']
NM_002972.3(SBF1):c.1249A>G (p.Met417Val)	587776986	SBF1	[]	[]	['Charcot-Marie-Tooth disease, type 4B3']
NM_000335.4(SCN5A):c.2551T>C (p.Phe851Leu)	199473586	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.2743T>C (p.Cys915Arg)	199473588	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.4046T>C (p.Ile1349Thr)	199473607	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.4453T>C (p.Phe1485Leu)	199473615	SCN5A			['Sudden infant death syndrome']
NM_000335.4(SCN5A):c.5111T>C (p.Phe1704Ser)	199473627	SCN5A			['Sudden infant death syndrome']
NM_000891.2(KCNJ2):c.650T>C (p.Leu217Pro)	199473656	KCNJ2			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.550T>C (p.Tyr184His)	199473661	KCNQ1	['AGCAAGBACGTGGG CCTCTGGG']	['CAGCAAGBACGTGGCCTCTGGG', 'AGCAAGBACGTGGCCTCTGGG', 'GCAAGBACGTGGCCTCTGGGGG']	['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2):c.206T>C (p.Leu69Pro)	199473665	KCNH2			['Congenital long QT syndrome']
NM_001130823.1(DNMT1):c.1531T>C (p.Tyr511His)	199473692	DNMT1			['Hereditary sensory neuropathy type IE']
NM_031226.2(CYP19A1):c.743+2T>C	786205107	-	['CTGTG YAAGT AATAC AACTT TGG']	['CTGTGYAAGTAA TACAACCTTTGG']	['Aromatase deficiency']
NM_016373.3(WWOX):c.872T>C (p.Leu291Pro)	119487098	WWOX	[]	[]	[]
NM_001287223.1(SCN11A):c.1142T>C (p.Ile381Thr)	606231280	SCN11A	['TTCAY TGTGG TCATTT TCCTGG']	['CTTCAYTGTGGT CATTTCCTGG', 'TTCAYTGTGGTC ATTTTCCTGGG']	['Episodic pain syndrome, familial, 3']
NM_003640.3(IKBKAP):c.2204+6T>C	111033171	IKBKAP	[]	[]	['Familial dysautonomia', 'not provided']

NM_000238.3(KCNH2):c.1282T>C (p.Ser428Pro)	794728368	KCNH2	[]	[]	['Cardiac arrhythmia']
NM_000441.1(SLC26A4):c.1588T>C (p.Tyr530His)	111033254	SLC26A4	[]	[]	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_206933.2(USH2A):c.10561T>C (p.Trp3521Arg)	111033264	USH2A	[]	[]	['Usher syndrome, type 2A']
NM_206933.2(USH2A):c.1606T>C (p.Cys536Arg)	111033273	USH2A	['ATATAGAYG CCTCT GCTCC CAGG']	['ATATAGAYGCCT CTGCTCCCAGG']	['Usher syndrome, type 2A']
NM_001363.4(DKC1):c.1049T>C (p.Met350Thr)	121912300	DKC1	[]	[]	['Dyskeratosis congenita X-linked']
NM_000274.3(OAT):c.1180T>C (p.Cys394Arg)	121965054	OAT	[]	[]	['Ornithine aminotransferase deficiency']
NM_001302946.1(TRNT1):c.497T>C (p.Leu166Ser)	606231289	TRNT1	['ACTTYATTTG ACTAC TTAA TGG']	['ACTTYATTTGAC TACTTTAATGG']	['Sideroblastic anemia with B-cell immunodeficiency, periodic fevers, and developmental delay']
NM_000454.4(SOD1):c.341T>C (p.Ile114Thr)	121912441	SOD1	[]	['CATCAYTGCCG CACACTGGTGG']	['Amyotrophic lateral sclerosis type 1']
NM_000454.4(SOD1):c.434T>C (p.Leu145Ser)	121912446	SOD1	[]	['CGTTYGGCTTGT GGTGTAATTGG', 'GTTYGGCTTGTG GTGTAATTGGG']	['Amyotrophic lateral sclerosis type 1']
NM_000454.4(SOD1):c.455T>C (p.Ile152Thr)	121912449	SOD1	[]	[]	['Amyotrophic lateral sclerosis type 1']
NM_000213.3(ITGB4):c.467T>C (p.Leu156Pro)	121912461	ITGB4	[]	[]	['Epidermolysis bullosa with pyloric atresia']
NM_000213.3(ITGB4):c.1684T>C (p.Cys562Arg)	121912463	ITGB4	[]	['GGCCAGYGTGT GTGTGAGCCTGG']	['Epidermolysis bullosa with pyloric atresia']
NM_000213.3(ITGB4):c.112T>C (p.Cys38Arg)	121912465	ITGB4	[]	[]	['Epidermolysis bullosa with pyloric atresia']
NM_002198.2(IRF1):c.31T>C (p.Trp11Arg)	121912470	IRF1	[]	[]	['Non-small cell lung cancer']
NM_000424.3(KRT5):c.20T>C (p.Val7Ala)	121912474	KRT5	['TCAA GTGYG TCCTC CGGAG CGG', 'CAAGT GYGTC CTTCC GGAGC GGG', 'AAGTG YGTCC TTCCG']	['TCAAGTGYGTCC TTCCGAGCGG', 'CAAGTGYGCCT TCCGAGCGGG', 'AAGTGYGCCTT CCGAGCGGGG', 'AGTGYGCCTTC CGGAGCGGGG']	['Epidermolysis bullosa simplex, Koebner type']

			GAGCG GGG']		
NM_002292.3(LAMB2):c.961T>C (p.Cys321Arg)	121912492	LAMB2	[]	['CCTCAACYGCGA GCAGTGTGAGG']	['Nephrotic syndrome, type 5, with or without ocular abnormalities']
NM_170707.3(LMNA):c.1139T>C (p.Leu380Ser)	121912495	LMNA	['TCTYGGAGGG GAGGG CGAGG AGGAG AGG']	['TCTYGGAGGGC GAGGAGGAGAGG']	['Congenital muscular dystrophy, LMNA-related', 'not provided']
NM_001399.4(EDA):c.2T>C (p.Met1Thr)	397516659	EDA	[]	['GGCCAYGGGCT ACCCGGAGGTGG']	['Hypohidrotic X-linked ectodermal dysplasia']
NM_000493.3(COL10A1):c.1951T>C (p.Trp651Arg)	111033549	-	[]	[]	['Metaphyseal chondrodysplasia, Schmid type']
NM_000233.3(LHCGR):c.1193T>C (p.Met398Thr)	121912526	-	[]	[]	['Gonadotropin-independent familial sexual precocity']
NM_000233.3(LHCGR):c.391T>C (p.Cys131Arg)	121912527	-	[]	[]	['Leydig cell hypoplasia, partial']
NM_000493.3(COL10A1):c.1798T>C (p.Ser600Pro)	111033555	-	[]	[]	['Metaphyseal chondrodysplasia, Schmid type']
NM_000233.3(LHCGR):c.1103T>C (p.Leu368Pro)	121912533	-	[]	[]	['Gonadotropin-independent familial sexual precocity']
NM_000233.3(LHCGR):c.1627T>C (p.Cys543Arg)	121912537	-	[]	[]	['Leydig cell agenesis']
NM_000233.3(LHCGR):c.1505T>C (p.Leu502Pro)	121912538	-	[]	[]	['Leydig cell agenesis']
NM_000901.4(NR3C2):c.2771T>C (p.Leu924Pro)	121912563	NR3C2	[]	[]	['Pseudohypoadosteronism type 1 autosomal dominant']
NM_021044.2(DHH):c.485T>C (p.Leu162Pro)	111033589	DHH	[]	['GTTGCGGCGCG CCTCGCAGTGG']	['46,XY gonadal dysgenesis, complete, dhh-related']
NM_000901.4(NR3C2):c.2936T>C (p.Leu979Pro)	121912567	NR3C2	[]	[]	['Pseudohypoadosteronism type 1 autosomal dominant']
NM_000517.4(HBA2):c.2T>C (p.Met1Thr)	111033603	HBA2	[]	[]	['alpha Thalassemia']
NM_000762.5(CYP2A6):c.670T>C (p.Ser224Pro)	111033610	-	[]	[]	['Tegafur response']
NM_000660.5(TGFB1):c.241T>C (p.Tyr81His)	111033611	TGFB1	[]	[]	['Diaphyseal dysplasia']
NM_001173464.1(KIF21A):c.1067T>C (p.Met356Thr)	121912588	KIF21A	[]	[]	['Fibrosis of extraocular muscles, congenital, 1']

NM_000206.2(IL2RG):c.343T>C (p.Cys115Arg)	111033622	IL2RG	[]	['TGGCYGTCAGTTGCAAAAAAAGG']	['X-linked severe combined immunodeficiency']
NM_001041.3(SI):c.1859T>C (p.Leu620Pro)	121912613	SI	[]	['ATGCGGAGTTCAGTTTGTGG']	['Sucrase-isomaltase deficiency']
NM_016180.4(SLC45A2):c.1082T>C (p.Leu361Pro)	121912619	SLC45A2	[]	['GAGTTTCYCATCTACGAAAGAGG']	['Oculocutaneous albinism type 4', 'not provided']
NM_000552.3(VWF):c.4837T>C (p.Ser1613Pro)	61750581	VWF	[]	['CTGCCYCTGATGAGATCAAGAGG']	['von Willebrand disease, type 2a', 'not provided']
NM_000552.3(VWF):c.4883T>C (p.Ile1628Thr)	61750584	VWF	[]	[]	['von Willebrand disease, type 2a', 'not provided']
NM_000180.3(GUCY2D):c.1694T>C (p.Phe565Ser)	61749755	GUCY2D	[]	[]	['Leber congenital amaurosis 1', 'not provided']
NM_003235.4(TG):c.3733T>C (p.Cys1245Arg)	121912647	TG	[]	[]	['Iodotyrosyl coupling defect']
NM_000546.5(TP53):c.755T>C (p.Leu252Pro)	121912653	TP53	[]	['CATCCYCACCATCATCACACTGG']	['Li-Fraumeni syndrome 1']
NM_000155.3(GALT):c.350T>C (p.Phe117Ser)	111033679	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.374T>C (p.Val125Ala)	111033680	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.386T>C (p.Met129Thr)	111033683	GALT	[]	['AGGTCAYGTGCTTCCACCCCTGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000546.5(TP53):c.1031T>C (p.Leu344Pro)	121912662	TP53	[]	[]	['Li-Fraumeni syndrome 1']
NM_000155.3(GALT):c.416T>C (p.Leu139Pro)	111033687	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.452T>C (p.Val151Ala)	111033701	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.499T>C (p.Trp167Arg)	111033708	GALT	['CCCTYGGGTGCAGGT TTGTGAGG']	['CCCTYGGGTGCAGGT TTGTGAGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.507+2T>C	111033710	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.512T>C (p.Phe171Ser)	111033715	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']

NM_000341.3(SLC3A1):c.2033T>C (p.Leu678Pro)	121912693	-	[]	[]	['Cystinuria']
NM_000540.2(RYR1):c.9242T>C (p.Met3081Thr)	147012990	RYR1	[]	[]	['Minicore myopathy with external ophthalmoplegia']
NM_000155.3(GALT):c.584T>C (p.Leu195Pro)	111033728	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.650T>C (p.Leu217Pro)	111033741	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.687+2T>C	111033748	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000039.1(APOA1):c.220T>C (p.Trp74Arg)	121912726	-	[]	[]	['Familial visceral amyloidosis, Ostertag type']
NM_000155.3(GALT):c.677T>C (p.Leu226Pro)	111033752	GALT	[]	['CAGGAGCYACT CAGGAAGGTGGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000039.1(APOA1):c.593T>C (p.Leu198Ser)	121912729	APOA1	[]	['GCGCTYGGCCGC GCGCCTTGAGG']	['Familial visceral amyloidosis, Ostertag type']
NM_000155.3(GALT):c.745T>C (p.Trp249Arg)	111033757	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_001681.3(ATP2A2):c.1678T>C (p.Cys560Arg)	121912734	ATP2A2	[]	[]	['Keratosis follicularis']
NM_000041.3(APOE):c.137T>C (p.Leu46Pro)	769452	APOE	[]	['AACYGGCACTG GGTCGCTTTTGG']	[]
NM_000342.3(SLC4A1):c.2317T>C (p.Ser773Pro)	121912753	SLC4A1	[]	[]	['Renal tubular acidosis, distal, with normal red cell morphology']
NM_003002.3(SDHD):c.284T>C (p.Leu95Pro)	80338846	SDHD	[]	[]	['Hereditary Paraganglioma-Pheochromocytoma Syndromes']
NM_016124.4(RHD):c.329T>C (p.Leu110Pro)	121912762	RHD	[]	['ACACYGTTTCAGG TATTGGGATGG']	[]
NM_003002.3(SDHD):c.416T>C (p.Leu139Pro)	80338847	SDHD	[]	[]	['Hereditary Paraganglioma-Pheochromocytoma Syndromes', 'Paragangliomas 1']
NM_001822.5(CHN1):c.427T>C (p.Tyr143His)	121912794	CHN1	[]	[]	['Duane syndrome type 2']
NM_000155.3(GALT):c.1138T>C	111033824	GALT	[]	['CGCCYGACCAC GCCGACCACAGG']	['Deficiency of UDPglucose-']

(p.Ter380Arg)				'GCCYGACCACGC CGACCACAGGG']	hexose-1-phosphate uridylyltransferase']
m.3271T>C	199474658	MT-TL1	[]	[]	['Juvenile myopathy, encephalopathy, lactic acidosis AND stroke']
NM_000155.3(GALT) :c.980T>C (p.Leu327Pro)	111033832	GALT	[]	['TCCYGCGCTCTG CCACTGTCCGG']	['Deficiency of UDPglucose- hexose-1-phosphate uridylyltransferase']
m.3290T>C	199474665	MT-TL1	[]	[]	['Sudden infant death syndrome']
NM_020549.4(CHAT) :c.629T>C (p.Leu210Pro)	121912820	CHAT	[]	[]	['Familial infantile myasthenia']
NM_020549.4(CHAT) :c.1007T>C (p.Ile336Thr)	121912823	CHAT	[]	[]	['Familial infantile myasthenia']
NM_000155.3(GALT) :c.328+2T>C	111033849	GALT	[]	[]	['Deficiency of UDPglucose- hexose-1-phosphate uridylyltransferase']
NM_000267.3(NF1):c. 1595T>C (p.Leu532Pro)	199474737	NF1	[]	[]	['Neurofibromatosis, type 1', 'not provided']
NM_000455.4(STK11) :c.545T>C (p.Leu182Pro)	730881974	STK11	[]	['GGGAACCYGCT GCTCACCACCGG', 'AACCYGCTGCTC ACCACCGGTGG']	['Hereditary cancer- predisposing syndrome']
NM_001042492.2(NF 1):c.2288T>C (p.Leu763Pro)	199474762	NF1	[]	[]	['Hereditary cancer- predisposing syndrome', 'not provided']
NM_001042492.2(NF 1):c.5858T>C (p.Leu1953Pro)	199474792	NF1	[]	[]	['Neurofibromatosis, type 1', 'not provided']
m.7512T>C	199474817	MT-TS1	[]	[]	['MERRF/MELAS overlap syndrome']
m.7510T>C	199474820	MT-TS1	[]	[]	['Deafness, nonsyndromic sensorineural, mitochondrial']
m.7511T>C	199474821	MT-TS1	[]	[]	['Deafness, nonsyndromic sensorineural, mitochondrial']
m.2991T>C	199474823	MT-RNR2	[]	[]	['Chloramphenicol resistance']
NM_201253.2(CRB1): c.3122T>C (p.Met1041Thr)	62635656	CRB1	[]	[]	['Retinitis pigmentosa 12', 'not provided']
m.7587T>C	199474825	MT-CO2	[]	[]	['Cytochrome-c oxidase deficiency']
NM_000400.3(ERCC2)):c.1454T>C (p.Leu485Pro)	121913025	ERCC2	[]	[]	['Xeroderma pigmentosum, group D']
NM_000157.3(GBA):c .703T>C	1064644	GBA	[]	['GGGYCACTCAA GGGACAGCCCGG']	['Gaucher disease']

(p.Ser235Pro)]	
NM_001113755.2(TYMP):c.854T>C (p.Leu285Pro)	121913042	TYMP	[]	[]		[]
NM_000122.1(ERCC3):c.296T>C (p.Phe99Ser)	121913045	ERCC3	[]	[]		['Xeroderma pigmentosum, complementation group b']
NM_000186.3(CFH):c.1606T>C (p.Cys536Arg)	121913052	CFH	[]	[]		['Factor H deficiency']
NM_138413.3(HOGA1):c.533T>C (p.Leu178Pro)	796052090	HOGA1	[]	['GGACCYGCCTGTGGATGCAGTGG']		['Primary hyperoxaluria, type III']
NM_004370.5(COL12A1):c.7001T>C (p.Ile2334Thr)	796052093	COL12A1	[]	[]		['BETHLEM MYOPATHY 2']
NM_000043.4(FAS):c.532T>C (p.Cys178Arg)	121913084	FAS	[]	[]		[]
NM_000208.2(INSR):c.779T>C (p.Leu260Pro)	121913141	INSR	[]	['CTACCYGGACGGCAGGTGTGTGG']		['Leprechaunism syndrome']
NM_000208.2(INSR):c.164T>C (p.Val55Ala)	121913152	INSR	[]	[]		['Leprechaunism syndrome']
NM_000026.2(ADSL):c.340T>C (p.Tyr114His)	374259530	ADSL	[]	[]		['Adenylosuccinate lyase deficiency', 'not provided']
NM_153490.2(KRT13):c.356T>C (p.Leu119Pro)	60440396	KRT13	[]	[]		['White sponge nevus 2', 'not provided']
NM_022455.4(NSD1):c.5885T>C (p.Ile1962Thr)	587784162	NSD1	[]	[]		['Sotos syndrome 1']
NM_006218.2(PIK3CA):c.1258T>C (p.Cys420Arg)	121913272	PIK3CA	[]	['GGAACACYGTCATTGGCATGGG', 'GAACACYGTCCA TTGGCATGGGG']		['Congenital lipomatous overgrowth, vascular malformations, and epidermal nevi', 'Neoplasm of ovary', 'PIK3CA Related Overgrowth Spectrum']
NM_000552.3(VWF):c.8317T>C (p.Cys2773Arg)	61751310	VWF	[]	['GCTCCYGCTGCTCTCCGACACGG']		['von Willebrand disease, type 2a', 'not provided']
NM_000335.4(SCN5A):c.5504T>C (p.Ile1835Thr)	45563942	SCN5A	[]	[]		['Primary dilated cardiomyopathy', 'Dilated cardiomyopathy 1E', 'not provided']
NM_183415.2(UBE3B):c.1741+2T>C	398123020	UBE3B	[]	[]		['Kaufman oculocerebrofacial syndrome']
NM_000021.3(PSEN1):c.1175T>C (p.Leu392Pro)	63750218	PSEN1	[]	[]		['Alzheimer disease, type 3', 'not provided']
NM_000350.2(ABCA4)	61751392	ABCA4	[]	[]		['Stargardt disease']

4):c.1622T>C (p.Leu541Pro)					1', 'Cone-rod dystrophy 3', 'not provided']
NM_007313.2(ABL1): c.1109T>C (p.Met370Thr)	121913457	ABL1	[]	[]	[]
NM_024408.3(NOTCH2): c.1117T>C (p.Cys373Arg)	312262793	NOTCH2	[]	[]	['Alagille syndrome 2']
NM_024408.3(NOTCH2): c.1438T>C (p.Cys480Arg)	312262799	NOTCH2	[]	['TTCACAYGTCTGTGCATGCCAGG']	['Alagille syndrome 2']
NM_003611.2(OFD1): c.111+2T>C	312262809	OFD1	[]	[]	['Oral-facial-digital syndrome', 'not provided']
NM_003611.2(OFD1): c.274T>C (p.Ser92Pro)	312262819	OFD1	[]	[]	['Oral-facial-digital syndrome']
NM_020631.4(PLEKHG5): c.1940T>C (p.Phe647Ser)	63750315	PLEKHG5	[]	[]	['Distal spinal muscular atrophy, autosomal recessive 4']
NM_001288953.1(TTC7A): c.1912T>C (p.Ser638Pro)	149602485	TTC7A	[]	[]	['Multiple gastrointestinal atresias']
NM_000391.3(TPP1): c.1093T>C (p.Cys365Arg)	119455953	TPP1	['GCCGGGYGTGCTGTCTCTGG']	['GCCGGGYGTGCTGTCTCTGG']	['Ceroid lipofuscinosis, neuronal, 2', 'not provided']
NM_000426.3(LAMA2): c.7691T>C (p.Leu2564Pro)	121913570	LAMA2	[]	['ATCATTCYTTTGGGAAGTGGAGG', 'TCATTCYTTTGGGAAGTGGAGG']	['Merosin deficient congenital muscular dystrophy']
NM_000426.3(LAMA2): c.2584T>C (p.Cys862Arg)	121913573	LAMA2	[]	[]	['Congenital muscular dystrophy due to partial LAMA2 deficiency']
NM_000530.6(MPZ): c.404T>C (p.Ile135Thr)	121913587	MPZ	[]	[]	['Charcot-Marie-Tooth disease type 1B']
NM_201253.2(CRB1): c.3541T>C (p.Cys1181Arg)	62636291	CRB1	[]	[]	['Retinitis pigmentosa 12', 'not provided']
NM_000257.3(MYH7): c.1046T>C (p.Met349Thr)	121913640	MYH7	[]	['AACTCCAYGTATAAGCTGACAGG']	['Familial hypertrophic cardiomyopathy 1', 'Cardiomyopathy']
NM_000257.3(MYH7): c.1594T>C (p.Ser532Pro)	121913642	MYH7	[]	['CATCATGYCCATCCTGGAAGAGG']	['Dilated cardiomyopathy 1S']
NM_000257.3(MYH7): c.5378T>C (p.Leu1793Pro)	121913654	MYH7	[]	[]	['Familial hypertrophic cardiomyopathy 1', 'Myosin storage myopathy', 'Left ventricular noncompaction 5', 'Cardiomyopathy']
NM_001127500.1(MET)	121913668	MET	[]	[]	['Renal cell

T):c.3446T>C (p.Met1149Thr)					carcinoma, papillary, 1']
NM_002443.3(MSMB)):c.-89T=	10993994	MSMB	[]	[]	['Prostate cancer, hereditary, 13']
NM_001079802.1(FK TN):c.527T>C (p.Phe176Ser)	119463996	FKTN	[]	['GTAGTCTYTCAT GAGAGGAGTGG']	['Limb-girdle muscular dystrophy- dystroglycanopathy, type C4']
NM_000169.2(GLA):c .865A>T (p.Ile289Phe)	140329381	-	[]	[]	['Fabry disease']
NM_000420.2(KEL):c .1790T>C (p.Leu597Pro)	8176038	KEL	[]	[]	[]
NM_003999.2(OSMR) :c.2072T>C (p.Ile691Thr)	63750567	OSMR	[]	[]	['Primary localized cutaneous amyloidosis 1']
NC_012920.1:m.9478 T>C	587776437	MT-CO3	['TCAG AAGYT TTTTTC TTCGC AGG']	['TCAGAAGYTTTT TTCTTCGCAGG']	['Leigh disease']
NM_002775.4(HTRA1)):c.1091T>C (p.Leu364Pro)	587776447	HTRA1	[]	[]	['Cerebral autosomal recessive arteriopathy with subcortical infarcts and leukoencephalopath y']
NM_002049.3(GATA 1):c.2T>C (p.Met1Thr)	587776451	GATA1	['TCCA YGGAG TTCCCT GGCCT GGG', 'CCAYG GAGTT CCCTG GCCTG GGG']	['CTCCAYGGAGTT CCCTGGCCTGG', 'TCCAYGGAGTTC CCTGGCCTGGG', 'CCAYGGAGTTCC CTGGCCTGGGG']	['GATA-1-related thrombocytopenia with dyserythropoiesis']
NM_000021.3(PSEN1)):c.254T>C (p.Leu85Pro)	63750599	PSEN1	[]	[]	['Alzheimer disease, familial, 3, with spastic paraparesis and apraxia', 'not provided']
NM_002049.3(GATA 1):c.1240T>C (p.Ter414Arg)	587776456	GATA1	[]	['GCTCAYGAGGG CACAGAGCATGG']	['GATA-1-related thrombocytopenia with dyserythropoiesis']
NM_006158.4(NEFL): c.281T>C (p.Leu94Pro)	62636505	NEFL	[]	[]	['Charcot-Marie- Tooth disease type 2E', 'not provided']
NM_000184.2(HBG2): c.-228T>C	63750654	HBG2	[]	['ATGCAAAYATCT GTCTGAAACGG']	['Fetal hemoglobin quantitative trait locus 1']
NM_000353.2(TAT):c .236-5A>G	587776512	TAT	[]	[]	['Tyrosinemia type 2']
NM_173560.3(RFX6): c.380+2T>C	587776514	RFX6	['CAGT GGYGA GACTC GCCCC CAGG',	['CAGTGGYGAGA CTCGCCCGCAGG', 'AGTGGYGAGACT CGCCCGCAGGG']	['Mitchell-Riley syndrome']

			'AGTGG YGAGA CTCGC CCGCA GGG']		
NM_001999.3(FBN2): c.3725-15A>G	587776519	FBN2	[]	['AGCAYTGCAAC CACATTGTCAGG']	['Congenital contractural arachnodactyly']
NM_000404.2(GLB1): c.1480-2A>G	587776526	GLB1	[]	[]	['GM1- GANGLIOSIDOSIS , TYPE I, WITH CARDIAC INVOLVEMENT']
NM_000402.4(G6PD): c.473T>C (p.Leu158Pro)	78365220	G6PD	[]	['TGCCCYCCACCT GGGGTCACAGG']	['Anemia, nonspherocytic hemolytic, due to G6PD deficiency', 'not provided']
NM_000179.2(MSH6) :c.1346T>C (p.Leu449Pro)	63750741	MSH6	[]	['CTGGGGCYGGT ATTCATGAAAGG']	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_145046.4(CALR3) :c.245A>G (p.Lys82Arg)	142951029	CALR3	['CGGT YTGAA GCGTG CAGAG ATGG']	['CGGTYTGAAGC GTGCAGAGATGG']	['Arrhythmogenic right ventricular cardiomyopathy', 'Familial hypertrophic cardiomyopathy 19', 'Hypertrophic cardiomyopathy']
NM_000260.3(MYO7 A):c.5573T>C (p.Leu1858Pro)	368657015	MYO7A	[]	[]	['Usher syndrome, type 1']
NM_024753.4(TTC21 B):c.2758-2A>G	766132877	TTC21B	[]	[]	['Nephronophthisis 12']
NM_001195129.1(PR SS56):c.1183T>C (p.Cys395Arg)	730882161	PRSS56	[]	[]	['Microphthalmia, isolated 6']
NM_001184.3(ATR):c .2022A>G (p.Gly674=)	587776690	ATR	[]	[]	['Seckel syndrome 1']
NM_000354.5(SERP INA7):c.623-2A>G	587776720	SERPINA7	[]	[]	[]
NM_000133.3(F9):c.2 77+2T>C	587776735	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_004006.2(DMD): c.9225-285A>G	587776747	DMD	[]	[]	['Becker muscular dystrophy']
NM_016835.4(MAPT) :c.1839T>C (p.Asn613=)	63750912	MAPT	['GGAT AAYAT CAAAC ACGTC CCGG', 'GATAA YATCA AACAC GTCCC GGG']	['GGATAAYATCA AACACGTCCCGG', 'GATAAYATCAAA CACGTCCCGG']	['Frontotemporal dementia', 'not provided']
NM_000321.2(RB1):c. 1960+2T>C	587776780	RB1	[]	[]	['Retinoblastoma']

NM_006517.4(SLC16A2):c.1253T>C (p.Leu418Pro)	367543059	SLC16A2	[]	[]	['Allan-Herndon-Dudley syndrome']
NM_000421.3(KRT10):c.1374-2A>G	587776815	-	[]	[]	['Erythroderma, ichthyosiform, congenital reticular']
NM_000251.2(MSH2):c.2089T>C (p.Cys697Arg)	63750961	MSH2	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_002618.3(PEX13):c.977T>C (p.Ile326Thr)	61752115	PEX13	[]	[]	['Peroxisome biogenesis disorder 11B']
NM_001079867.1(PEX2):c.739T>C (p.Cys247Arg)	61752128	PEX2	[]	[]	['Peroxisome biogenesis disorder 5A']
NM_017929.5(PEX26):c.134T>C (p.Leu45Pro)	61752132	PEX26	[]	[]	['Peroxisome biogenesis disorder 7B']
NG_012088.1:g.2209A>G	587776843	IL10	['ACCY TATGA TCCGC CCGCC TTGG']	['ACCYTATGATCC GCCCGCCTTGG']	[]
NM_001735.2(C5):c.1115A>G (p.Lys372Arg)	587776846	C5	[]	[]	['Leiner disease']
NM_002087.3(GRN):c.2T>C (p.Met1Thr)	63751006	GRN	['CCAY GTGGA CCCTG GTGAG CTGG']	['CCAYGTGGACCC TGGTGAGCTGG']	['Frontotemporal dementia, ubiquitin-positive', 'not provided']
NM_004656.3(BAP1):c.2057-2A>G	587776878	BAP1	[]	[]	['Tumor predisposition syndrome']
NM_004656.3(BAP1):c.438-2A>G	587776879	BAP1	['GCCY GGGGA AAAAC AGAGT CAGG']	['GCCYGGGAAA AACAGAGTCAGG']	['Tumor predisposition syndrome']
NM_004329.2(BMPRI1A):c.370T>C (p.Cys124Arg)	199476087	BMPRI1A	[]	[]	['Juvenile polyposis syndrome', 'Hereditary cancer-predisposing syndrome']
NM_000510.2(FSHB):c.298T>C (p.Cys100Arg)	5030777	FSHB	[]	[]	['Follicle-stimulating hormone deficiency, isolated']
NM_001009944.2(PKD1):c.2534T>C (p.Leu845Ser)	199476100	PKD1	[]	[]	['Polycystic kidney disease, adult type']
NM_004963.3(GUCY2C):c.1160A>G (p.Asp387Gly)	587776905	GUCY2C	[]	[]	['Meconium ileus']
m.14487T>C	199476109	MT-ND6	[]	[]	['Leigh disease', 'Leigh syndrome due to mitochondrial complex I deficiency']

NM_017565.3(FAM20A):c.813-2A>G	587776912	-	[]	[]	['Enamel-renal syndrome']
NM_017565.3(FAM20A):c.590-2A>G	587776914	FAM20A	[]	['GTAATCYGCAA AGGAGGAGAAGG , 'TAATCYGCAAAG GAGGAGAAGGG']	['Enamel-renal syndrome']
NM_014165.3(NDUF AF4):c.194T>C (p.Leu65Pro)	63751061	NDUF4F4	[]	[]	['Mitochondrial complex I deficiency']
m.4160T>C	199476119	MT-ND1	[]	[]	['Leber optic atrophy']
NM_000551.3(VHL):c.292T>C (p.Tyr98His)	5030809	VHL	[]	['CCCYACCCAACG CTGCCGCCTGG']	['Von Hippel-Lindau syndrome', 'Hereditary cancer-predisposing syndrome']
m.3949T>C	199476124	MT-ND1	[]	[]	['Juvenile myopathy, encephalopathy, lactic acidosis AND stroke']
m.6742T>C	199476126	MT-CO1	[]	[]	[]
m.6721T>C	199476127	MT-CO1	[]	[]	[]
m.5692T>C	199476131	MT-TN	[]	[]	[]
m.5728T>C	199476132	MT-TN	[]	['CAATCYACTTCT CCCGCCGCCGG', 'AATCYACTTCTC CCGCCGCCGGG']	['Cytochrome-c oxidase deficiency', 'Mitochondrial complex I deficiency']
m.9101T>C	199476134	MT-ATP6	[]	[]	['Leber optic atrophy']
m.8851T>C	199476136	MT-ATP6	[]	[]	['Leigh disease', 'Striatonigral degeneration, infantile, mitochondrial']
m.9185T>C	199476138	MT-ATP6	[]	[]	['Leigh disease']
NM_000277.1(PAH):c.143T>C (p.Leu48Ser)	5030841	PAH	[]	[]	['Phenylketonuria', 'not provided']
NM_000277.1(PAH):c.691T>C (p.Ser231Pro)	5030845	PAH	[]	[]	['Phenylketonuria', 'not provided']
NM_000251.2(MSH2):c.595T>C (p.Cys199Arg)	63751110	MSH2	['AAGG AAYGT GTTTT ACCCG GAGG']	['AAGGAAYGTGT TTTACCCGGAGG']	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_000039.1(APOA1):c.341T>C (p.Leu114Pro)	28931575	-	[]	[]	[]
NM_001288953.1(TTC7A):c.2366T>C (p.Leu789Pro)	587776972	TTC7A	[]	[]	['Multiple gastrointestinal atresias']
NM_014336.4(AIPL1):c.715T>C (p.Cys239Arg)	62637012	AIPL1	[]	['CTGCCAGYGCCT GCTGAAGAAGG', 'CCAGYGCCTGCT GAAGAAGGAGG']	['Leber congenital amaurosis 4', 'not provided']
NM_000155.3(GALT)	367543254	GALT	[]	[]	['Deficiency of

:c.336T>C (p.Ser112=)					UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_207352.3(CYP4V2):c.655T>C (p.Tyr219His)	199476191	CYP4V2	[]	[]	['Bietti crystalline corneoretinal dystrophy']
NM_207352.3(CYP4V2):c.1021T>C (p.Ser341Pro)	199476199	CYP4V2	[]	['AAACTGGYCCTT ATACCTGTTGG', 'AACTGGYCCTTA TACCTGTTGGG']	['Bietti crystalline corneoretinal dystrophy']
NM_001142519.1(FAM111A):c.1531T>C (p.Tyr511His)	587777012	FAM111A	[]	[]	['Kenny-Caffey syndrome type 2']
NM_000435.2(NOTCH3):c.4556T>C (p.Leu1519Pro)	367543285	NOTCH3	[]	[]	['Infantile myofibromatosis 1', 'Infantile myofibromatosis 2']
NM_000021.3(PSEN1):c.749T>C (p.Leu250Ser)	63751163	PSEN1	[]	[]	['Alzheimer disease, type 3', 'not provided']
NM_001283009.1(RTEL1):c.3730T>C (p.Cys1244Arg)	587777037	-	['CTGTG TGYGC CAGGG CTGTG GGG']	['CTGTGTGYGCCA GGGCTGTGGGG']	['Dyskeratosis congenita, autosomal recessive, 5']
NM_001135021.1(ELMOD3):c.794T>C (p.Leu265Ser)	587777040	ELMOD3	[]	[]	['Deafness, autosomal recessive 88']
NM_001001557.2(GDF6):c.866T>C (p.Leu289Pro)	63751220	GDF6	[]	[]	['Klippel-Feil syndrome 1, autosomal dominant']
NM_014754.2(PTDSS1):c.794T>C (p.Leu265Pro)	587777090	PTDSS1	[]	[]	['Lenz-Majewski hyperostosis syndrome']
NM_052844.3(WDR34):c.1307A>G (p.Lys436Arg)	587777098	WDR34	[]	[]	['Short-rib thoracic dysplasia 11 with or without polydactyly']
NM_001290048.1(ATL3):c.521A>G (p.Tyr174Cys)	587777108	ATL3	[]	[]	['Hereditary sensory neuropathy type IF']
NM_001018005.1(TPM1):c.515T>C (p.Ile172Thr)	199476312	TPM1	[]	[]	['Primary familial hypertrophic cardiomyopathy', 'Cardiomyopathy', 'not provided']
NM_018849.2(ABCB4):c.523A>G (p.Thr175Ala)	58238559	ABCB4	[]	[]	['Cholecystitis']
NM_001018005.1(TPM1):c.842T>C (p.Met281Thr)	199476321	TPM1	[]	[]	['Cardiomyopathy', 'not specified', 'not provided']
NM_005763.3(AASS):c.874A>G (p.Ile292Val)	587777122	AASS	[]	[]	['Hyperlysinemia']
NM_194442.2(LBR):c.1639A>G (p.Asn547Asp)	587777171	LBR	[]	[]	['Greenberg dysplasia']

NM_006702.4(PNPLA6):c.3053T>C (p.Phe1018Ser)	587777183	PNPLA6	[]	['CCTYTAACCGCAGCATCCATCGG']	['Boucher Neuhauser syndrome']
NM_000487.5(ARSA):c.899T>C (p.Leu300Ser)	199476389	ARSA	[]	['GGTCTCTYGC GG TGTGGAAAGGG']	['Metachromatic leukodystrophy', 'not provided']
NM_016599.4(MYOZ2):c.142T>C (p.Ser48Pro)	199476398	MYOZ2	[]	['TTAYCCCATCTCAGTAACCGTGG']	['Familial hypertrophic cardiomyopathy 16', 'not provided']
NM_014740.3(EIF4A3):c.809A>G (p.Asp270Gly)	587777204	EIF4A3	[]	[]	['Richieri Costa Pereira syndrome']
NM_001040436.2(YARS2):c.1303A>G (p.Ser435Gly)	587777215	YARS2	[]	[]	['Myopathy, lactic acidosis, and sideroblastic anemia 2']
NM_001278503.1(STT3A):c.1877T>C (p.Val626Ala)	587777216	STT3A	[]	[]	['Congenital disorder of glycosylation type 1w']
NM_001037633.1(SIL1):c.1370T>C (p.Leu457Pro)	119456967	SIL1	[]	['TTGCYGAAGGAGCTGAGATGAGG']	['Marinesco-Sjogren syndrome']
NM_006888.4(CALM1):c.268T>C (p.Phe90Leu)	730882253	CALM1	[]	['GGCAYTCCGAGTCTTTGACAAGG']	['Long QT syndrome 14']
NM_001003811.1(TEX11):c.511A>G (p.Met171Val)	143246552	TEX11	['TCCAYGGTCAAGYGGTC AAGTC AGCCT CAGG']	['TCCAYGGTCAAGTCAGCCTCAGG', 'CCAYGGTCAAGTCAGCCTCAGGG']	['Spermatogenic failure, X-linked, 2']
NM_033419.4(PGAP3):c.914A>G (p.Asp305Gly)	587777252	PGAP3	[]	[]	['Hyperphosphatasia with mental retardation syndrome 4']
NM_000021.3(PSEN1):c.338T>C (p.Leu113Pro)	63751399	PSEN1	[]	[]	['Alzheimer disease, type 3', 'Frontotemporal dementia', 'not provided']
NM_000097.5(CPOX):c.980A>G (p.His327Arg)	587777271	CPOX	[]	[]	['Harderoporphyria']
NM_005654.5(NR2F1):c.755T>C (p.Leu252Pro)	587777276	NR2F1	[]	[]	['Bosch-boonstra-schaaf optic atrophy syndrome']
NM_012338.3(TSPAN12):c.413A>G (p.Tyr138Cys)	587777283	TSPAN12	[]	['TAATCCAYAATTGTCATCCTGG']	['Exudative vitreoretinopathy 5']
NM_003181.3(T):c.512A>G (p.His171Arg)	587777303	T	[]	[]	['Sacral agenesis with vertebral anomalies']
NM_015884.3(MBTPS2):c.1391T>C (p.Phe464Ser)	587777306	MBTPS2	[]	['GCTYTGCTTTGGATGGACAATGG']	['Palmoplantar keratoderma, mutilating, with periorificial keratotic plaques, X-linked']
NM_020435.3(GJC2):	74315311	GJC2	['TGAG']	['TGAGAYGGCCC']	['Leukodystrophy,

c.857T>C (p.Met286Thr)			AYGGC CCACC TGGGC TTGG']	ACCTGGGCTTGG', 'GAGAYGGCCCAC CTGGGCTTGGG']	hypomyelinating, 2']
NM_005356.4(LCK):c .1022T>C (p.Leu341Pro)	587777335	LCK	[]	[]	['Immunodeficiency 22']
NM_005861.3(STUB1 (p.Met240Thr)	587777345	-	[]	[]	['Spinocerebellar ataxia, autosomal recessive 16']
NM_000250.1(MPO):c .752T>C (p.Met251Thr)	56378716	MPO	[]	['TCACTCAYGTTC ATGCAATGGGG']	['Myeloperoxidase deficiency']
NM_017890.4(VPS13 B):c.11119+2T>C	587777382	VPS13B	[]	[]	['Cohen syndrome']
NM_005026.3(PIK3C D):c.1246T>C (p.Cys416Arg)	587777390	PIK3CD	[]	['GCAGGACYGCC CCATTGCCTGGG']	['Activated PI3K- delta syndrome']
NM_002633.2(PGM1) :c.1547T>C (p.Leu516Pro)	587777401	PGM1	[]	[]	['Congenital disorder of glycosylation type 1t']
NM_000261.1(MYOC (p.Tyr437His)	74315328	MYOC	[]	[]	['Primary open angle glaucoma juvenile onset 1']
NM_001159287.1(TPI 1):c.833T>C (p.Phe278Ser)	587777440	TPI1	[]	[]	['Triosephosphate isomerase deficiency']
NM_000414.3(HSD17 B4):c.1547T>C (p.Ile516Thr)	587777443	HSD17B4	[]	[]	['Gonadal dysgenesis with auditory dysfunction, autosomal recessive inheritance']
NM_005359.5(SMAD 4):c.970T>C (p.Cys324Arg)	377767339	SMAD4	[]	[]	['Juvenile polyposis syndrome']
NM_000211.4(ITGB2) :c.1877+2T>C	483352818	ITGB2	['CATG YGAGT GCAGG CGGAG CAGG']	['CATGYGAGTGC AGGCGGAGCAGG']	['Leukocyte adhesion deficiency type 1']
NM_005359.5(SMAD 4):c.1087T>C (p.Cys363Arg)	377767348	SMAD4	[]	[]	['Juvenile polyposis syndrome']
NM_001128159.2(VP S53):c.2084A>G (p.Gln695Arg)	587777465	VPS53	[]	[]	['Pontocerebellar hypoplasia, type 2e']
NM_000249.3(MLH1) :c.1745T>C (p.Leu582Pro)	63751616	MLH1	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_003108.3(SOX11 (p.Ser60Pro)	587777480	SOX11	[]	['TATGGYCCAAG ATCGAACGCAGG']	['Mental retardation, autosomal dominant 27']
NM_020630.4(RET):c. 1888T>C (p.Cys630Arg)	377767404	RET	[]	[]	[]
NM_017565.3(FAM20 A):c.720-2A>G	587777530	-	[]	[]	['Enamel-renal syndrome']

NM_015599.2(PGM3) :c.737A>G (p.Asn246Ser)	587777562	PGM3	['TAAA TGAYT GAGTT TGCCC TTGG']	['TAAATGAYTGA GTTTGCCCTTGG']	['Immunodeficiency 23']
NM_015599.2(PGM3) :c.1352A>G (p.Gln451Arg)	587777565	PGM3	[]	[]	['Immunodeficiency 23']
NM_000206.2(IL2RG) :c.452T>C (p.Leu151Pro)	137852511	IL2RG	[]	[]	['X-linked severe combined immunodeficiency']
NM_198282.3(TMEM 173):c.461A>G (p.Asn154Ser)	587777609	TMEM173	[]	[]	['Sting-associated vasculopathy, infantile-onset']
NM_000329.2(RPE65) :c.1022T>C (p.Leu341Ser)	61752909	RPE65	[]	[]	['Retinitis pigmentosa 20', 'not provided']
NM_001127899.3(CL CN5):c.1768T>C (p.Ser590Pro)	151340623	CLCN5	[]	[]	['Dent disease 1']
NM_005027.3(PIK3R 2):c.1202T>C (p.Leu401Pro)	587777624	PIK3R2	[]	[]	['Megalencephaly polymicrogyria- polydactyly hydrocephalus syndrome']
NM_007315.3(STAT1 (p.Gln285Arg)	587777629	STAT1	[]	[]	['Immunodeficiency 31C']
NM_139276.2(STAT3 (p.Lys392Arg)	587777648	STAT3	[]	[]	['Autoimmune disease, multisystem, infantile-onset']
NM_001037811.2(HS D17B10):c.257A>G (p.Asp86Gly)	587777651	HSD17B10	[]	[]	['2-methyl-3- hydroxybutyric aciduria']
NM_001288767.1(AR MC5):c.1928T>C (p.Leu643Pro)	587777661	ARMC5	[]	[]	['Acth-independent macronodular adrenal hyperplasia 2']
NM_001288767.1(AR MC5):c.1379T>C (p.Leu460Pro)	587777663	ARMC5	[]	['GCCCGACYGCG GGATGCTGGTGG']	['Acth-independent macronodular adrenal hyperplasia 2']
NM_007315.3(STAT1 (p.Lys673Arg)	587777704	STAT1	[]	[]	['Immunodeficiency 31a']
NM_007315.3(STAT1 (p.Lys637Glu)	587777705	STAT1	[]	[]	['Immunodeficiency 31a']
NM_014845.5(FIG4):c .50T>C (p.Leu17Pro)	587777713	FIG4	[]	[]	['Charcot-Marie- Tooth disease, type 4J']
NM_000350.2(ABCA 4):c.5819T>C (p.Leu1940Pro)	61753033	ABCA4	[]	['AAGGCYACATG AACTAACCAAGG']	['Stargardt disease', 'Stargardt disease 1', 'Cone-rod dystrophy 3', 'not provided']
NM_002972.3(SBF1): c.4768A>G (p.Thr1590Ala)	200488568	SBF1	[]	['CAGGCGYCCTCT TGCTCAGCCGG']	['Charcot-Marie- Tooth disease, type 4B3']

NM_000377.2(WAS): c.244T>C (p.Ser82Pro)	132630272	WAS	[]	[]	[]
NM_000377.2(WAS): c.809T>C (p.Leu270Pro)	132630274	WAS	[]	['CGGAGTCYGTTC TCCAGGGCAGG']	['Severe congenital neutropenia X- linked']
NM_001128834.2(PLP 1):c.487T>C (p.Trp163Arg)	132630279	PLP1	[]	[]	['Pelizaeus- Merzbacher disease', 'not provided']
NM_001128834.2(PLP 1):c.671T>C (p.Leu224Pro)	132630283	PLP1	[]	[]	['Pelizaeus- Merzbacher disease']
NM_001128834.2(PLP 1):c.560T>C (p.Ile187Thr)	132630288	PLP1	[]	[]	['Spastic paraplegia 2']
NM_001128834.2(PLP 1):c.710T>C (p.Phe237Ser)	132630291	PLP1	[]	[]	['Spastic paraplegia 2']
NM_001015877.1(PH F6):c.2T>C (p.Met1Thr)	132630300	PHF6	[]	[]	['Borjeson- Forssman-Lehmann syndrome']
NM_001399.4(EDA):c .181T>C (p.Tyr61His)	132630308	EDA	[]	['CTGCYACCTAGA GTTGCGCTCGG']	['Hypohidrotic X- linked ectodermal dysplasia']
NM_001205019.1(GK):c.1525T>C (p.Trp509Arg)	132630330	GK	[]	[]	['Deficiency of glycerol kinase']
NM_000076.2(CDKN 1C):c.*5+2T>C	587777866	CDKN1C	['CCAAGY GYGAG TACAG CGCAC CTGG', 'CAAGY GAGTA CAGCG CACCT GGG', 'AAGYG AGTAC AGCGC ACCTG GGG']	['CCAAGYGAGTA CAGCGCACCTGG', 'CAAGYGAGTACA GCGCACCTGGG', 'AAGYGAGTACAG CGCACCTGGGG']	['Beckwith- Wiedemann syndrome']
NM_000271.4(NPC1): c.2054T>C (p.Ile685Thr)	483352888	NPC1	[]	[]	['Niemann-Pick disease type C1']
NM_170707.3(LMNA):c.1589T>C (p.Leu530Pro)	60934003	LMNA	[]	['ACGGCTCYCATC AACTCCACTGG', 'CGGCTCYCATCA ACTCCACTGGG', 'GGCTCYCATCAA CTCCACTGGGG']	['Benign scapuloperoneal muscular dystrophy with cardiomyopathy', 'not provided']
NM_001165963.1(SC N1A):c.5536A>T (p.Lys1846Ter)	372098964	-	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_130838.1(UBE3A):c.2485T>C (p.Tyr829His)	587784526	UBE3A	[]	[]	['Angelman syndrome']
NM_014588.5(VSX1): c.50T>C (p.Leu17Pro)	74315436	VSX1	[]	[]	['Keratoconus 1']
NM_000404.2(GLB1):	398123354	GLB1	[]	[]	['Mucopolysaccharid

c.457+2T>C					osis, MPS-IV-B', 'Infantile GM1 gangliosidosis', 'Juvenile GM>1< gangliosidosis', 'Gangliosidosis GM1 type 3', 'not provided']
NM_003159.2(CDKL5):c.145+2T>C	267608430	CDKL5	[]	[]	['Atypical Rett syndrome', 'not provided']
NM_002764.3(PRPS1):c.869T>C (p.Ile290Thr)	180177153	PRPS1	[]	[]	['Deafness, high-frequency sensorineural, X-linked']
NM_000030.2(AGXT):c.1076T>C (p.Leu359Pro)	180177160	AGXT	[]	['GGTGCYGCGGA TCGGCCTGCTGG', 'GTGCYGCGGATC GGCCTGCTGGG']	['Primary hyperoxaluria, type I']
NM_000030.2(AGXT):c.1151T>C (p.Leu384Pro)	180177165	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_000030.2(AGXT):c.449T>C (p.Leu150Pro)	180177222	AGXT	[]	['GTGCYGCTGTTC TTAACCCACGG', 'TGCYGCTGTTCTT AACCCACGGG']	['Primary hyperoxaluria, type I']
NM_000268.3(NF2):c.1079T>C (p.Leu360Pro)	74315492	NF2	[]	[]	['Neurofibromatosis, type 2']
NM_000030.2(AGXT):c.661T>C (p.Ser221Pro)	180177254	AGXT	[]	['GCTCATCYCCTT CAGTGACAAGG']	['Primary hyperoxaluria, type I']
NM_000030.2(AGXT):c.757T>C (p.Cys253Arg)	180177264	AGXT	[]	['GGGGCYGTGAC GACCAGCCCAGG']	['Primary hyperoxaluria, type I']
NM_000030.2(AGXT):c.77T>C (p.Leu26Pro)	180177268	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_000030.2(AGXT):c.851T>C (p.Leu284Pro)	180177287	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_000030.2(AGXT):c.893T>C (p.Leu298Pro)	180177293	AGXT	[]	['GTATCYGCATGG GCGCCTGCAGG']	['Primary hyperoxaluria, type I']
NM_012203.1(GRHPR):c.203T>C (p.Leu68Pro)	180177305	GRHPR	[]	[]	['Primary hyperoxaluria, type II']
NM_000017.3(ACADS):c.1057T>C (p.Ser353Pro)	796051904	ACADS	[]	[]	['not provided']
NM_000406.2(GNRHR):c.392T>C (p.Met131Thr)	606231406	GNRHR	[]	[]	['Hypogonadotropic hypogonadism']
NM_000255.3(MUT):c.842T>C (p.Leu281Ser)	796052007	MUT	[]	[]	['not provided']
NM_000030.2(AGXT):c.947T>C (p.Leu316Pro)	796052063	AGXT	[]	[]	['Primary hyperoxaluria, type I']

NM_138413.3(HOGA1):c.875T>C (p.Met292Thr)	796052087	HOGA1	[]	[]	['Primary hyperoxaluria, type III']
NM_013382.5(POMT2):c.1997A>G (p.Tyr666Cys)	200198778	POMT2	['GGAAGYAGTGGGTGG AAGTA GAGG']	['GGAAGYAGTGG TGG AAGTAGAGG']	['Congenital muscular dystrophy', 'Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies, type A2', 'Muscular dystrophy', 'Congenital muscular dystrophy-dystroglycanopathy with mental retardation, type B2', 'not provided']
NM_015909.3(NBAS):c.3164T>C (p.Leu1055Pro)	796052121	NBAS	[]	[]	['Infantile liver failure syndrome 2']
NM_000263.3(NAGLU):c.1208T>C (p.Ile403Thr)	796052122	NAGLU	[]	[]	['Charcot-Marie-Tooth disease, axonal type 2V']
NM_203290.2(POLR1C):c.436T>C (p.Cys146Arg)	796052125	POLR1C	[]	[]	['LEUKODYSTROPHY, HYPOMYELINATING, 11']
NM_018359.3(UFSP2):c.868T>C (p.Tyr290His)	796052130	-	[]	[]	['Hip dysplasia, beukes type']
NM_000053.3(ATP7B):c.122A>G (p.Asn41Ser)	201738967	ATP7B	[]	[]	['Wilson disease']
NM_001356.4(DDX3X):c.704T>C (p.Leu235Pro)	796052224	DDX3X	[]	[]	['not provided']
NM_001356.4(DDX3X):c.1541T>C (p.Ile514Thr)	796052226	DDX3X	[]	[]	['not provided']
NM_001356.4(DDX3X):c.1175T>C (p.Leu392Pro)	796052232	DDX3X	[]	[]	['not provided']
NM_000321.2(RB1):c.2663+2T>C	587778839	RB1	[]	[]	['Retinoblastoma']
NM_000321.2(RB1):c.1472T>C (p.Leu491Pro)	587778848	RB1	[]	[]	['Retinoblastoma']
NM_006894.5(FMO3):c.1079T>C (p.Leu360Pro)	28363581	FMO3	[]	[]	['Trimethylaminuria']
NM_172107.2(KCNQ2):c.583T>C (p.Ser195Pro)	796052620	KCNQ2	[]	[]	['not provided']
NM_001282227.1(CECR1):c.1232A>G (p.Tyr411Cys)	376785840	CECR1	[]	['GAAATCAYAGG ACAAGCCTTTGG']	['Polyarteritis nodosa']
NM_170707.3(LMNA):c.644T>C	61295588	LMNA	[]	[]	['Dilated cardiomyopathy 1A']

(p.Leu215Pro)					'not provided']
NM_005249.4(FOXG1):c.673T>C (p.Trp225Arg)	796052482	FOXG1	[]	[]	['not provided']
NM_000806.5(GABRA1):c.788T>C (p.Met263Thr)	796052491	GABRA1	[]	[]	['not provided']
NM_000251.2(MSH2):c.1319T>C (p.Leu440Pro)	587779084	MSH2	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_003401.3(XRCC4):c.127T>C (p.Trp43Arg)	587779351	XRCC4	[]	[]	['Ateleiotic dwarfism']
NM_198056.2(SCN5A):c.1247A>G (p.Tyr416Cys)	372395294	SCN5A	['CTCAYAGGCCAT YAGGC CATTG CGACC ACGG']	['CTCAYAGGCCAT TGCGACCACGG']	['not provided']
NM_000257.3(MYH7):c.4835T>C (p.Leu1612Pro)	587779392	-	[]	[]	['Myopathy, distal, 1']
NM_000257.3(MYH7):c.4937T>C (p.Leu1646Pro)	587779393	-	[]	['GAGCCYCCAGA GCTTGTGAAGG']	['Myopathy, distal, 1']
NM_000404.2(GLB1):c.922T>C (p.Phe308Leu)	587779404	GLB1	[]	[]	['Infantile GM1 gangliosidosis']
NM_012434.4(SLC17A5):c.500T>C (p.Leu167Pro)	587779410	SLC17A5	[]	['ATTGTACYCAGA GCACTAGAAGG']	['Sialic acid storage disease, severe infantile type']
NM_000257.3(MYH7):c.4442T>C (p.Leu1481Pro)	587779414	-	[]	[]	['Myopathy, distal, 1']
NM_000090.3(COL3A1):c.2022+2T>C (p.Gly660_Lys674del)	587779429	COL3A1	[]	[]	['Ehlers-Danlos syndrome, type 4']
NM_004453.3(ETFDH):c.1852T>C (p.Ter618Gln)	765742496	ETFDH	[]	[]	['not provided']
NM_000090.3(COL3A1):c.2337+2T>C (p.Gly762_Lys779del)	587779513	COL3A1	[]	['AGGYAACCTTA ATACTACCTGG']	['Ehlers-Danlos syndrome, type 4']
NM_000310.3(PPT1):c.2T>C (p.Met1Thr)	796052927	PPT1	[]	[]	['not provided']
NM_020376.3(PNPLA2):c.757+2T>C	777539013	PNPLA2	[]	['GAACGGYGCGC GGACCCGGGCGG' , 'AACGGYGCGCGG ACCCGGGCGGG']	['Neutral lipid storage disease with myopathy']
NM_000090.3(COL3A1):c.3039+6T>C (p.Asp1013_Gly1014insVSSSFYSTSQ)	587779532	COL3A1	[]	[]	['Ehlers-Danlos syndrome, type 4']
NM_012452.2(TNFRSF13B):c.310T>C (p.Cys104Arg)	34557412	TNFRSF13B	[]	['ACTTCYGTGAGA ACAAGCTCAGG']	['Immunoglobulin A deficiency 2', 'Common variable immunodeficiency 2']

NM_000570.4(FCGR3B):c.244A=(p.Asn82=)	147574249	FCGR3B	[]	[]	[]
NM_001165963.1(SCN1A):c.1094T>C (p.Phe365Ser)	796052970	SCN1A	[]	['CAAGCTYTGATACCTTCAGTTGG', 'AAGCTYTGATACCTTCAGTTGGG']	['not provided']
NC_012920.1:m.7505T>C	724159989	MT-TS1	[]	['CCTCCAYGACTTTTCAAAAAGG']	['Deafness, nonsyndromic sensorineural, mitochondrial']
NM_000663.4(ABAT):c.1433T>C (p.Leu478Pro)	724159991	ABAT	[]	[]	['Gamma-aminobutyric acid transaminase deficiency']
NM_153818.1(PEX10):c.890T>C (p.Leu297Pro)	724160000	PEX10	[]	[]	['Peroxisome biogenesis disorder 6B']
NM_153818.1(PEX10):c.2T>C (p.Met1Thr)	724160002	PEX10	[]	[]	['Peroxisome biogenesis disorder 6B']
NM_000090.3(COL3A1):c.4399T>C (p.Ter1467Gln)	587779618	COL3A1	[]	[]	['Ehlers-Danlos syndrome, type 4']
NM_002485.4(NBN):c.511A>G (p.Ile171Val)	61754966	NBN	[]	[]	['Microcephaly, normal intelligence and immunodeficiency', 'Aplastic anemia', 'Hereditary cancer-predisposing syndrome', 'Leukemia, acute lymphoblastic, susceptibility to', 'not specified', 'not provided']
NM_000090.3(COL3A1):c.2553+2T>C (p.Gly816_Ala851del)	587779684	COL3A1	[]	[]	['Ehlers-Danlos syndrome, type 4']
NM_021007.2(SCN2A):c.4308+2T>C	796053139	SCN2A	['CGAATGYAAGTCTAGTTA GAGG', 'GAAATGYAAGTCTAGTTAGAGGG']	['CGAATGYAAGTCTAGTTAGAGG', 'GAAATGYAAGTCTAGTTAGAGGG']	['not provided']
NM_021007.2(SCN2A):c.4718T>C (p.Leu1573Pro)	796053152	SCN2A	[]	[]	['not provided']
NM_001101.3(ACTB):c.356T>C (p.Met119Thr)	587779773	ACTB	['GAGAAGAYGACCCCA GGTGAGTGG']	['GAGAAGAYGACCCAGGTGAGTGG']	['Baraitser-Winter syndrome 1']
NM_021007.2(SCN2A):c.2306T>C	796053191	SCN2A	[]	[]	['not provided']

(p.Ile769Thr)					
NM_014191.3(SCN8A):c.4889T>C (p.Leu1630Pro)	796053222	SCN8A	[]	['CGTCYGATCAAA GGCGCCAAAGG', 'GTCYGATCAAAG GCGCCAAAGGG']	['not provided']
NM_012415.3(RAD54B):c.1778A>G (p.Asn593Ser)	114216685	RAD54B	[]	[]	['Malignant lymphoma, non-Hodgkin']
NM_007215.3(POLG2):c.1105A>G (p.Arg369Gly)	201936720	POLG2	[]	[]	['Autosomal dominant progressive external ophthalmoplegia with mitochondrial DNA deletions 4', 'not specified']
NM_004006.2(DMD):c.6982A>T (p.Lys2328Ter)	754896795	DMD	['GCTTT TYTTC AAGCT GCCCA AGG']	['GCTTTTYTTCAA GCTGCCCAAGG']	['Duchenne muscular dystrophy', 'Becker muscular dystrophy', 'Dilated cardiomyopathy 3B']
NM_001061.4(TBXAS1):c.248T>C (p.Leu83Pro)	140005285	TBXAS1	[]	[]	['not provided']
NM_000540.2(RYR1):c.10817T>C (p.Leu3606Pro)	118192127	RYR1	[]	['TACTACCYGGAC CAGGTGGGTGG', 'ACTACCYGGACC AGGTGGGTGGG', 'CTACCYGGACCA GGTGGGTGGG']	['Central core disease', 'not provided']
NM_000138.4(FBN1):c.7754T>C (p.Ile2585Thr)	727503054	FBN1	[]	[]	['Thoracic aortic aneurysms and aortic dissections', 'Marfan syndrome']
NM_001128227.2(GNE):c.2228T>C (p.Met743Thr)	28937594	GNE	[]	[]	['Inclusion body myopathy 2', 'Nonaka myopathy']
NM_000540.2(RYR1):c.14693T>C (p.Ile4898Thr)	118192170	RYR1	[]	['AGGCAYTGGGG ACGAGATCGAGG']	['Malignant hyperthermia susceptibility type 1', 'Central core disease', 'not provided']
NM_005247.2(FGF3):c.466T>C (p.Ser156Pro)	121917703	FGF3	[]	['GTACGTGYCTGT GAACGGCAAGG', 'TACGTGYCTGTG AACGGCAAGGG']	['Deafness with labyrinthine aplasia microtia and microdontia (LAMM)']
NM_005247.2(FGF3):c.17T>C (p.Leu6Pro)	121917706	FGF3	[]	[]	['Deafness with labyrinthine aplasia microtia and microdontia (LAMM)']
NM_005211.3(CSF1R):c.2450T>C (p.Leu817Pro)	690016549	CSF1R	[]	['CCGCCYGCCTGT GAAGTGGATGG']	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_005211.3(CSF1R):c.2480T>C (p.Ile827Thr)	690016550	CSF1R	[]	[]	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_005211.3(CSF1R)	690016552	CSF1R	[]	['GAATCCCYACCC']	['Hereditary diffuse

);c.2566T>C (p.Tyr856His)				TGGCATCCTGG']	leukoencephalopathy with spheroids']
NM_001098668.2(SFTPA2):c.593T>C (p.Phe198Ser)	121917738	SFTPA2	[]	['GGAGACTYCCGCTACTCAGATGG', 'GAGACTYCCGCTACTCAGATGGG']	['Idiopathic fibrosing alveolitis, chronic form']
NM_005211.3(CSF1R):c.1957T>C (p.Cys653Arg)	690016559	CSF1R	[]	['AGCCYGTACCCA TGGAGGTAAGG', 'GCCYGTACCCATGGAGGTAAGGG']	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_005211.3(CSF1R):c.2717T>C (p.Ile906Thr)	690016560	CSF1R	[]	['GCAGAYCTGCTCCTTCCTCAGG']	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_000540.2(RYR1):c.14378T>C (p.Leu4793Pro)	118192179	RYR1	[]	[]	['Central core disease', 'not provided']
NM_199292.2(TH):c.707T>C (p.Leu236Pro)	121917763	TH	[]	[]	['Segawa syndrome, autosomal recessive']
NM_000256.3(MYBPC3):c.2994+2T>C	727503176	MYBPC3	[]	[]	['Familial hypertrophic cardiomyopathy 4']
NM_003361.3(UMOD):c.376T>C (p.Cys126Arg)	121917769	UMOD	[]	['GGCCACAYGTGTCAATGTGGTGG', 'GCCACAYGTGTC AATGTGGTGGG']	['Familial juvenile gout']
NM_003361.3(UMOD):c.943T>C (p.Cys315Arg)	121917773	UMOD	[]	['ATGGCACYGCCAGTGCAAACAGG']	['Glomerulocystic kidney disease with hyperuricemia and isosthenuria']
NM_024649.4(BBS1):c.1553T>C (p.Leu518Pro)	121917778	-	[]	[]	['Bardet-Biedl syndrome 1']
NM_172107.2(KCNQ2):c.2T>C (p.Met1Thr)	118192186	KCNQ2	[]	[]	['Benign familial neonatal seizures 1']
NM_014324.5(AMACR):c.154T>C (p.Ser52Pro)	121917814	-	[]	[]	['Alpha-methylacyl-CoA racemase deficiency', 'Bile acid synthesis defect, congenital, 4']
NM_014324.5(AMACR):c.320T>C (p.Leu107Pro)	121917816	-	[]	[]	['Bile acid synthesis defect, congenital, 4']
NM_007255.2(B4GALT7):c.617T>C (p.Leu206Pro)	121917818	B4GALT7	[]	['TGCYCTCCAAGCAGCACTACCGG']	['Ehlers-Danlos syndrome progeroid type']
NM_021615.4(CHST6):c.827T>C (p.Leu276Pro)	121917824	CHST6	[]	['GGACCYGGCGCGGGAGCCGCTGG']	['Macular corneal dystrophy Type I']
NM_006261.4(PROP1):c.263T>C (p.Phe88Ser)	121917841	PROP1	[]	[]	['Pituitary hormone deficiency, combined 2']
NM_000452.2(SLC10A2):c.728T>C (p.Leu243Pro)	121917848	SLC10A2	[]	['TTTCYTCTGGCTAGAATTGCTGG']	['Bile acid malabsorption, primary']
NM_000322.4(PRP2):c.637T>C (p.Cys213Arg)	61755802	PRP2	[]	[]	['Patterned dystrophy of retinal pigment epithelium',

					'not provided', 'Leber congenital amaurosis 18']
NM_000517.4(HBA2):c.89T>C (p.Leu30Pro)	41341344	HBA2	[]	[]	['Hemoglobin H disease, nondeletional']
NM_002181.3(IHH):c.569T>C (p.Val190Ala)	121917857	IHH	[]	[]	['Acrocapitofemoral dysplasia']
NM_000322.4(PRPH2):c.736T>C (p.Trp246Arg)	61755817	PRPH2	['ACCTGYGGGTGC GTGGCTGCAGG', 'CCTGYGGGTGCG TGGCTGCAGGG']	['ACCTGYGGGTGC GTGGCTGCAGG', 'CCTGYGGGTGCG TGGCTGCAGGG']	['Retinitis pigmentosa', 'not provided']
NM_005413.3(SIX3):c.749T>C (p.Val250Ala)	121917880	SIX3	[]	[]	['Holoprosencephaly 2']
NM_000124.3(ERCC6):c.2960T>C (p.Leu987Pro)	121917905	ERCC6	['TGCYAAAAGACCCCAAAACA AAGG']	['TGCYAAAAGACCCAAAACA AAGG']	['Cerebro-oculo-facio-skeletal syndrome']
NM_006920.4(SCN1A):c.4729T>C (p.Cys1577Arg)	121917919	-	[]	[]	['Severe myoclonic epilepsy in infancy', 'not provided']
NM_006920.4(SCN1A):c.5113T>C (p.Cys1705Arg)	121917926	-	[]	[]	['Severe myoclonic epilepsy in infancy', 'not provided']
NM_006920.4(SCN1A):c.3577T>C (p.Trp1193Arg)	121917930	-	['AACAAYGGT GGAACCTGAG AAGG']	['AACAAYGGTGG AACCTGAGAAGG']	['Generalized epilepsy with febrile seizures plus, type 1', 'Generalized epilepsy with febrile seizures plus, type 2']
NM_006920.4(SCN1A):c.838T>C (p.Trp280Arg)	121917938	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_000478.4(ALPL):c.1306T>C (p.Tyr436His)	121918006	ALPL	[]	['TGGACYATGGTGAGACCTCCAGG']	['Infantile hypophosphatasia']
NM_000478.4(ALPL):c.979T>C (p.Phe327Leu)	121918010	ALPL	[]	['CAAAGGCYTCTTCTTGCTGGTGG', 'GGCYTCTTCTTGC TGGTGAAGG']	['Infantile hypophosphatasia']
NM_000301.3(PLG):c.1771T>C (p.Ser591Pro)	121918029	PLG	[]	[]	['Dysplasminogenemia']
NM_000174.4(GP9):c.20T>C (p.Leu7Pro)	121918038	GP9	[]	[]	['Bernard-Soulier syndrome type C']
NM_000371.3(TTR):c.224T>C (p.Leu75Pro)	121918079	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis']
NM_000371.3(TTR):c.88T>C (p.Cys30Arg)	121918083	TTR	[]	[]	['Amyloidogenic transthyretin']

					amyloidosis', 'Cardiomyopathy']
NM_000371.3(TTR):c.272T>C (p.Val91Ala)	121918084	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis']
NM_000371.3(TTR):c.400T>C (p.Tyr134His)	121918088	TTR	[]	['CCCCYACTCCTA TTCCACCACGG']	[]
NM_012275.2(IL36RN):c.115+6T>C	148755083	IL36RN	[]	[]	['Pustular psoriasis, generalized']
NM_000371.3(TTR):c.95T>C (p.Leu32Pro)	121918094	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis']
NM_000371.3(TTR):c.265T>C (p.Tyr89His)	121918100	TTR	[]	[]	['AMYLOIDOSIS, LEPTOMENINGEAL, TRANSTHYRETIN-RELATED']
NM_001042465.1(PSAP):c.1055T>C (p.Leu352Pro)	121918110	PSAP	[]	['GAAGCYGCCGA AGTCCCTGTCCG']	['Gaucher disease, atypical, due to saposin C deficiency']
NM_013251.3(TAC3):c.269T>C (p.Met90Thr)	121918123	TAC3	[]	[]	['not provided']
NM_199069.1(NDUFAF3):c.2T>C (p.Met1Thr)	121918136	NDUFAF3	[]	[]	['Mitochondrial complex I deficiency']
NM_003730.4(RNASET2):c.550T>C (p.Cys184Arg)	121918137	RNASET2	[]	['CCAGYGCCTTCC ACCAAGCCAGG']	['Leukoencephalopathy, cystic, without megalencephaly']
NM_203395.2(IYD):c.347T>C (p.Ile116Thr)	121918139	IYD	[]	[]	['Iodotyrosine deiodination defect']
NM_001127628.1(FBP1):c.581T>C (p.Phe194Ser)	121918191	FBP1	[]	['GGAGTYCATTTT GGTGGACAAGG']	['Fructose-biphosphatase deficiency']
NM_015506.2(MMACHC):c.347T>C (p.Leu116Pro)	121918240	MMACHC	[]	[]	['Methylmalonic acidemia with homocystinuria']
NM_000255.3(MUT):c.313T>C (p.Trp105Arg)	121918249	MUT	[]	[]	['METHYLMALONIC ACIDURIA, mut(0) TYPE']
NM_022370.3(ROBO3):c.14T>C (p.Leu5Pro)	121918275	ROBO3	[]	[]	['Gaze palsy, familial horizontal, with progressive scoliosis']
NM_018400.3(SCN3B):c.29T>C (p.Leu10Pro)	121918282	SCN3B	[]	[]	['Brugada syndrome', 'Brugada syndrome 7', 'Cardiac arrhythmia', 'Atrial fibrillation, familial, 16']
NM_004183.3(BEST1):c.122T>C (p.Leu41Pro)	121918288	BEST1	[]	[]	['Bestrophinopathy, autosomal recessive', 'not provided']
NM_020166.4(MCCC1):c.640-2A>G	772395858	MCCC1	[]	[]	['3 Methylcrotonyl-CoA carboxylase 1 deficiency']
NM_004817.3(TJP2):c.143T>C (p.Val48Ala)	121918299	TJP2	[]	[]	['Hypercholanemia, familial']

NM_006946.2(SPTBN2):c.758T>C (p.Leu253Pro)	121918306	SPTBN2	[]	['ACCAAGCYGCTGGATCCCCGAAGG', 'AAGCYGCTGGATCCCCGAAGGTGG', 'AGCYGCTGGATCCCCGAAGGTGGG']	['Spinocerebellar ataxia 5']
NM_000214.2(JAG1):c.110T>C (p.Leu37Ser)	121918352	JAG1	[]	[]	['Alagille syndrome 1']
NM_007194.3(CHEK2):c.470T>C (p.Ile157Thr)	17879961	CHEK2	[]	[]	['Familial cancer of breast', 'Hereditary cancer-predisposing syndrome', 'Li-Fraumeni syndrome 2', 'not specified']
NM_015384.4(NIPBL):c.7637T>C (p.Leu2546Pro)	727503772	NIPBL	[]	[]	['Cornelia de Lange syndrome 1']
NM_001083112.2(GPD2):c.1904T>C (p.Phe635Ser)	121918407	GPD2	['AAGTYTGATGCA GACCAGAAAGG']	['AAGTYTGATGCA GACCAGAAAGG']	['Diabetes mellitus type 2']
NM_021957.3(GYS2):c.1447T>C (p.Ser483Pro)	121918424	GYS2	[]	[]	['Hypoglycemia with deficiency of glycogen synthetase in the liver']
NM_018849.2(ABCB4):c.1207T>C (p.Tyr403His)	121918443	ABCB4	[]	[]	['Progressive familial intrahepatic cholestasis 3']
NM_000212.2(ITGB3):c.2332T>C (p.Ser778Pro)	121918447	-	[]	[]	['Glanzmann thrombasthenia']
NM_000506.3(F2):c.1139T>C (p.Met380Thr)	121918481	F2	[]	[]	['Hereditary factor II deficiency disease']
NM_000141.4(FGFR2):c.1018T>C (p.Tyr340His)	121918489	FGFR2	['TGGGGAAYATACGTGCTTGCGCGG', 'GGGGAAYATACGTGCTTGCGGG']	['TGGGGAAYATACGTGCTTGCGCGG', 'GGGGAAYATACGTGCTTGCGGG']	['Crouzon syndrome']
NM_000141.4(FGFR2):c.799T>C (p.Ser267Pro)	121918505	FGFR2	[]	['AATGCCYCCACAGTGGTCGGAGG']	['Pfeiffer syndrome', 'Neoplasm of stomach']
NM_002739.3(PRKG):c.355T>C (p.Ser119Pro)	121918512	PRKCG	[]	[]	['Spinocerebellar ataxia 14']
NM_002739.3(PRKG):c.1927T>C (p.Phe643Leu)	121918516	PRKCG	[]	[]	['Spinocerebellar ataxia 14']
NM_015107.2(PHF8):c.836T>C (p.Phe279Ser)	121918524	PHF8	[]	[]	['Siderius X-linked mental retardation syndrome']
NM_000311.3(PRNP):c.593T>C	74315405	PRNP	[]	[]	['Gerstmann-Straussler-Scheinker']

(p.Phe198Ser)					syndrome', 'Genetic prion diseases']
NM_015665.5(AAAS):c.787T>C (p.Ser263Pro)	121918550	AAAS	[]	[]	['Glucocorticoid deficiency with achalasia']
NM_003018.3(SFTPC):c.581T>C (p.Leu194Pro)	121918560	SFTPC	[]	[]	['Surfactant metabolism dysfunction, pulmonary, 2']
NM_000322.4(PRPH2):c.554T>C (p.Leu185Pro)	121918563	PRPH2	[]	[]	['Patterned dystrophy of retinal pigment epithelium', 'Retinitis pigmentosa 7, digenic', 'not provided', 'Leber congenital amaurosis 18']
NM_000322.4(PRPH2):c.2T>C (p.Met1Thr)	121918565	PRPH2	[]	[]	['Macular dystrophy, vitelliform, adult-onset', 'not provided']
NM_001035.2(RYR2):c.1298T>C (p.Leu433Pro)	121918602	RYR2	[]	[]	['Arrhythmogenic right ventricular cardiomyopathy, type 2', 'Catecholaminergic polymorphic ventricular tachycardia', 'Long QT syndrome']
NM_001199138.1(NLRC4):c.1022T>C (p.Val341Ala)	587781260	NLRC4	[]	[]	['Syndrome of enterocolitis and autoinflammation caused by mutation of NLRC4 (SCAN4)', 'Autoinflammation with infantile enterocolitis']
NM_000702.3(ATP1A2):c.857T>C (p.Ile286Thr)	121918617	ATP1A2	[]	[]	['Familial hemiplegic migraine type 2']
NM_006920.4(SCN1A):c.4250T>C (p.Val1417Ala)	121918627	-	[]	[]	['Generalized epilepsy with febrile seizures plus, type 1', 'Generalized epilepsy with febrile seizures plus, type 2']
NM_006920.4(SCN1A):c.434T>C (p.Met145Thr)	121918631	SCN1A	[]	[]	['Generalized epilepsy with febrile seizures plus, type 2']
NM_006920.4(SCN1A):c.4462T>C (p.Phe1488Leu)	121918632	-	[]	[]	['Familial hemiplegic migraine type 3']
NM_003126.2(SPTA1):c.781T>C (p.Ser261Pro)	121918636	SPTA1	[]	[]	['Elliptocytosis 2']

NM_003126.2(SPTA1):c.620T>C (p.Leu207Pro)	121918643	SPTA1	[]	['GTGGAGCYGGT AGCTAAAGAAGG', 'TGGAGCYGGTAG CTAAAGAAGGG']	['Hereditary pyropoikilocytosis', 'Elliptocytosis 2']
NM_001024858.2(SPTB):c.604T>C (p.Trp202Arg)	121918646	SPTB	[]	['CTCCAGCYGGA AGGATGGCTTGG']	['Spherocytosis type 2']
NM_001024858.2(SPTB):c.6055T>C (p.Ser2019Pro)	121918648	SPTB	[]	['ATGCCYCTGTGG CTGAGGCGTGG']	[]
NM_001128177.1(THRB):c.929T>C (p.Met310Thr)	121918699	THRB	[]	[]	[]
NM_001128177.1(THRB):c.1373T>C (p.Val458Ala)	121918704	THRB	[]	[]	['Thyroid hormone resistance, generalized, autosomal recessive']
NM_000421.3(KRT10):c.482T>C (p.Leu161Ser)	60118264	-	[]	[]	['Bullous ichthyosiform erythroderma', 'not provided']
NM_006920.4(SCN1A):c.269T>C (p.Phe90Ser)	121918733	SCN1A	['ACTTY TATAG TATTG AATAA AGG', 'CTTYT ATAGT ATTGA ATAAA GGG']	['ACTTYTATAGTA TTGAATAAAGG', 'CTTYTATAGTATT GAATAAAGGG']	['Severe myoclonic epilepsy in infancy']
NM_006920.4(SCN1A):c.272T>C (p.Ile91Thr)	121918734	SCN1A	['ACTTT TAYAG TATTG AATAA AGG', 'CTTTT AYAGT ATTGA ATAAA GGG']	['ACTTTTAYAGTA TTGAATAAAGG', 'CTTTTAYAGTATT GAATAAAGGG']	['Severe myoclonic epilepsy in infancy']
NM_006920.4(SCN1A):c.382T>C (p.Leu1276Pro)	121918740	-	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_000543.4(SMPD1):c.475T>C (p.Cys159Arg)	727504166	SMPD1	[]	['TGAGGCCYGTG GCCTGCTCCTGG', 'GAGGCCYGTGGC CTGCTCCTGGG']	['Niemann-Pick disease, type A', 'Niemann-Pick disease, type B']
NM_006920.4(SCN1A):c.568T>C (p.Trp190Arg)	121918773	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_006920.4(SCN1A):c.552T>C (p.Met1841Thr)	121918783	-	[]	[]	['Severe myoclonic epilepsy in infancy', 'Generalized epilepsy with febrile seizures plus, type 1']

NM_002538.3(OCLN):c.656T>C (p.Phe219Ser)	267606926	OCLN	[]	[]	['Band-like calcification with simplified gyration and polymicrogyria']
NM_000434.3(NEU1):c.1088T>C (p.Leu363Pro)	193922915	NEU1	[]	['CAGCYATGGCCAGGCCCCAGTGG']	['Sialidosis, type II']
NM_198578.3(LRRK2):c.6059T>C (p.Ile2020Thr)	35870237	LRRK2	[]	[]	['Parkinson disease 8, autosomal dominant']
NM_000501.3(ELN):c.889+2T>C	727504419	ELN	[]	['CAGGYAACATCTGTCCCAGCAGG', 'AGGYAACATCTGTCCCAGCAGGG']	['Supravalvar aortic stenosis']
NM_001085.4(SERPINA3):c.233T>C (p.Leu78Pro)	1800463	SERPINA3	[]	[]	[]
NM_000238.3(KCNH2):c.1736T>C (p.Met579Thr)	199473425	KCNH2			['Congenital long QT syndrome']
NM_001211.5(BUB1B):c.3035T>C (p.Leu1012Pro)	28989185	-	[]	[]	['Mosaic variegated aneuploidy syndrome', 'Premature chromatid separation trait']
NM_000256.3(MYBPC3):c.26-2A>G	376395543	MYBPC3	[]	['GAGACYGAAGGGCCAGGTGGAGG']	['Primary familial hypertrophic cardiomyopathy', 'Familial hypertrophic cardiomyopathy 4', 'Cardiomyopathy']
NM_000051.3(ATM):c.4776+2T>C	587781927	ATM	[]	[]	['Ataxia-telangiectasia syndrome', 'Hereditary cancer-predisposing syndrome']
NM_006412.3(AGPAT2):c.589-2A>G	116807569	AGPAT2	[]	[]	['Congenital generalized lipodystrophy type 1']
NM_000545.6(HNF1A):c.1720G>A (p.Gly574Ser)	1169305	HNF1A	[]	['GATGCYGGCAGGGTCCTGGCTGG', 'ATGCYGGCAGGGTCCTGGCTGGG', 'TGCYGGCAGGGTCCTGGCTGGGG']	['Maturity-onset diabetes of the young, type 3']
NM_024514.4(CYP2R1):c.296T>C (p.Leu99Pro)	61495246	CYP2R1	[]	[]	['Vitamin d hydroxylation-deficient rickets, type 1b']
NM_012213.2(MLYCD):c.119T>C (p.Met40Thr)	28937908	MLYCD	[]	[]	['Deficiency of malonyl-CoA decarboxylase']
NM_001101.3(ACTB):c.224T>C (p.Ile75Thr)	587779771	ACTB	[]	[]	['Baraitser-Winter syndrome 1']

NM_021007.2(SCN2A):c.1271T>C (p.Val424Ala)	796053181	SCN2A	['TGTG GYGGC CATGG CCTAT GAGG']	['TGTGGYGGCCAT GGCCTATGAGG']	['not provided']
NM_002880.3(RAF1):c.769T>C (p.Ser257Pro)	727505017	RAF1	[]	[]	['Rasopathy', 'not specified']
NM_000527.4(LDLR):c.1468T>C (p.Trp490Arg)	730880130	LDLR	[]	['CTACYGGACCG ACTCTGTCCTGG', 'TACYGGACCGAC TCTGTCCTGGG']	['Familial hypercholesterolemia']
NM_170707.3(LMNA):c.710T>C (p.Phe237Ser)	730880132	LMNA	['TGAG TYTGA GAGCC GGCTG GCGG']	['TGAGTYTGAGA GCCGGCTGGCGG']	['Primary dilated cardiomyopathy']
NM_000080.3(CHRNE):c.223T>C (p.Trp75Arg)	193919341	-	[]	[]	['MYASTHENIC SYNDROME, CONGENITAL, 4B, FAST-CHANNEL']
NM_005211.3(CSF1R):c.2297T>C (p.Met766Thr)	281860270	CSF1R	[]	[]	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_005211.3(CSF1R):c.2381T>C (p.Ile794Thr)	281860274	CSF1R	['CAAG AYTGG GGACT TCGGG CTGG']	['CAAGAYTGGGG ACTTCGGGCTGG']	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_005211.3(CSF1R):c.2546T>C (p.Phe849Ser)	281860277	CSF1R	[]	[]	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_005211.3(CSF1R):c.2624T>C (p.Met875Thr)	281860279	CSF1R	[]	[]	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_018713.2(SLC30A10):c.266T>C (p.Leu89Pro)	281860284	SLC30A10	[]	[]	['Hypermannesemia with dystonia, polycythemia and cirrhosis']
NM_018713.2(SLC30A10):c.500T>C (p.Phe167Ser)	281860286	SLC30A10	[]	['GGCGCTTYCGGG GGCCTCAGGG']	['Hypermannesemia with dystonia, polycythemia and cirrhosis']
NM_018713.2(SLC30A10):c.1046T>C (p.Leu349Pro)	281860291	SLC30A10	[]	[]	['Hypermannesemia with dystonia, polycythemia and cirrhosis']
NM_016218.2(POLK):c.2287T>A (p.Tyr763Asn)	772307321	POLK	[]	[]	['Malignant tumor of prostate']
NM_000570.4(FCGR3B):c.194A>G (p.Asn65Ser)	448740	FCGR3B	[]	[]	[]
NM_030653.3(DDX11):c.2271+2T>C	730880279	DDX11	['TCCA GGYGC GGGCG TCATG CTGG']	['TCCAGGYGCGG GCGTCATGCTGG', 'CCAGGYGCGGGC GTCATGCTGGG']	['Warsaw breakage syndrome']
NM_145693.2(LPIN1)	730880306	LPIN1	[]	['AAGGYACCGCG']	['Myoglobinuria,

:c.1441+2T>C				GGCCTCGCGCGG', 'AGGYACCGCGGG CCTCGCGCGGG']	acute recurrent, autosomal recessive']
NM_002546.3(TNFRSF11B):c.226A>C (p.Thr76Pro)	200071478	TNFRSF11B	[]	[]	['Hyperphosphatase mia with bone disease']
NM_000166.5(GJB1): c.145T>C (p.Ser49Pro)	116840817	GJB1	[]	[]	['X-linked hereditary motor and sensory neuropathy']
NM_005159.4(ACTC1)):c.755T>C (p.Ile252Thr)	730880398	-	[]	[]	['Cardiomyopathy']
NM_020166.4(MCCC1):c.205A>T (p.Lys69Ter)	147741073	MCCC1	[]	[]	['3 Methylcrotonyl- CoA carboxylase 1 deficiency']
NM_000454.4(SOD1): c.338T>C (p.Ile113Thr)	74315452	SOD1	[]	['TTGCAYCATTGG CCGCACACTGG']	['Amyotrophic lateral sclerosis type 1']
NM_000169.2(GLA):c. 41T>C (p.Leu14Pro)	730880455	-	[]	['CGCGCYTGCGCT TCGCTTCCTGG']	['not provided']
NM_152743.3(BRAT1)):c.176T>C (p.Leu59Pro)	727505363	BRAT1	[]	[]	['Rigidity and multifocal seizure syndrome, lethal neonatal']
NM_005633.3(SOS1): c.2104T>C (p.Tyr702His)	727505381	SOS1	[]	[]	['Noonan syndrome', 'Rasopathy']
m.1095T>C	267606618	MT-RNR1	[]	[]	['Aminoglycoside- induced deafness', 'Auditory neuropathy', 'Deafness, nonsyndromic sensorineural, mitochondrial', 'not specified']
m.1291T>C	267606620	MT-RNR1	[]	[]	['Deafness, nonsyndromic sensorineural, mitochondrial']
NM_020247.4(ADCK3):c.1398+2T>C	606231138	ADCK3	[]	[]	['Coenzyme Q10 deficiency, primary, 4']
NM_000256.3(MYBPC3):c.467T>C (p.Leu156Pro)	730880616	MYBPC3	[]	[]	['Cardiomyopathy']
NM_022458.3(LMBR1):c.423+4842T>C	606231149	LMBR1	[]	[]	['Triphalangeal thumb', 'Preaxial polydactyly 2']
NM_022458.3(LMBR1):c.423+4808T>C	606231152	LMBR1	[]	[]	['Triphalangeal thumb', 'Preaxial polydactyly 2']
NM_021102.3(SPINT2):c.337+2T>C	606231155	SPINT2	[]	[]	['Diarrhea 3, secretory sodium, congenital, syndromic']
NM_001004127.2(ALG11):c.257T>C (p.Leu86Ser)	267606652	ALG11	[]	[]	['Congenital disorder of glycosylation type 1P']

NM_054027.4(ANKH) :c.1015T>C (p.Cys339Arg)	267606656	ANKH	[]	['AGCTCYGTTTCG TGATGTTTTGG']	['Craniometaphyseal dysplasia, autosomal dominant']
NM_054027.4(ANKH) :c.1172T>C (p.Leu391Pro)	267606658	-	[]	[]	['Craniometaphyseal dysplasia, autosomal dominant']
NM_175073.2(APTX): c.668T>C (p.Leu223Pro)	267606665	APTX	[]	[]	['Adult onset ataxia with oculomotor apraxia']
NM_004183.3(BEST1)):c.704T>C (p.Val235Ala)	267606679	BEST1	['CACT GGYGT ATACA CAGGT GAGG']	['CACTGGYGTATA CACAGGTGAGG']	['Vitreoretinchoroid opathy dominant']
NM_004183.3(BEST1)):c.614T>C (p.Ile205Thr)	267606680	BEST1	[]	[]	['Retinitis pigmentosa 50']
NM_033409.3(SLC52 A3):c.670T>C (p.Phe224Leu)	267606685	SLC52A3	[]	[]	['Brown-Vialetto- Van laere syndrome']
NM_033409.3(SLC52 A3):c.1238T>C (p.Val413Ala)	267606687	SLC52A3	[]	['AGTTACGYCAA GGTGATGCTGGG']	['Brown-Vialetto- Van laere syndrome']
NM_004056.4(CA8):c. 298T>C (p.Ser100Pro)	267606695	CA8	[]	[]	['Cerebellar ataxia, mental retardation, and dysequilibrium syndrome 3']
NM_000256.3(MYBP C3):c.3713T>C (p.Leu1238Pro)	730880702	MYBPC3	[]	[]	['Cardiomyopathy']
NM_001928.2(CFD):c .640T>C (p.Cys214Arg)	267606721	CFD	[]	['GGTGYGCGGGG GCGTGCTCGAGG', 'GTGYGCGGGGGC GTGCTCGAGGG']	['Complement factor d deficiency']
NM_005740.2(DNAL 4):c.153+2T>C	606231254	DNAL4	['CGAG GYATT GCCAG CAGTG CAGG']	['CGAGGYATTGCC AGCAGTGCAGG']	['Mirror movements 3']
NM_020975.4(RET):c. 2753T>C (p.Met918Thr)	74799832	RET	[]	[]	['Multiple endocrine neoplasia, type 2a', 'Multiple endocrine neoplasia, type 2b', 'Multiple endocrine neoplasia, type 2', 'Pheochromocytoma' , 'not provided']
NM_001849.3(COL6A 2):c.2329T>C (p.Cys777Arg)	267606747	COL6A2	[]	['CGCCYGCGACA AGCCACAGCAGG']	['Ullrich congenital muscular dystrophy']
NM_001006657.1(WD R35):c.781T>C (p.Trp261Arg)	431905505	WDR35	[]	[]	['Short rib polydactyly syndrome 5']
NM_003764.3(STX11) :c.173T>C (p.Leu58Pro)	431905512	STX11	['TGCY GGTGG CCGAC GTGAA GCGG']	['TGCYGGTGGCCG ACGTGAAGCGG']	['Hemophagocytic lymphohistiocytosis, familial, 4']
NM_001044.4(SLC6A	431905515	SLC6A3	[]	['CTGCACCYCCAC	['Infantile

3):c.671T>C (p.Leu224Pro)				CAGAGCCATGG']	Parkinsonism- dystonia']
NM_000277.1(PAH):c .764T>C (p.Leu255Ser)	62642930	PAH	[]	[]	['Phenylketonuria', 'not provided']
NM_000277.1(PAH):c .932T>C (p.Leu311Pro)	62642936	PAH	[]	[]	['Phenylketonuria', 'not provided']
NM_000118.3(ENG):c .2T>C (p.Met1Thr)	267606783	ENG	[]	[]	['Osler hemorrhagic telangiectasia syndrome']
NM_000129.3(F13A1) :c.728T>C (p.Met243Thr)	267606788	F13A1	['TGTG AYGGA CAGAG CACAA ATGG']	['TGTGAYGGACA GAGCACAAATGG']	['Factor xiii, a subunit, deficiency of']
NM_000257.3(MYH7) :c.1952A>G (p.His651Arg)	606231328	MYH7	[]	[]	['Familial cardiomyopathy']
NM_014053.3(FLVCR 1):c.574T>C (p.Cys192Arg)	267606821	FLVCR1	[]	[]	['Posterior column ataxia with retinitis pigmentosa']
NM_005249.4(FOXG1)):c.643T>C (p.Phe215Leu)	267606828	FOXG1	[]	[]	['Rett syndrome, congenital variant']
NM_015474.3(SAMH D1):c.1153A>G (p.Met385Val)	515726140	SAMHD1	[]	[]	['Aicardi Goutieres syndrome 5']
NM_015474.3(SAMH D1):c.1411-2A>G	515726141	SAMHD1	[]	[]	['Aicardi Goutieres syndrome 5']
NM_000180.3(GUCY 2D):c.2846T>C (p.Ile949Thr)	267606857	GUCY2D	[]	['AGAGAYCGCCA ACATGTCACTGG']	['Cone-rod dystrophy 6']
NM_022489.3(INF2):c .556T>C (p.Ser186Pro)	267606877	INF2	[]	[]	['Focal segmental glomerulosclerosis 5']
NM_000257.3(MYH7) :c.1048T>C (p.Tyr350His)	730880863	MYH7	[]	[]	['Cardiomyopathy']
NM_022489.3(INF2):c .125T>C (p.Leu42Pro)	267606880	INF2	[]	['GCTGCYCCAGAT GCCCTCTGTGG']	['Focal segmental glomerulosclerosis 5']
m.4681T>C	267606889	MT-ND2	[]	[]	['Leigh disease', 'Leigh syndrome due to mitochondrial complex I deficiency']
m.10191T>C	267606890	MT-ND3	[]	[]	['Leigh disease', 'Mitochondrial complex I deficiency']
m.10563T>C	267606892	MT-ND4L	[]	[]	['Familial colorectal cancer']
m.12706T>C	267606893	MT-ND5	[]	[]	['Leigh disease', 'Leigh syndrome due to mitochondrial complex I deficiency']
NM_015713.4(RRM2)	515726190	RRM2B	[]	[]	['RRM2B-related

B):c.556A>G (p.Arg186Gly)					mitochondrial disease']
NM_015713.4(RRM2B):c.581A>G (p.Glu194Gly)	515726191	RRM2B	[]	['AACTCCTYCTACAGCAGCAAAGG']	['RRM2B-related mitochondrial disease']
NM_000315.2(PTH):c.52T>C (p.Cys18Arg)	104894271	PTH	['AATTYGTTCCTTACAAAATTCGG']	['AATTYGTTCCTTACAAAATTCGG']	['Hypoparathyroidism familial isolated']
NM_001136271.2(NKX2-6):c.451T>C (p.Phe151Leu)	267606914	NKX2-6	[]	[]	['Persistent truncus arteriosus']
NM_004646.3(NPHS1):c.793T>C (p.Cys265Arg)	267606917	NPHS1	[]	['GCTGCCGYGCGTGGCCCGAGGGG', 'CTGCCGYGCGTG GCCCGAGGGG']	['Finnish congenital nephrotic syndrome']
NM_000406.2(GNRHR):c.94A>G (p.Thr32Ala)	515726219	GNRHR	[]	[]	['Hypogonadotropic hypogonadism']
NM_152296.4(ATP1A3):c.2270T>C (p.Leu757Pro)	606231436	ATP1A3	[]	[]	['Alternating hemiplegia of childhood 2']
NM_000513.2(OPN1MW):c.529T>C (p.Trp177Arg)	267606927	OPN1MW	[]	[]	['Cone dystrophy 5, X-linked']
NM_152296.4(ATP1A3):c.1144T>C (p.Trp382Arg)	606231448	ATP1A3	[]	[]	['Dystonia 12']
NM_024411.4(PDYN):c.632T>C (p.Leu211Ser)	267606940	PDYN	[]	[]	['Spinocerebellar ataxia 23']
NM_000444.5(PHEX):c.755T>C (p.Phe252Ser)	267606945	PHEX	[]	[]	['Familial X-linked hypophosphatemic vitamin D refractory rickets']
NM_001543.4(NDST1):c.1918T>C (p.Phe640Leu)	606231458	NDST1	[]	[]	['Mental retardation, autosomal recessive 46']
NM_013382.5(POMT2):c.2242T>C (p.Trp748Arg)	267606964	POMT2	[]	[]	['Congenital muscular dystrophy-dystroglycanopathy with mental retardation, type B2']
NM_006121.3(KRT1):c.482T>C (p.Leu161Pro)	57695159	KRT1	[]	[]	['Bullous ichthyosiform erythroderma', 'not provided']
NM_016203.3(PRKAG2):c.1459T>C (p.Tyr487His)	267606976	PRKAG2	[]	[]	['Familial hypertrophic cardiomyopathy 6', 'not provided']
NM_016203.3(PRKAG2):c.1642T>C (p.Ser548Pro)	267606979	PRKAG2	[]	[]	['Familial hypertrophic cardiomyopathy 6', 'not provided']
NM_198965.1(PTHLH):c.179T>C	267606985	PTHLH	[]	[]	['Brachydactyly type E2']

(p.Leu60Pro)					
NM_198965.1(PTHLH):c.131T>C (p.Leu44Pro)	267606986	PTHLH	[]	[]	['Brachydactyly type E2']
NM_004990.3(MARS):c.1108T>C (p.Phe370Leu)	140467171	MARS	[]	[]	['Interstitial lung and liver disease']
NM_173560.3(RFX6):c.649T>C (p.Ser217Pro)	267607012	RFX6	[]	[]	['Mitchell-Riley syndrome']
NM_002942.4(ROBO2):c.2834T>C (p.Ile945Thr)	267607014	ROBO2	['GAGAYTGGAAA TTTTGGCCGTGG']	['GAGAYTGGAAA TTTTGGCCGTGG']	['Vesicoureteral reflux 2']
NM_178857.5(RPIL1):c.2878T>C (p.Trp960Arg)	267607018	RPIL1	[]	[]	['Occult macular dystrophy']
NM_002880.3(RAF1):c.1423T>C (p.Phe475Leu)	730881003	RAF1	[]	[]	['Rasopathy']
NM_015272.3(RPGRIPI1):c.1975T>C (p.Ser659Pro)	267607020	RPGRIPI1	[]	[]	['Joubert syndrome 7', 'COACH syndrome']
NM_015559.2(SETBP1):c.2612T>C (p.Ile871Thr)	267607038	SETBP1	[]	[]	['Schinzel-Giedion syndrome']
NM_000433.3(NCF2):c.481A>G (p.Lys161Glu)	137878529	NCF2	[]	[]	['Chronic granulomatous disease, autosomal recessive cytochrome b-positive, type 2']
NM_001041.3(SI):c.1022T>C (p.Leu341Pro)	267607049	SI	[]	[]	['Sucrase-isomaltase deficiency']
NM_005633.3(SOS1):c.1294T>C (p.Trp432Arg)	267607080	SOS1	['GGTYGGGAG GGGAGGGAAA AGACA TTGG']	['GGTYGGGAGGG AAAAGACATTGG']	['Noonan syndrome 4', 'Rasopathy']
NM_018136.4(ASPM):c.2419+2T>C	587783225	ASPM	[]	[]	['Primary autosomal recessive microcephaly 5']
NM_001199107.1(TBC1D24):c.751T>C (p.Phe251Leu)	267607104	TBC1D24	[]	['CAAGTTCYTCCA CAAGGTGAGGG', 'TTCYTCCACAAG GTGAGGGCCGG']	['Myoclonic epilepsy, familial infantile']
NM_153704.5(TMEM67):c.1769T>C (p.Phe590Ser)	267607115	TMEM67	[]	[]	['Joubert syndrome 6', 'COACH syndrome']
NM_153704.5(TMEM67):c.2498T>C (p.Ile833Thr)	267607119	TMEM67	[]	[]	['Joubert syndrome 6', 'COACH syndrome']
NM_133378.4(TTN):c.100163T>C (p.Leu33388Pro)	267607156	-	[]	[]	['Distal myopathy Markesbery-Griggs type']
m.12811T>C	199974018	MT-ND5	[]	[]	['Leber optic atrophy']
NM_144631.5(ZNF51)	267607182	ZNF513	[]	['TGGGCGCYGCAT']	['Retinitis']

3):c.1015T>C (p.Cys339Arg)				GCGAGGAGAGG', 'CGCYGCATGCCGA GGAGAGGCTGG']	pigmentosa 58']
NM_004737.4(LARGE):c.1483T>C (p.Trp495Arg)	267607209	LARGE	[]	[]	['Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies, type A6']
NM_000229.1(LCAT):c.508T>C (p.Trp170Arg)	267607211	LCAT	[]	['TATGACYGGCG GCTGGAGCCCGG']	['Norum disease']
NM_016269.4(LEF1):c.181T>C (p.Ser61Pro)	267607215	-	[]	['GAACGAGYCTG AAATCATCCCGG']	['Sebaceous tumors, somatic']
NM_139248.2(LIPH):c.322T>C (p.Trp108Arg)	267607219	LIPH	[]	[]	['Woolly hair, autosomal recessive 2, with or without hypotrichosis']
NM_004268.4(MED17):c.1112T>C (p.Leu371Pro)	267607232	MED17	[]	[]	['Microcephaly, postnatal progressive, with seizures and brain atrophy']
NM_000530.6(MPZ):c.341T>C (p.Ile114Thr)	267607241	MPZ	[]	[]	[]
NM_000489.4(ATRX):c.4840T>C (p.Cys1614Arg)	122445094	ATRX	[]	[]	['ATR-X syndrome']
NM_000489.4(ATRX):c.6250T>C (p.Tyr2084His)	122445097	ATRX	[]	[]	['ATR-X syndrome']
NM_000489.4(ATRX):c.1226T>C (p.Leu409Ser)	122445109	ATRX	[]	[]	[]
NM_000489.4(ATRX):c.6149T>C (p.Ile2050Thr)	122445110	ATRX	[]	[]	['Multiple congenital anomalies']
NM_178151.2(DCX):c.272T>C (p.Leu91Pro)	587783536	DCX	[]	[]	['Heterotopia']
NM_178151.2(DCX):c.2T>C (p.Met1Thr)	587783539	DCX	['CAAA ATAYG GAACT TGATT TTGG']	['CAAAATAYGGA ACTTGATTTTGG']	['Heterotopia']
NM_178151.2(DCX):c.412T>C (p.Tyr138His)	587783551	DCX	[]	[]	['Heterotopia']
NM_000212.2(ITGB3):c.2231T>C (p.Leu744Pro)	398122374	-	[]	[]	['Platelet-type bleeding disorder 16']
NM_178151.2(DCX):c.641T>C (p.Ile214Thr)	587783574	DCX	[]	[]	['Heterotopia']
NM_178151.2(DCX):c.683T>C (p.Leu228Pro)	587783580	DCX	[]	['AAAAAACYCTA CACTCTGGATGG']	['Heterotopia']
NM_001005360.2(DNM2):c.1862T>C (p.Leu621Pro)	587783597	DNM2	[]	[]	['Myopathy, centronuclear']
NM_006579.2(EBP):c.310T>C (p.Tyr104His)	587783609	EBP	[]	[]	['Chondrodysplasia punctata 2 X-linked

					dominant']
NM_004004.5(GJB2): c.107T>C (p.Leu36Pro)	587783644	GJB2	[]	['GATCCYCGTTGT GGCTGCAAAGG']	['Hearing impairment']
NM_005682.6(ADGR G1):c.1460T>C (p.Leu487Pro)	587783653	ADGRG1	[]	['CCCTGCYCACCT GCCTTTCCTGG']	['Polymicrogyria, bilateral frontoparietal']
NM_000525.3(KCNJ1 1):c.988T>C (p.Tyr330His)	587783675	KCNJ11	[]	[]	['Diabetes mellitus']
NM_170707.3(LMNA (p.Tyr267His)	267607593	LMNA	[]	[]	['Dilated cardiomyopathy 1A', 'not provided']
NM_000252.2(MTM1) :c.1353+2T>C	587783780	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1) :c.1367T>C (p.Phe456Ser)	587783783	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1) :c.1433T>C (p.Phe478Ser)	587783794	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1) :c.1495T>C (p.Trp499Arg)	587783801	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1) :c.260T>C (p.Leu87Pro)	587783816	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1) :c.683T>C (p.Leu228Pro)	587783851	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1) :c.958T>C (p.Ser320Pro)	587783863	MTM1	[]	['GGAAYCTTTAAA AAAAGTGAAGG']	['Severe X-linked myotubular myopathy']
NM_000526.4(KRT14 (p.Leu384Pro)	59629244	KRT14	[]	[]	['Epidermolysis bullosa simplex, Koebner type', 'not provided']
NM_000249.3(MLH1) :c.453+2T>C	267607751	MLH1	[]	['ATCACGGYAAG AATGGTACATGG', 'TCACGGYAAGAA TGGTACATGGG']	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_002764.3(PRPS1) :c.455T>C (p.Leu152Pro)	80338676	PRPS1	[]	[]	['Arts syndrome', 'not provided']
NM_022132.4(MCCC 2):c.499T>C (p.Cys167Arg)	119103222	MCCC2	[]	[]	['3-methylcrotonyl CoA carboxylase 2 deficiency']
NM_000411.6(HLCS): c.710T>C (p.Leu237Pro)	119103227	HLCS	[]	['CTATCYTTCTCA GGGAGGGAAGG']	['Holocarboxylase synthetase deficiency']
NM_005787.5(ALG3): c.211T>C (p.Trp71Arg)	119103237	ALG3	[]	['GATTGACYGGA AGGCCTACATGG']	['Congenital disorder of glycosylation type 1D']
NM_005609.2(PYGM) :c.1187T>C (p.Leu396Pro)	119103254	PYGM	[]	[]	['Glycogen storage disease, type V']
m.3250T>C	199474664	MT-TL1	[]	[]	[]

NM_002764.3(PRPS1):c.344T>C (p.Met115Thr)	80338732	PRPS1	['GCAAATAYGCTATCTGTAGCAGG']	['GCAAATAYGCTATCTGTAGCAGG']	['Charcot-Marie-Tooth disease, X-linked recessive, type 5']
NM_003172.3(SURF1):c.679T>C (p.Trp227Arg)	398122806	SURF1	[]	['CCACYGGCATTATCGAGACCTGG']	['Congenital myasthenic syndrome, acetazolamide-responsive']
NM_004525.2(LRP2):c.7564T>C (p.Tyr2522His)	80338747	LRP2	[]	['GTACCTGYACTGGCTGACTGGG']	['Donnai Barrow syndrome']
NM_006329.3(FBLN5):c.649T>C (p.Cys217Arg)	80338766	FBLN5	[]	[]	['Autosomal recessive cutis laxa type IA']
NM_001271723.1(FBXO38):c.616T>C (p.Cys206Arg)	398122838	FBXO38	[]	['TTCCTYGTATCCCAATGCTAAGG']	['Distal hereditary motor neuropathy 2D']
NM_133433.3(NIPBL):c.7062+2T>C	587784032	NIPBL	[]	[]	['Cornelia de Lange syndrome 1']
NM_000334.4(SCN4A):c.4468T>C (p.Phe1490Leu)	80338790	SCN4A	[]	[]	['Hyperkalemic Periodic Paralysis Type 1']
NM_058216.2(RAD51C):c.404+2T>C	730881931	RAD51C	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_001111.4(ADAR):c.2615T>C (p.Ile872Thr)	398122897	ADAR	[]	[]	['Aicardi-goutieres syndrome 6']
NM_005334.2(HCFC1):c.-970T>C	398122908	HCFC1	['CAAGAYGGCCTCCAGGGAGG']	['CAAGAYGGCCTCCAGGGAGG']	['Mental retardation 3, X-linked']
NM_000431.3(MVK):c.1039+2T>C	398122910	MVK	['CCAGGYATCCCGGGGTAGGTGG']	['CCAGGYATCCCGGGGTAGGTGG', 'CAGGYATCCCGGGGTAGGTGGG']	['Porokeratosis, disseminated superficial actinic 1']
NM_000431.3(MVK):c.1094T>C (p.Phe365Ser)	398122911	MVK	[]	[]	['Porokeratosis, disseminated superficial actinic 1']
NM_005050.3(ABCD4):c.956A>G (p.Tyr319Cys)	201777056	ABCD4	['GATGAGGYAATGACACAAAGG']	['GATGAGGYAATGACACAAAGG']	['METHYLMALONIC ACIDURIA AND HOMOCYSTINURIA, cblJ TYPE', 'not provided']
NM_000251.2(MSH2):c.2005+2T>C	267607987	MSH2	['CTGGYAAAAAACCTGGTTTGG', 'TGGYAAAAAACCTGGTTTTGG']	['CTGGYAAAAAACCTGGTTTGG', 'TGGYAAAAAACCTGGTTTTGG']	['Hereditary Nonpolyposis Colorectal Neoplasms']

NM_000518.4(HBB):c.257T>C (p.Phe86Ser)	35693898	HBB	[]	[]	['Hemoglobinopathy']
NM_001194998.1(CEP152):c.3149T>C (p.Leu1050Pro)	398122977	CEP152	[]	[]	['Primary autosomal recessive microcephaly 9']
NM_022455.4(NSD1):c.5989T>C (p.Tyr1997His)	587784171	NSD1	[]	[]	['Sotos syndrome 1']
NM_014495.3(ANGPTL3):c.883T>C (p.Phe295Leu)	398122989	-	[]	['ACAAAACYTCAATGAAACGTGGG']	['Hypobetalipoproteinemia, familial, 2']
NM_024577.3(SH3TC2):c.1982T>C (p.Leu661Pro)	80338927	SH3TC2	[]	[]	['Charcot-Marie-Tooth disease, type 4C']
NM_000551.3(VHL):c.227T>C (p.Phe76Ser)	730882033	VHL	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_004004.5(GJB2):c.269T>C (p.Leu90Pro)	80338945	GJB2	[]	['GCTCCYAGTGGCCATGCACGTGG']	['Deafness, autosomal recessive 1A', 'Hearing impairment', 'not provided']
NM_000334.4(SCN4A):c.2078T>C (p.Ile693Thr)	80338956	SCN4A	[]	['AAGATCAYTGGCAATTCAGTGGG', 'AGATCAYTGGCAATTCAGTGGGG', 'GATCAYTGGCAATTCAGTGGGGG']	['Hyperkalemic Periodic Paralysis Type 1', 'Paramyotonia congenita of von Eulenburg']
NM_004523.3(KIF11):c.2547+2T>C	730882063	KIF11	['GGAGGYAATAACTTGTAA GTGG']	['GGAGGYAATAACTTTGTAAGTGG']	['Microcephaly with or without chorioretinopathy, lymphedema, or mental retardation']
NM_003060.3(SLC22A5):c.1051T>C (p.Trp351Arg)	68018207	SLC22A5	[]	[]	['Renal carnitine transport defect']
NM_001070.4(TUBG1):c.1160T>C (p.Leu387Pro)	398123045	TUBG1	[]	[]	['Cortical dysplasia, complex, with other brain malformations 4']
NM_000441.1(SLC26A4):c.-103T>C	60284988	-	[]	[]	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_000016.5(ACADM):c.233T>C (p.Ile78Thr)	398123074	ACADM	[]	[]	['Medium-chain acyl-coenzyme A dehydrogenase deficiency', 'not provided']
NM_000179.2(MSH6):c.4001+2T>C	267608131	MSH6	[]	['CGGYAACTAACTAACTATAATGG']	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_000019.3(ACAT1):c.730+2T>C	398123096	ACAT1	[]	[]	['Deficiency of acetyl-CoA acetyltransferase', 'not provided']
NM_000430.3(PAFAH1B1):c.841T>C (p.Cys281Arg)	587784288	PAFAH1B1	[]	[]	['Lissencephaly 1']

NM_000899.4(KITLG):c.98T>C (p.Val33Ala)	730882156	KITLG	[]	[]	['Familial progressive hyperpigmentation with or without hypopigmentation']
NM_015599.2(PGM3):c.248T>C (p.Leu83Ser)	267608260	PGM3	['AATGTYGGC ACCAT CCTGG GAGG']	['AATGTYGGCACC ATCCTGGGAGG']	['Immunodeficiency 23']
NM_000169.2(GLA):c.899T>C (p.Leu300Pro)	398123223	-	[]	[]	['Fabry disease']
NM_001256047.1(C19orf12):c.391A>G (p.Lys131Glu)	146170087	C19orf12	[]	[]	['Neurodegeneration with brain iron accumulation 4']
NM_172337.2(OTX2):c.674A>G (p.Asn225Ser)	370761964	OTX2	[]	[]	['Pituitary hormone deficiency, combined 6']
NM_000202.6(IDS):c.587T>C (p.Leu196Ser)	398123250	IDS	[]	[]	['Mucopolysaccharidosis, MPS-II', 'not provided']
NM_000252.2(MTM1):c.688T>C (p.Trp230Arg)	398123274	MTM1	[]	[]	['Severe X-linked myotubular myopathy', 'not provided']
NM_022445.3(TPK1):c.656A>G (p.Asn219Ser)	371271054	TPK1	[]	[]	['THIAMINE METABOLISM DYSFUNCTION SYNDROME 5 (EPISODIC ENCEPHALOPATHY TYPE)']
NM_014139.2(SCN11A):c.2432T>C (p.Leu811Pro)	483352920	SCN11A	[]	[]	['NEUROPATHY, HEREDITARY SENSORY AND AUTONOMIC, TYPE VII']
NM_000733.3(CD3E):c.520+2T>C	483352928	CD3E	[]	[]	['Immunodeficiency 18']
NM_017653.3(DYM):c.621-2A>G	775414124	DYM	[]	[]	['Dyggve-Melchior-Clausen syndrome']
NM_001253816.1(SLC52A2):c.1016T>C (p.Leu339Pro)	148234606	SLC52A2	[]	[]	['Brown-Vialetto-Van Laere syndrome 2']
NM_004963.3(GUCY2C):c.2782T>C (p.Cys928Arg)	587784573	-	[]	['TCCCYGTGCTGCTGGAGTTGTGG', 'CCCYGTGCTGCTGGAGTTGTGG']	['Meconium ileus']
NM_003159.2(CDKL5):c.659T>C (p.Leu220Pro)	267608511	CDKL5	[]	['CCAACYTTTTAC TATTCAGAAGG']	['Early infantile epileptic encephalopathy 2']
NM_000528.3(MAN2B1):c.2436+2T>C	398123457	MAN2B1	[]	[]	['not provided']
NM_002136.2(HNRNPA1):c.841T>C (p.Phe281Leu)	483353031	HNRNPA1	['AATYTTGGA GGCAG AAGCT CTGG']	['AATYTTGGAGGC AGAAGCTCTGG']	['Chronic progressive multiple sclerosis']

NM_006920.4(SCN1A):c.4251+2T>C	398123595	-	[]	[]	['Severe myoclonic epilepsy in infancy', 'Generalized epilepsy with febrile seizures plus, type 2']
NM_002225.3(IVD):c.465+2T>C	398123683	IVD	[]	[]	['Isovaleryl-CoA dehydrogenase deficiency', 'not provided']
NM_000118.3(ENG):c.1273-2A>G	373842615	ENG	[]	['CCGCCYGC GGG GATAAAGCCAGG', 'CGCCYGC GGGGA TAAAGCCAGG']	['Haemorrhagic telangiectasia 1']
NM_005859.4(PURA):c.218T>C (p.Phe73Ser)	793888535	PURA	[]	[]	['not provided']
NM_003494.3(DYSF):c.1284+2T>C	398123765	DYSF	['ATGG YAAGG AGCAA GGGAG CAGG']	['ATGGYAAGGAG CAAGGGAGCAGG']	['Limb-girdle muscular dystrophy, type 2B', 'not provided']
NM_004006.2(DMD):c.2380+2T>C	398123885	DMD	[]	[]	['Dilated cardiomyopathy 3B']
NM_006364.2(SEC23A):c.2104A>G (p.Met702Val)	138568622	SEC23A	[]	[]	['Craniolenticulosutural dysplasia']
NM_000335.4(SCN5A):c.376A>G (p.Lys126Glu)	185492581	SCN5A	[]	['GAATCTYCACAG CCGCTCTCCGG']	['Brugada syndrome']
NM_015865.6(SLC14A1):c.871T>C (p.Ser291Pro)	78242949	SLC14A1	[]	[]	[]
NM_003995.3(NPR2):c.226T>C (p.Ser76Pro)	796065355	NPR2	[]	[]	['SHORT STATURE WITH NONSPECIFIC SKELETAL ABNORMALITIES']
NM_012463.3(ATP6V0A2):c.825+2T>C	398124257	ATP6V0A2	['CACT GYGAG TAAGC TGGAA GTGG']	['CACTGYGAGTA AGCTGGAAGTGG']	['Cutis laxa with osteodystrophy', 'not provided']
NM_014795.3(ZEB2):c.73+2T>C	398124282	ZEB2	[]	[]	['Mowat-Wilson syndrome']
NM_000424.3(KRT5):c.1388T>C (p.Leu463Pro)	57599352	KRT5	[]	[]	['Epidermolysis bullosa simplex, Koebner type', 'not provided']
NM_133499.2(SYN1):c.1699A>G (p.Thr567Ala)	200533370	SYN1	[]	['GATGYCTGACG GGTAGCCTGTGG', 'ATGYCTGACGGG TAGCCTGTGGG']	['Epilepsy, X-linked, with variable learning disabilities and behavior disorders', 'not specified']
NM_148960.2(CLDN19):c.269T>C (p.Leu90Pro)	118203981	CLDN19	[]	['GCTCCYGGGCTT CGTGGCCATGG']	['Hypomagnesemia 5, renal, with ocular involvement']

NM_018006.4(TRMU):c.229T>C (p.Tyr77His)	118203990	TRMU	[]	[]	['Liver failure acute infantile']
NM_000056.3(BCKDHB):c.752T>C (p.Val251Ala)	398124593	BCKDHB	[]	[]	['Maple syrup urine disease', 'not provided']
NM_182680.1(AMELX):c.2T>C (p.Met1Thr)	104894737	-	[]	[]	['Amelogenesis imperfecta, type 1E']
NM_018105.2(THAP1):c.241T>C (p.Phe81Leu)	118204013	THAP1	[]	[]	['Dystonia 6, torsion']
NM_001235.3(SERP1NH1):c.233T>C (p.Leu78Pro)	137853892	SERP1NH1	[]	['GTCGCYAGGGCTCGTGTCGCTGG', 'TCGCYAGGGCTCGTGTCGCTGGG']	['Osteogenesis imperfecta type 10']
NM_004482.3(GALNT3):c.516_688del	761396172	GALNT3	[]	[]	['Tumoral calcinosis, familial, hyperphosphatemic']
NM_000263.3(NAGLU):c.142T>C (p.Phe48Leu)	118204024	NAGLU	[]	['GGCCGACYTCTCCGTGTCGGTGG']	['Mucopolysaccharidosis, MPS-III-B']
NM_000559.2(HBG1):c.-251T>C	35710727	HBG1	[]	[]	['Fetal hemoglobin quantitative trait locus 1']
NM_000527.4(LDLR):c.694+2T>C	200238879	LDLR	['CGGYATGGGCCGGGCCAGG', 'CGGGCCAGG', 'GTGG']	['ACTGCGGYATGGCGGGGCCAGG', 'CTGCGGYATGGGCCGGGCCAGG', 'CGGYATGGGCCGGGCCAGGGTGG']	['Familial hypercholesterolemia']
NM_001012515.2(FECH):c.1268T>C (p.Phe423Ser)	118204039	FECH	[]	[]	['Erythropoietic protoporphyria']
NM_005211.3(CSF1R):c.1745T>C (p.Leu582Pro)	690016563	CSF1R	[]	['CAACCYGCAGTTGGTGAGATGG']	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_000526.4(KRT14):c.1243T>C (p.Tyr415His)	58380626	KRT14	[]	['CGCCACCYACCGCCGCTGCTGG', 'CACCYACCGCCGCTGCTGGAGG', 'ACCYACCGCCGCTGCTGGAGGG']	['Epidermolysis bullosa herpetiformis, Dowling-Meara', 'not provided']
NM_006493.2(CLN5):c.2T>C (p.Met1Thr)	201615354	CLN5	[]	[]	['not provided']
NM_002863.4(PYGL):c.2461T>C (p.Tyr821His)	113993988	PYGL	['AAGAYATGCCCAAAACATCTGG']	['AAGAAYATGCCCAAAACATCTGG']	['Glycogen storage disease, type VI']
NM_016038.2(SBDS):c.258+2T>C	113993993	SBDS	[]	[]	['Shwachman syndrome', 'Aplastic anemia, susceptibility to']
NM_000110.3(DPYD):c.85T>C (p.Cys29Arg)	1801265	DPYD	[]	[]	['Dihydropyrimidine dehydrogenase deficiency']
NM_001034116.1(EIF)	113994038	EIF2B4	[]	[]	['Ovarioleukodystro

2B4):c.1393T>C (p.Cys465Arg)					phy']
NM_001165963.1(SCN1A):c.323T>C (p.Leu108Pro)	794726793	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_001034116.1(EIF2B4):c.1465T>C (p.Tyr489His)	113994040	EIF2B4	[]	[]	['Ovarioleukodystrophy']
NM_004304.4(ALK):c.3749T>C (p.Ile1250Thr)	113994092	ALK	[]	[]	['Neuroblastoma 3']
NM_207346.2(TSEN54):c.277T>C (p.Ser93Pro)	113994151	TSEN54	[]	['TTGAAGYCTCCC GCGGTGAGCGG', 'AAGYCTCCCGCG GTGAGCGGCGG']	['Pontocerebellar hypoplasia type 4']
NM_000018.3(ACADVL):c.848T>C (p.Val283Ala)	113994167	ACADVL	['TTTGY GGTGG AGAGG GGCTT CGG', 'TTGYG GTGGA GAGGG GCTTC GGG']	['TTTGYGGTGGAG AGGGGCTTCGG', 'TTGYGGTGGAGA GGGGCTTCGGG']	['Very long chain acyl-CoA dehydrogenase deficiency', 'not provided']
NM_000430.3(PAFAH1B1):c.569-10T>C	113994202	PAFAH1B1	[]	[]	['Lissencephaly 1']
NM_004937.2(CTNS):c.473T>C (p.Leu158Pro)	113994206	CTNS	[]	['TGGTCYAGCTT CGACTTCGTGG']	['Cystinosis']
NM_000546.5(TP53):c.488A>G (p.Tyr163Cys)	148924904	TP53	['GCTTG YAGAT GGCCA TGGCG CGG']	['GCTTGYAGATGG CCATGGCGCGG']	['Hereditary cancer-predisposing syndrome']
NM_004211.3(SLC6A5):c.1444T>C (p.Trp482Arg)	281864925	SLC6A5	[]	[]	['Hyperekplexia 3']
NM_024312.4(GNPTAB):c.1208T>C (p.Ile403Thr)	281864973	GNPTAB	[]	[]	['Pseudo-Hurler polydystrophy']
NM_024312.4(GNPTAB):c.3002T>C (p.Leu1001Pro)	281865006	GNPTAB	[]	[]	['I cell disease']
NM_000540.2(RYR1):c.7358T>C (p.Ile2453Thr)	118192123	RYR1	[]	[]	['Central core disease', 'not provided']
NM_000748.2(CHRN2):c.923T>C (p.Val308Ala)	281865070	CHRN2	[]	[]	['Epilepsy, nocturnal frontal lobe, type 3']
NM_000526.4(KRT14):c.356T>C (p.Met119Thr)	28928893	KRT14	[]	[]	['Epidermolysis bullosa herpetiformis, Dowling-Meara', 'not provided']
NM_000093.4(COL5A1):c.5137-11T>A	183495554	-	[]	[]	['Ehlers-Danlos syndrome, classic type']
NM_000277.1(PAH):c	62516109	PAH	[]	['CCTTCYTGAA	['Phenylketonuria',

.638T>C (p.Leu213Pro)				AAGTACTGTGG']	'not provided']
NM_000531.5(OTC):c .663+2T>C	72558427	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c .717+2T>C	72558431	OTC	[]	[]	['Ornithine carbamoyltransferas e deficiency', 'not provided']
NM_000531.5(OTC):c .793T>C (p.Trp265Arg)	72558445	OTC	[]	[]	['not provided']
NM_000493.3(COL10 A1):c.1841T>C (p.Leu614Pro)	111033545	-	[]	[]	['Metaphyseal chondrodysplasia, Schmid type']
NM_000493.3(COL10 A1):c.1771T>C (p.Cys591Arg)	111033546	-	[]	[]	['Metaphyseal chondrodysplasia, Schmid type']
NM_000493.3(COL10 A1):c.2011T>C (p.Ser671Pro)	111033552	-	[]	[]	['Metaphyseal chondrodysplasia, Schmid type']
NM_004614.4(TK2):c. 173A>G (p.Asn58Ser)	138439950	TK2	[]	[]	['Mitochondrial DNA depletion syndrome 2']
NM_004614.4(TK2):c. 644T>C (p.Leu215Pro)	281865497	TK2	[]	[]	['Mitochondrial DNA depletion syndrome 2']
NM_004614.4(TK2):c. 156+2T>C	281865499	TK2	[]	[]	['Mitochondrial DNA depletion syndrome 2']
NM_153026.2(PRICK LE1):c.1414T>C (p.Tyr472His)	281865564	PRICKLE1	[]	[]	['Progressive myoclonus epilepsy with ataxia']
NM_017882.2(CLN6): c.767A>G (p.Asp256Gly)	143781303	CLN6	[]	[]	['not provided']
NM_130838.1(UBE3A):c.389T>C (p.Ile130Thr)	111033597	UBE3A	[]	[]	['Angelman syndrome']
NM_001198799.2(AS CC1):c.953A>G (p.Asn318Ser)	146370051	ASCC1	[]	[]	[]
NM_000174.4(GP9):c. 70T>C (p.Cys24Arg)	28933378	GP9	['CCCA YGTA CTGCC GCGCC CTGG']	['CCCAYGTACCTG CCGCGCCCTGG']	['Bernard Soulier syndrome', 'Bernard- Soulier syndrome type C']
NM_001173464.1(KIF 21A):c.3029T>C (p.Ile1010Thr)	121912587	KIF21A	[]	[]	['Fibrosis of extraocular muscles, congenital, 1']
NM_001302946.1(TR NT1):c.668T>C (p.Ile223Thr)	370011798	TRNT1	[]	['GCAAYTGCAGA AAATGCAAAAAGG']	['Sideroblastic anemia with B-cell immunodeficiency, periodic fevers, and developmental delay']
NM_000371.3(TTR):c. 250T>C (p.Phe84Leu)	121918091	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis']
m.5814T>C	200077222	MT-TC	[]	[]	['Juvenile myopathy,

					encephalopathy, lactic acidosis AND stroke']
NM_000277.1(PAH):c.293T>C (p.Leu98Ser)	62517167	PAH	[]	['AAGATCTYGAGGCATGACATTGG']	['Mild non-PKU hyperphenylalanemia', 'not provided']
NM_017882.2(CLN6):c.486+2T>C	796052355	CLN6	[]	[]	['not provided']
NM_001813.2(CENPE):c.4063A>G (p.Lys1355Glu)	141488085	CENPE	[]	[]	['Primary autosomal recessive microcephaly 13']
NM_001918.3(DBT):c.1150G>A (p.Gly384Ser)	12021720	DBT	[]	['GACYCACAGAGCCCAATTTCTGG']	['Intermediate maple syrup urine disease type 2']
NM_000495.4(COL4A5):c.4699T>C (p.Cys1567Arg)	104886288	COL4A5	['AGTAYGTGAAGCTCAGCTGTGG']	['AGTAYGTGAAGCTCAGCTGTGG']	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.4756T>C (p.Cys1586Arg)	104886289	COL4A5	[]	['TCCCCATYGTCC TCAGGGATGGG']	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.5032T>C (p.Cys1678Arg)	104886310	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000155.3(GALT):c.482T>C (p.Leu161Pro)	111033700	GALT	['AGCYGGGTGCCAGTACCCTTGG']	['AGCYGGGTGCCAGTACCCTTGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_001199252.2(SGOL1):c.67A>G (p.Lys23Glu)	199815268	-	[]	[]	['Chronic atrial and intestinal dysrhythmia']
NC_012920.1:m.5559A>G	370471013	MT-TW	[]	['CAACYTACTGAGGGCTTTGAAGG']	['Leigh disease']
NM_000487.5(ARSA):c.410T>C (p.Leu137Pro)	121434215	ARSA	[]	['GCCTTCCYGCCCCCCATCAGGG']	['Metachromatic leukodystrophy, adult type']
NM_000051.3(ATM):c.7967T>C (p.Leu2656Pro)	121434218	-	[]	[]	[]
NM_000096.3(CP):c.650T>C (p.Phe217Ser)	386134125	CP	[]	[]	['Deficiency of ferroxidase']
NM_000096.3(CP):c.1123T>C (p.Tyr375His)	386134128	CP	[]	['ACACTACYACAT TGCCGCTGAGG']	['Deficiency of ferroxidase']
NM_000268.3(NF2):c.185T>C (p.Phe62Ser)	121434261	NF2	[]	[]	['Neurofibromatosis, type 2']
NM_000495.4(COL4A5):c.4803+121T>C	104886423	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_024529.4(CDC73):c.191T>C (p.Leu64Pro)	121434264	CDC73	[]	[]	['Hyperparathyroidism 1']
NM_001061.4(TBXAS1):c.1463T>C (p.Leu488Pro)	199422114	TBXAS1	[]	[]	[]
NM_001127328.2(ACADM):c.1136T>C (p.Ile379Thr)	121434275	ACADM	[]	['GTGCAGAYACTTGGAGGCAATGG']	['Medium-chain acyl-coenzyme A dehydrogenase deficiency']

NM_001127328.2(ACADM):c.742T>C (p.Cys248Arg)	121434276	ACADM	[]	['CAGCGAYGTTCA GATACTAGAGG']	['Medium-chain acyl-coenzyme A dehydrogenase deficiency']
NM_000016.5(ACADM):c.199T>C (p.Tyr67His)	121434280	ACADM	[]	[]	['Medium-chain acyl-coenzyme A dehydrogenase deficiency', 'not provided']
NM_002225.3(IVD):c.134T>C (p.Leu45Pro)	121434284	IVD	[]	['ATGGGCYAAGC GAGGAGCAGAGG']	['ISOVALERIC ACIDEMIA, TYPE I']
NM_005957.4(MTHFR):c.968T>C (p.Leu323Pro)	121434297	MTHFR	[]	[]	['Homocystinuria due to MTHFR deficiency']
NM_000136.2(FANCC):c.1661T>C (p.Leu554Pro)	104886458	-	[]	[]	['Fanconi anemia, complementation group C', 'not provided']
NM_005908.3(MANBA):c.1513T>C (p.Ser505Pro)	121434334	MANBA	[]	['ATTACGYCCAGT CCTACAAATGG', 'TTACGYCCAGTC CTACAAATGGG', 'TACGYCCAGTCC TACAAATGGGG']	['Beta-D-mannosidosis']
NM_000244.3(MEN1):c.518T>C (p.Leu173Pro)	386134256	MEN1	[]	[]	['Multiple endocrine neoplasia, type 1']
NM_199242.2(UNC13D):c.1208T>C (p.Leu403Pro)	121434353	UNC13D	[]	[]	['Hemophagocytic lymphohistiocytosis, familial, 3']
NM_152783.4(D2HGDH):c.1331T>C (p.Val444Ala)	121434360	D2HGDH	[]	[]	['D-2-hydroxyglutaric aciduria 1']
NM_207118.2(GTF2H5):c.62T>C (p.Leu21Pro)	121434365	GTF2H5	[]	[]	['Photosensitive trichothiodystrophy']
NM_000159.3(GCDH):c.883T>C (p.Tyr295His)	121434366	GCDH	[]	['CGCCGGYACG GCATCGCGTGGG', 'GCCCGGYACGGC ATCGCGTGGGG']	['Glutaric aciduria, type 1']
NM_018668.4(VPS33B):c.89T>C (p.Leu30Pro)	121434385	VPS33B	[]	[]	['Arthrogryposis renal dysfunction cholestasis syndrome']
NM_000424.3(KRT5):c.541T>C (p.Ser181Pro)	60715293	KRT5	[]	['GTTTGCCYCCTT CATCGACAAGG']	['Epidermolysis bullosa herpetiformis, Dowling-Meara', 'not provided']
NM_001003722.1(GLE1):c.2051T>C (p.Ile684Thr)	121434409	GLE1	[]	['AAGGACAYTCCT GTCCCAAGGG']	['Lethal arthrogryposis with anterior horn cell disease']
NM_003659.3(AGPS):c.1406T>C (p.Leu469Pro)	121434413	AGPS	[]	[]	['Rhizomelic chondrodysplasia punctata type 3']
NM_004550.4(NDUFS2):c.1237T>C	121434429	NDUFS2	[]	[]	['Mitochondrial complex I']

(p.Ser413Pro)					deficiency', 'not provided']
NM_001287.5(CLCN7):c.2297T>C (p.Leu766Pro)	121434434	CLCN7	[]	['GGGCCYGCGGC ACCTGGTGGTGG']	['Osteopetrosis autosomal recessive 4']
m.14709T>C	121434453	MT-TE	[]	[]	['Diabetes-deafness syndrome maternally transmitted']
NM_000466.2(PEX1):c.1991T>C (p.Leu664Pro)	121434455	PEX1	[]	['GATGACCYTGAC CTCATTGCTGG']	['Zellweger syndrome']
NM_198253.2(TERT):c.3043T>C (p.Cys1015Arg)	199422307	TERT	[]	[]	['Aplastic anemia']
m.4290T>C	121434469	MT-TI	['ACTYT GATAG AGTAA ATAAT AGG']	['ACTYTGATAGAG TAAATAATAGG']	[]
m.4291T>C	121434471	MT-TI	['ACTTY GATAG AGTAA ATAAT AGG']	['ACTTYGATAGAG TAAATAATAGG']	['Hypertension, hypercholesterolemia, and hypomagnesemia, mitochondrial']
m.9997T>C	121434475	MT-TG	[]	[]	['Primary familial hypertrophic cardiomyopathy']
NM_001099274.1(TINF2):c.860T>C (p.Leu287Pro)	199422316	TINF2	[]	[]	['Dyskeratosis congenita autosomal dominant']
NM_001099274.1(TINF2):c.862T>C (p.Phe288Leu)	199422317	TINF2	[]	['CTGYTCCCTTT AGGAATCTCGG']	['Aplastic anemia']
NM_000430.3(PAFAH1B1):c.505T>C (p.Ser169Pro)	121434484	PAFAH1B1	[]	[]	['Subcortical band heterotopia']
NM_000430.3(PAFAH1B1):c.92T>C (p.Phe31Ser)	121434486	PAFAH1B1	[]	[]	['Lissencephaly 1']
NM_005535.2(IL12RB1):c.592T>C (p.Cys198Arg)	121434495	IL12RB1	[]	[]	['Immunodeficiency 30']
NM_030662.3(MAP2K2):c.400T>C (p.Tyr134His)	121434499	MAP2K2	[]	[]	['Cardiofaciocutaneous syndrome 4', 'Rasopathy', 'Noonan syndrome and Noonan-related syndrome']
NM_001065.3(TNFRSF1A):c.349T>C (p.Cys117Arg)	104895221	TNFRSF1A	[]	['CTCTTCTYGCAC AGTGGACCGGG']	['TNF receptor-associated periodic fever syndrome (TRAPS)']
NM_000123.3(ERCC5):c.2573T>C (p.Leu858Pro)	121434575	-	[]	[]	['Xeroderma pigmentosum, group G']
NM_001493.2(GDI1):c.275T>C (p.Leu92Pro)	121434607	GDI1	[]	[]	['X-Linked Mental Retardation 41']

NM_020061.5(OPN1LW):c.607T>C (p.Cys203Arg)	121434621	OPN1LW	[]	[]	['Cone monochromatism']
NM_024420.2(PLA2G4A):c.331T>C (p.Ser111Pro)	121434634	PLA2G4A	[]	[]	[]
NM_005557.3(KRT16):c.395T>C (p.Leu132Pro)	60944949	KRT16	[]	[]	['Pachyonychia congenita, type 1', 'not provided']
NM_000485.2(APRT):c.407T>C (p.Met136Thr)	28999113	APRT	[]	[]	['Adenine phosphoribosyltransferase deficiency', 'APRT deficiency, Japanese type']
NM_005270.4(GLI2):c.4663T>C (p.Ser1555Pro)	144372453	GLI2	[]	[]	['Holoprosencephaly 9', 'not specified']
NM_024753.4(TTC21B):c.2384T>C (p.Leu795Pro)	387907060	TTC21B	[]	[]	['Asphyxiating thoracic dystrophy 4']
NM_000155.3(GALT):c.680T>C (p.Leu227Pro)	111033846	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000138.4(FBN1):c.4987T>C (p.Cys1663Arg)	137854459	FBN1	[]	['GGGACAYGTTA CAACACCGTTGG']	['Marfan syndrome']
NM_032446.2(MEGF10):c.976T>C (p.Cys326Arg)	387907073	MEGF10	[]	[]	['Myopathy, areflexia, respiratory distress, and dysphagia, early-onset, mild variant']
NM_024027.4(COLEC11):c.505T>C (p.Ser169Pro)	387907075	COLEC11	[]	['CAGCTGYCCTGC CAGGGCCGCGG', 'AGCTGYCCTGCC AGGGCCGCGGG', 'GCTGYCCTGCCA GGGCCGCGGG', 'CTGYCCTGCCAG GGCCGCGGGG']	['Carnevale syndrome']
NM_001946.3(DUSP6):c.566A>G (p.Asn189Ser)	143946794	DUSP6	['CACT AYTGG GGTCT CGGTC AAGG']	['CACTAYTGGGGT CTCGGTCAAGG']	['Hypogonadotropic hypogonadism 19 with or without anosmia']
NM_000138.4(FBN1):c.3793T>C (p.Cys1265Arg)	137854474	FBN1	['CTTGY GTTAT GATGG ATTCA TGG']	['CTTGYGTTATGA TGGATTCATGG']	['Marfan syndrome']
NM_022068.3(PIEZO2):c.2134A>G (p.Met712Val)	587777453	PIEZO2	[]	[]	['Oculomegaly amyoplasia']
NM_000570.4(FCGR3B):c.316A= (p.Ile106=)	2290834	FCGR3B	[]	[]	[]
NM_000352.4(ABCC8):c.674T>C (p.Leu225Pro)	1048095	ABCC8	[]	['TGCYGTCCAAAG GCACCTACTGG']	['Permanent neonatal diabetes mellitus']

NM_153490.2(KRT13):c.332T>C (p.Leu111Pro)	59897026	KRT13	[]	[]	['White sponge nevus 2', 'not provided']
NM_000132.3(F8):c.1174T>C (p.Ser392Pro)	28933669	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000492.3(CFTR):c.1021T>C (p.Ser341Pro)	397508144	CFTR	[]	[]	['Cystic fibrosis']
NM_000133.3(F9):c.1058T>C (p.Val353Ala)	137852255	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_001202.3(BMP4):c.362A>G (p.His121Arg)	376960358	BMP4	['TTCGTGGYGG AAGCTCCTCA CCG']	['TTCGTGGYGGAA GCTCCTCACGG']	['Microphthalmia syndromic 6']
NM_000133.3(F9):c.1328T>C (p.Ile443Thr)	137852268	F9	['GAAYATATA CCAAGGTATC CCG']	['GAAYATATACC AAGGTATCCCG']	['Hereditary factor IX deficiency disease']
NM_000133.3(F9):c.1357T>C (p.Trp453Arg)	137852269	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_000435.2(NOTCH3):c.1363T>C (p.Cys455Arg)	28933698	NOTCH3	['ACCYGTATC TGTATGGCAG GTGG']	['TTCACCYGTATC TGTATGGCAGG', 'ACCYGTATCTGT ATGGCAGGTGG']	['Cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy']
NM_019074.3(DLL4):c.1168T>C (p.Cys390Arg)	796065347	DLL4	[]	['GAAYGTCCCCC AACTCACCG']	['Adams-Oliver syndrome', 'ADAMS-OLIVER SYNDROME 6']
NM_000032.4(ALAS2):c.595T>C (p.Tyr199His)	137852310	ALAS2	[]	[]	['Hereditary sideroblastic anemia']
NM_019074.3(DLL4):c.583T>C (p.Phe195Leu)	796065351	DLL4	[]	[]	['Adams-Oliver syndrome']
NM_000402.4(G6PD):c.1054T>C (p.Tyr352His)	137852347	G6PD	[]	['AGGGYACCTGG ACGACCCACGG']	['Anemia, nonspherocytic hemolytic, due to G6PD deficiency']
NM_007325.4(GRIA3):c.2117T>C (p.Met706Thr)	137852352	GRIA3	[]	[]	['Mental retardation, X-linked, syndromic, wu type']
NM_000132.3(F8):c.6554T>C (p.Leu2185Ser)	137852365	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.5372T>C (p.Met1791Thr)	137852375	F8	['TCAYGGTGA GTTAA GGACA GTGG']	['TCAYGGTGAGTT AAGGACAGTGG']	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.1754T>C (p.Ile585Thr)	137852376	F8	['AACA GAYAA']	['AACAGAYAATG TCAGACAAGAGG']	['Hereditary factor VIII deficiency']

			TGTCA GACAA GAGG']]	disease']
NM_001127695.1(CTSA):c.707T>C (p.Leu236Pro)	137854546	CTSA	[]	[]	['Galactosialidosis, early infantile']
NM_000132.3(F8):c.935T>C (p.Phe312Ser)	137852405	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.980T>C (p.Leu327Pro)	137852407	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000308.2(CTSA):c.1271T>C (p.Met424Thr)	137854548	CTSA	[]	[]	['Galactosialidosis, late infantile']
NM_000132.3(F8):c.1481T>C (p.Ile494Thr)	137852413	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_001250.5(CD40):c.247T>C (p.Cys83Arg)	28931586	CD40	[]	[]	['Immunodeficiency with hyper IgM type 3']
NM_000165.4(GJA1):c.32T>C (p.Leu1Pro)	121912969	GJA1	[]	[]	['Oculodentodigital dysplasia']
NM_000132.3(F8):c.1958T>C (p.Val653Ala)	137852430	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_001972.2(ELANE):c.211T>C (p.Cys71Arg)	28931611	ELANE	[]	[]	['Severe congenital neutropenia autosomal dominant']
NM_000098.2(CPT2):c.1342T>C (p.Phe448Leu)	74315297	CPT2	[]	[]	['CARNITINE PALMITOYLTRANSFERASE II DEFICIENCY, LATE-ONSET', 'not provided']
NM_213653.3(HFE2):c.842T>C (p.Ile281Thr)	74315326	HFE2	[]	[]	['Hemochromatosis type 2A']
NM_213653.3(HFE2):c.302T>C (p.Leu101Pro)	74315327	HFE2	[]	['GGACCYCGCCTT CCATTCGGCGG']	['Hemochromatosis type 2A']
NM_000194.2(HPRT1):c.122T>C (p.Leu41Pro)	137852480	HPRT1	[]	[]	['Lesch-Nyhan syndrome']
NM_000261.1(MYOC):c.1297T>C (p.Cys433Arg)	74315338	MYOC	[]	[]	['Primary open angle glaucoma juvenile onset 1']
NM_000194.2(HPRT1):c.170T>C (p.Met57Thr)	137852495	HPRT1	[]	[]	['Lesch-nyhan syndrome, neurologic variant']
NM_000267.3(NF1):c.3728T>C (p.Leu1243Pro)	137854564	NF1	[]	[]	['Neurofibromatosis, type 1']
NM_000291.3(PGK1):c.263T>C (p.Leu88Pro)	137852531	PGK1	[]	[]	['Phosphoglycerate kinase 1 deficiency']
NM_000291.3(PGK1):c.946T>C	137852533	PGK1	[]	[]	['Phosphoglycerate kinase 1 deficiency']

(p.Cys316Arg)					
NM_000291.3(PGK1): c.758T>C (p.Ile253Thr)	137852534	PGK1	[]	[]	['Phosphoglycerate kinase 1 deficiency']
NM_170784.2(MKKS) :c.830T>C (p.Leu277Pro)	74315398	MKKS	[]	[]	['Bardet-Biedl syndrome 6', 'not provided']
NM_000451.3(SHOX) :c.877T>C (p.Ter293Arg)	137852559	SHOX	[]	[]	['Leri Weill dyschondrosteosis']
NM_001029871.3(RSPO4): c.319T>C (p.Cys107Arg)	74315421	RSPO4	[]	[]	['Anonychia']
NM_000044.3(AR):c.2033T>C (p.Leu678Pro)	137852579	AR	[]	['GTCCYGGAAGC CATTGAGCCAGG']	[]
NM_000044.3(AR):c.2423T>C (p.Met808Thr)	137852592	AR	[]	[]	['Reifenstein syndrome']
NM_000044.3(AR):c.2596T>C (p.Ser866Pro)	137852597	AR	[]	[]	['Androgen resistance syndrome']
NM_172201.1(KCNE2) :c.161T>C (p.Met54Thr)	74315447	KCNE2	[]	[]	['Long QT syndrome 6', 'Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_172201.1(KCNE2) :c.170T>C (p.Ile57Thr)	74315448	KCNE2	[]	[]	['Long QT syndrome 6', 'Cardiac arrhythmia', 'not provided']
NM_000211.4(ITGB2) :c.446T>C (p.Leu149Pro)	137852611	ITGB2	['AGCYAGGTGGC GCGAC CTGCT CCGG']	['AGCYAGGTGGC GACCTGCTCCGG']	['Leukocyte adhesion deficiency']
NM_000211.4(ITGB2) :c.412T>C (p.Ser138Pro)	137852617	ITGB2	[]	[]	['Leukocyte adhesion deficiency']
NM_000023.2(SGCA) :c.524T>C (p.Val175Ala)	137852622	SGCA	[]	[]	['Limb-girdle muscular dystrophy, type 2D']
NM_001166107.1(HMGCS2): c.520T>C (p.Phe174Leu)	137852636	HMGCS2	[]	['CCCTCYTCAATG CTGCCAACTGG']	['mitochondrial 3-hydroxy-3-methylglutaryl-CoA synthase deficiency']
NM_001886.2(CRYBA4): c.281T>C (p.Phe94Ser)	74315486	CRYBA4	[]	[]	['Cataract 23']
NM_001886.2(CRYBA4): c.206T>C (p.Leu69Pro)	74315487	CRYBA4	[]	[]	['Cataract 23']
NM_002047.2(GARS) :c.548T>C (p.Leu183Pro)	137852644	GARS	[]	[]	['Distal hereditary motor neuropathy type 5']
NM_000268.3(NF2):c.1604T>C (p.Leu535Pro)	74315493	NF2	[]	[]	['Neurofibromatosis, type 2']
NM_000095.2(COMP) :c.982T>C	137852653	COMP	[]	[]	['Pseudoachondroplastic']

(p.Cys328Arg)					spondyloepiphyseal dysplasia syndrome']
NM_000095.2(COMP):c.1042T>C (p.Cys348Arg)	137852656	COMP	[]	[]	['Pseudoachondroplastic spondyloepiphyseal dysplasia syndrome']
NM_017929.5(PEX26):c.2T>C (p.Met1Thr)	74315506	PEX26	[]	[]	['Peroxisome biogenesis disorder 7B']
NM_033163.3(FGF8):c.118T>C (p.Phe40Leu)	137852661	FGF8	[]	['TTCCCTGYTCCG GGCTGGCCGGG']	['Kallmann syndrome 6']
NM_007315.3(STAT1):c.2117T>C (p.Leu706Ser)	137852677	STAT1	[]	[]	['Immunodeficiency 31a']
NM_007315.3(STAT1):c.1799T>C (p.Leu600Pro)	137852678	STAT1	[]	[]	['Mycobacterial and viral infections, susceptibility to, autosomal recessive']
NM_000336.2(SCNN1B):c.1858T>C (p.Tyr620His)	137852707	SCNN1B	[]	[]	['Pseudoprimary hyperaldosteronism']
NM_005215.3(DCC):c.503T>C (p.Met168Thr)	121912967	DCC	[]	['AGCCCA YGCCA ACAATCCACTGG']	[]
NM_001034850.2(FAM134B):c.873+2T>C	137852738	FAM134B	[]	[]	['Hereditary sensory and autonomic neuropathy type IIA']
NM_000182.4(HADHA):c.1025T>C (p.Leu342Pro)	137852772	HADHA	[]	[]	['Mitochondrial trifunctional protein deficiency']
NM_001165974.1(UROCI):c.209T>C (p.Leu70Pro)	137852796	UROCI	[]	[]	['Urocanate hydratase deficiency']
NM_000405.4(GM2A):c.412T>C (p.Cys138Arg)	137852797	GM2A	[]	[]	['Tay-Sachs disease, variant AB']
NM_001039523.2(CHRNA1):c.901T>C (p.Phe301Leu)	137852806	CHRNA1	[]	['TGTGYTCCTTCT GGTCATCGTGG']	['Myasthenic syndrome, congenital, fast-channel']
NM_003688.3(CASK):c.802T>C (p.Tyr268His)	137852817	CASK	[]	[]	['FG syndrome 4']
NM_003688.3(CASK):c.2740T>C (p.Trp914Arg)	137852819	CASK	['CACA GYGGG TCCCT GTCTC CTGG', 'ACAGY GGGTC CCTGT CTCCT GGG']	['CACAGYGGGTC CCTGTCTCCTGG', 'ACAGYGGGTCCC TGTCTCCTGGG']	['FG syndrome 4']
NM_182760.3(SUMF1):c.1006T>C (p.Cys336Arg)	137852848	SUMF1	[]	[]	['Multiple sulfatase deficiency']

NM_182760.3(SUMF1):c.463T>C (p.Ser155Pro)	137852850	SUMF1	[]	['GGCGACYCCTTTGTCTTTGAAGG']	['Multiple sulfatase deficiency', 'not provided']
NM_000158.3(GBE1):c.671T>C (p.Leu224Pro)	137852886	GBE1	[]	['AATGTACYACCAAGAATCAAAGG']	['Glycogen storage disease, type IV', 'GLYCOGEN STORAGE DISEASE IV, NONPROGRESSIVE HEPATIC']
m.8356T>C	118192099	MT-TK	[]	[]	['Myoclonus with epilepsy with ragged red fibers', 'MERRF/MELAS overlap syndrome']
NM_024312.4(GNPTAB):c.1120T>C (p.Phe374Leu)	137852900	GNPTAB	[]	[]	['Pseudo-Hurler polydystrophy', 'I cell disease']
NM_031924.4(RSPH3):c.631-2A>G	142800871	RSPH3	[]	[]	['Kartagener syndrome']
NM_058172.5(ANTXR2):c.566T>C (p.Ile189Thr)	137852905	ANTXR2	[]	[]	['Hyaline fibromatosis syndrome']
NM_000419.3(ITGA2B):c.641T>C (p.Leu214Pro)	137852911	ITGA2B	[]	['CTGGTGTCYTGCGGCTCCTGGCGG']	['Glanzmann thrombasthenia']
NM_001171507.2(MCFD2):c.407T>C (p.Ile136Thr)	137852914	MCFD2	[]	[]	['Factor v and factor viii, combined deficiency of, 2']
NM_000540.2(RYR1):c.1205T>C (p.Met402Thr)	118192117	RYR1	['CGCAYGATCCACAGCACCAATGG']	['CGCAYGATCCACAGCACCAATGG']	['Congenital myopathy with fiber type disproportion', 'Central core disease', 'not provided']
NM_000540.2(RYR1):c.13703T>C (p.Leu4568Pro)	118192131	RYR1	[]	[]	['Central core disease', 'not provided']
NM_000540.2(RYR1):c.13949T>C (p.Leu4650Pro)	118192138	RYR1	[]	[]	['Central core disease', 'not provided']
NM_138694.3(PKHD1):c.10658T>C (p.Ile3553Thr)	137852948	PKHD1	[]	['GAGCCCAYTGA AATACGCTCAGG']	['Polycystic kidney disease, infantile type']
NM_012464.4(TLL1):c.713T>C (p.Val238Ala)	137852952	TLL1	['GGGATTGTTGTTGTTCA TGAAT TGGG']	['GGGATTGTTGTT CATGAATTGGG']	['Atrial septal defect 6']
NM_000540.2(RYR1):c.14762T>C (p.Phe4921Ser)	118192156	RYR1	[]	[]	['Central core disease', 'not provided']
NM_024960.4(PANK2):c.178T>C (p.Ser60Pro)	137852964	PANK2	[]	['ATTGACYCAGTC GGATTCAATGG']	[]
NM_001001486.1(ATP2C1):c.1751T>C (p.Leu584Pro)	137853015	ATP2C1	[]	[]	['Familial benign pemphigus']
NM_014363.5(SACS):	137853017	SACS	[]	[]	['Spastic ataxia

c.5836T>C (p.Trp1946Arg)					Charlevoix-Saguenay type']
NM_014363.5(SACS): c.9742T>C (p.Trp3248Arg)	137853018	SACS	[]	[]	['Spastic ataxia Charlevoix-Saguenay type']
NM_014363.5(SACS): c.3161T>C (p.Phe1054Ser)	137853019	SACS	[]	[]	['Spastic ataxia Charlevoix-Saguenay type']
NM_006899.3(IDH3B)):c.395T>C (p.Leu132Pro)	137853020	IDH3B	[]	['TGCGGCYGAGG TAGGTGGTCTGG', 'GCGGCYGAGGTA GGTGGTCTGGG']	['Retinitis pigmentosa 46']
NM_001139.2(ALOX12B): c.1277T>C (p.Leu426Pro)	137853023	ALOX12B	[]	[]	['Autosomal recessive congenital ichthyosis 2']
NM_001080463.1(DYNC2H1): c.3719T>C (p.Ile1240Thr)	137853028	DYNC2H1	[]	[]	['Short-rib thoracic dysplasia 3 with or without polydactyly']
NM_006009.3(TUBA1A): c.1190T>C (p.Leu397Pro)	137853048	TUBA1A	[]	[]	['Lissencephaly 3']
NM_004519.3(KCNQ3): c.925T>C (p.Trp309Arg)	118192249	KCNQ3	[]	[]	['Benign familial neonatal seizures 2']
NM_000531.5(OTC): c.332T>C (p.Leu111Pro)	1800324	OTC	[]	[]	['Ornithine carbamoyltransferase deficiency']
NM_001172567.1(MYD88): c.317T>C (p.Leu106Pro)	137853065	MYD88	[]	[]	['Myd88 deficiency']
NM_002241.4(KCNJ10): c.418T>C (p.Cys140Arg)	137853068	KCNJ10	[]	[]	['SeSAME syndrome']
NM_000455.4(STK11): c.200T>C (p.Leu67Pro)	137853077	STK11	[]	[]	['Peutz-Jeghers syndrome']
NM_000518.4(HBB): c.332T>C (p.Leu111Pro)	35256489	HBB	[]	[]	['Beta thalassemia major']
NM_005094.3(SLC27A4): c.739T>C (p.Ser247Pro)	137853133	SLC27A4	[]	[]	['Ichthyosis prematurity syndrome']
NM_001145308.4(LRTOMT): c.313T>C (p.Trp105Arg)	137853186	LRTOMT	[]	[]	['Deafness, autosomal recessive 63']
NM_178012.4(TUBB2B): c.514T>C (p.Ser172Pro)	137853194	TUBB2B	[]	[]	['Polymicrogyria, asymmetric']
NM_178012.4(TUBB2B): c.683T>C (p.Leu228Pro)	137853195	TUBB2B	[]	[]	['Polymicrogyria, asymmetric']
NM_178012.4(TUBB2B): c.793T>C (p.Phe265Leu)	137853196	TUBB2B	[]	[]	['Polymicrogyria, asymmetric']
NM_001082971.1(DDC): c.925T>C (p.Phe309Leu)	137853209	DDC	[]	[]	['Deficiency of aromatic-L-amino-acid decarboxylase']
NM_006121.3(KRT1):	137853225	KRT1	[]	[]	['Bullous

c.1424T>C (p.Leu475Pro)					ichthyosiform erythroderma']
NM_033500.2(HK1):c.1550T>C (p.Leu517Ser)	137853249	HK1	[]	['GACTTCTYGGCCCTGGATCTTGG', 'TTCTYGGCCCTG GATCTTGGAGG']	['Hemolytic anemia due to hexokinase deficiency']
NM_000249.3(MLH1):c.229T>C (p.Cys77Arg)	63749859	MLH1	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_000444.5(PHEX):c.1664T>C (p.Leu555Pro)	137853270	PHEX	[]	['AGCYCCAGAAG CCTTTCCTTTGG']	['Familial X-linked hypophosphatemic vitamin D refractory rickets']
NM_000321.2(RB1):c.2134T>C (p.Cys712Arg)	137853296	RB1	[]	[]	['Retinoblastoma']
NM_007313.2(ABL1):c.988T>C (p.Phe330Leu)	137853304	ABL1	[]	[]	[]
NM_003639.4(IKBKG):c.1249T>C (p.Cys417Arg)	137853325	IKBKG	[]	['TGGAGYGCATTG AGTAGGGCCGG']	['Hypohidrotic ectodermal dysplasia with immune deficiency', 'Hyper-IgM immunodeficiency, X-linked, with hypohidrotic ectodermal dysplasia']
NM_017534.5(MYH2):c.5609T>C (p.Leu1870Pro)	786201023	-	[]	[]	['Inclusion body myopathy 3']
NM_000314.6(PTEN):c.406T>C (p.Cys136Arg)	786201044	PTEN	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_001257988.1(TYMP):c.665A>G (p.Lys222Arg)	149977726	TYMP	['CACGAGTYTCTTACTGAGAATGG']	['CACGAGTYTCTTACTGAGAATGGAGG']	[]
NM_016725.2(FOLR1):c.493+2T>C	144637717	FOLR1	['AGGYGAGGGCTGGG GTGGG CAGG']	['CTTCAGGYGAGGGCTGGGGTGGG', 'AGGYGAGGGCTGGGGTGGGCAGG']	['not provided']
NM_002225.3(IVD):c.295+2T>C	748026507	IVD	[]	[]	['not provided']
NM_000175.3(GPI):c.1016T>C (p.Leu339Pro)	137853587	GPI	[]	[]	[]
NM_002055.4(GFAP):c.1055T>C (p.Leu352Pro)	28932769	GFAP	[]	['GGACCYGCTCAATGTCAAGCTGG']	['Alexander disease', 'not provided']
NM_020921.3(NIN):c.3665A>G (p.Gln1222Arg)	187464517	NIN	[]	[]	['Seckel syndrome 7']
NM_005603.4(ATP8B1):c.2097+2T>C	387906381	ATP8B1	[]	[]	['Progressive intrahepatic

					cholestasis']
NM_005144.4(HR):c.-320T>C	387906382	HR	[]	[]	['Marie Unna hereditary hypotrichosis 1']
NM_001303.3(COX10):c.2T>C (p.Met1Thr)	387906383	COX10	[]	[]	['Congenital myasthenic syndrome, acetazolamide-responsive']
NM_004415.2(DSP):c.4961T>C (p.Leu1654Pro)	749730642	DSP	[]	[]	['not provided']
NM_000392.4(ABCC2):c.1967+2T>C	387906396	ABCC2	[]	[]	['Dubin-Johnson syndrome']
NM_002769.4(PRSS1):c.116T>C (p.Val39Ala)	397507439	-	[]	['TACCAGGYGTCCCTGAATTCTGG']	['Hereditary pancreatitis']
NM_130439.3(MXI1):c.552+2T>C	387906417	MXI1	[]	[]	['Malignant tumor of prostate']
m.8528T>C	387906422	-	[]	[]	['Cardiomyopathy, infantile hypertrophic']
NM_000132.3(F8):c.985T>C (p.Cys329Arg)	387906430	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.1417T>C (p.Tyr473His)	387906443	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.1729T>C (p.Ser577Pro)	387906446	F8	[]	['AAAGAAYCTGTAGATCAAAGAGG']	['Hereditary factor VIII deficiency disease']
NM_004333.4(BRAF):c.1403T>C (p.Phe468Ser)	397507473	BRAF	['ATCATYTGGACAGTCTACAAGG', 'TCATYTGGAA CAGTC TACAA GGG']	['ATCATYTGGAACAGTCTACAAGG', 'TCATYTGGAA CAGTC TACAA GGG']	['Cardiofaciocutaneous syndrome', 'Rasopathy']
NM_004333.4(BRAF):c.1454T>C (p.Leu485Ser)	397507475	BRAF	[]	[]	['Cardiofaciocutaneous syndrome', 'Rasopathy']
NM_005188.3(CBL):c.1201T>C (p.Cys401Arg)	397507492	CBL	[]	[]	['Rasopathy']
NM_000133.3(F9):c.52T>C (p.Cys18Arg)	387906474	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_005249.4(FOXG1):c.700T>C (p.Ser234Pro)	786205008	FOXG1	[]	[]	['Rett syndrome, congenital variant', 'not provided']
NM_000133.3(F9):c.82T>C (p.Cys28Arg)	387906481	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_000133.3(F9):c.1031T>C (p.Ile344Thr)	387906482	F9	[]	['ACGAACAYCTTCCTCAAATTTGG']	['Hereditary factor IX deficiency disease']

m.11253T>C	200145866	MT-ND4	[]	[]	['Leber optic atrophy']
NM_000131.4(F7):c.38T>C (p.Leu13Pro)	387906507	F7	[]	[]	['Factor VII deficiency']
NM_000131.4(F7):c.983T>C (p.Phe328Ser)	387906508	F7	[]	['GACGTYCTCTGAGAGGACGCTGG']	['Factor VII deficiency']
NM_000422.2(KRT17):c.296T>C (p.Leu99Pro)	28933089	KRT17	[]	[]	['Pachyonychia congenita type 2', 'not provided']
NM_001040113.1(MYH11):c.3791T>C (p.Leu1264Pro)	387906532	MYH11	[]	['GAAGCYGGAGGCGCAGGTGCAGG']	['Aortic aneurysm, familial thoracic 4']
NM_013246.2(CLCF1):c.46T>C (p.Cys16Arg)	137853934	-	[]	[]	['Cold-induced sweating syndrome 2']
NM_013246.2(CLCF1):c.676T>C (p.Ter226Arg)	137853935	-	[]	[]	['Cold-induced sweating syndrome 2']
NM_001077620.2(PRC1):c.2T>C (p.Met1Thr)	527236092	-	[]	[]	['Retinitis pigmentosa']
NM_000488.3(SERPINC1):c.68T>C (p.Leu23Pro)	387906575	SERPINC1	[]	[]	['Antithrombin III deficiency']
NM_206933.2(USH2A):c.9751T>C (p.Cys3251Arg)	527236118	USH2A	[]	[]	['Retinitis pigmentosa']
NM_001458.4(FLNC):c.752T>C (p.Met251Thr)	387906586	FLNC	[]	[]	['Myopathy, distal, 4']
NM_000781.2(CYP11A1):c.665T>C (p.Leu222Pro)	387906601	CYP11A1	[]	[]	['Adrenal insufficiency, congenital, with 46,XY sex reversal, partial or complete']
NM_000548.3(TSC2):c.2410T>C (p.Cys804Arg)	137853995	TSC2	[]	[]	['Tuberous sclerosis syndrome', 'Tuberous sclerosis 2', 'not provided']
NM_001145661.1(GATA2):c.1117T>C (p.Cys373Arg)	387906633	GATA2	[]	[]	['Lymphedema, primary, with myelodysplasia']
NM_002693.2(POLG):c.3470A>G (p.Asn1157Ser)	548076633	POLG	['CAAGAGGYTGGTCTGC AAGG']	['CAAGAGGYTGGTGATCTGCAAGG']	['not provided']
NM_002465.3(MYBPC1):c.706T>C (p.Trp236Arg)	387906657	MYBPC1	[]	[]	['Distal arthrogyriposis type 1B']
NM_002465.3(MYBPC1):c.2566T>C (p.Tyr856His)	387906658	MYBPC1	[]	['CAAACCYATATCGCAGAGTTGG']	['Distal arthrogyriposis type 1B']
NM_003392.4(WNT5A):c.544T>C (p.Cys182Arg)	387906663	WNT5A	[]	[]	['Robinow syndrome']
NM_005188.3(CBL):c.1186T>C (p.Cys396Arg)	387906665	CBL	[]	[]	['Rasopathy']

NM_006902.4(PRRX1):c.338T>C (p.Phe113Ser)	387906667	PRRX1	[]	[]	['Dysgnathia complex']
NM_001111035.1(ACP5):c.602T>C (p.Leu201Pro)	387906672	-	[]	[]	['Spondyloenchondrodysplasia with immune dysregulation']
NM_002734.4(PRKARIA):c.1117T>C (p.Tyr373His)	387906693	PRKARIA	[]	[]	['Acrodysostosis 1 with or without hormone resistance']
NM_002734.4(PRKARIA):c.980T>C (p.Ile327Thr)	387906695	PRKARIA	[]	[]	['Acrodysostosis 1 with or without hormone resistance']
NM_003491.3(NAA10):c.109T>C (p.Ser37Pro)	387906701	NAA10	[]	['TGGCCTTYCCTG GCCCAGGTGG', 'GGCCTTYCCTGG CCCAGGTGGG']	['N-terminal acetyltransferase deficiency']
NM_006306.3(SMC1A):c.2351T>C (p.Ile784Thr)	387906702	SMC1A	['AGAY TGGTG TGC GC AACAT CCGG']	['AGAYTGGTGTGC GCAACATCCGG']	['Congenital muscular hypertrophy-cerebral syndrome']
NM_000377.2(WAS):c.814T>C (p.Ser272Pro)	387906716	WAS	[]	[]	['Severe congenital neutropenia X-linked']
NM_000377.2(WAS):c.881T>C (p.Ile294Thr)	387906717	WAS	[]	['GACTTCAYTGAG GACCAGGGTGG', 'ACTTCAYTGAGG ACCAGGGTGGG']	['Severe congenital neutropenia X-linked']
m.12201T>C	387906733	MT-TH	[]	[]	['Deafness, nonsyndromic sensorineural, mitochondrial']
NM_139125.3(MASP1):c.1888T>C (p.Cys630Arg)	387906753	MASP1	[]	[]	['Michels syndrome']
NM_007315.3(STAT1):c.520T>C (p.Cys174Arg)	387906763	STAT1	[]	[]	['Immunodeficiency 31C']
NM_053025.3(MYLK):c.5275T>C (p.Ser1759Pro)	387906781	-	[]	[]	['Aortic aneurysm, familial thoracic 7']
NM_000287.3(PEX6):c.1601T>C (p.Leu534Pro)	387906809	PEX6	[]	['CTTCYGGGCCGG GACCGTGATGG', 'TTCYGGGCCGGG ACCGTGATGGG']	['Peroxisome biogenesis disorder 4B']
NM_000174.4(GP9):c.167T>C (p.Leu56Pro)	28933377	GP9	[]	[]	['Bernard-Soulier syndrome type C']
NM_004153.3(ORC1):c.266T>C (p.Phe89Ser)	387906827	ORC1	[]	[]	['Meier-Gorlin syndrome 1']
NM_021252.4(RAB18):c.619T>C (p.Ter207Gln)	387906833	RAB18	[]	[]	['Warburg micro syndrome 3']
NM_000702.3(ATP1A2):c.2291T>C (p.Leu764Pro)	28933398	ATP1A2	[]	[]	['Familial hemiplegic migraine type 2']
NM_000702.3(ATP1A2):c.2659T>C	28933399	ATP1A2	[]	[]	['Familial hemiplegic migraine']

(p.Trp887Arg)					type 2']
NM_000702.3(ATP1A2):c.2192T>C (p.Met731Thr)	28933400	ATP1A2	[]	[]	['Familial hemiplegic migraine type 2']
NM_020433.4(JPH2):c.421T>C (p.Tyr141His)	387906897	JPH2	[]	[]	['Familial hypertrophic cardiomyopathy 17']
NM_173170.1(IL36RN):c.80T>C (p.Leu27Pro)	387906914	IL36RN	[]	[]	['Pustular psoriasis, generalized']
NM_004990.3(MARS):c.1568T>C (p.Ile523Thr)	201555303	MARS	[]	[]	['Interstitial lung and liver disease']
NM_020191.2(MRPS22):c.644T>C (p.Leu215Pro)	387906924	MRPS22	['ATCYT AGGGT AAGGT GACTT AGG']	['ATCYTAGGGTAA GGTGACTTAGG']	['Combined oxidative phosphorylation deficiency 5']
NM_022445.3(TPK1):c.119T>C (p.Leu40Pro)	387906936	TPK1	[]	[]	['THIAMINE METABOLISM DYSFUNCTION SYNDROME 5 (EPISODIC ENCEPHALOPATHY TYPE)']
NM_020634.1(GDF3):c.914T>C (p.Leu305Pro)	387906945	GDF3	[]	[]	['Congenital ocular coloboma', 'Microphthalmia, isolated 7']
NM_024513.3(FYCO1):c.4127T>C (p.Leu1376Pro)	387906965	FYCO1	[]	['CAGCCYGATCCC CATCACTGTGG']	['Cataract, autosomal recessive congenital 2']
NM_006147.3(IRF6):c.65T>C (p.Leu22Pro)	387906967	IRF6	[]	['GCCYCTACCCTG GGCTCATCTGG']	['Van der Woude syndrome', 'Popliteal pterygium syndrome']
NM_025132.3(WDR19):c.2129T>C (p.Leu710Ser)	387906980	WDR19	[]	[]	['Cranioectodermal dysplasia 4', 'SENIOR-LOKEN SYNDROME 8']
NM_025132.3(WDR19):c.20T>C (p.Leu7Pro)	387906982	WDR19	[]	['TCTCACYGCTAG AAAAGACTTGG']	['Asphyxiating thoracic dystrophy 5']
NM_014874.3(MFN2):c.647T>C (p.Phe216Ser)	387906990	MFN2	[]	[]	[]
NM_016097.4(IER3IP1):c.233T>C (p.Leu78Pro)	387907012	IER3IP1	[]	[]	['Microcephaly, epilepsy, and diabetes syndrome']
NM_022489.3(INF2):c.310T>C (p.Cys104Arg)	387907034	INF2	[]	[]	['Charcot-Marie-Tooth disease, dominant intermediate E']
NM_022489.3(INF2):c.383T>C (p.Leu128Pro)	387907037	INF2	[]	[]	['Charcot-Marie-Tooth disease, dominant intermediate E']
NM_003235.4(TG):c.229T>C	137854433	TG	[]	[]	['Iodotyrosyl coupling defect']

(p.Cys1077Arg)					
NM_058246.3(DNAJB6):c.277T>C (p.Phe93Leu)	387907046	DNAJB6	[]	[]	['Limb-girdle muscular dystrophy, type 1E']
NM_032446.2(MEGF10):c.2320T>C (p.Cys774Arg)	387907072	MEGF10	[]	['GGGCAGYGTAC TTGCCGCACTGG']	['Myopathy, areflexia, respiratory distress, and dysphagia, early-onset', 'Myopathy, areflexia, respiratory distress, and dysphagia, early-onset, mild variant']
NM_005502.3(ABCA1):c.4429T>C (p.Cys1477Arg)	137854494	ABCA1	['CCTGT GYGTC CCCCA GGGGC AGG', 'CTGTG YGTCC CCCAG GGGCA GGG']	['CCTGTGYGTCCC CCAGGGGCAGG', 'CTGTGYGTCCCC CAGGGGCAGGG', 'TGTGYGTCCCC AGGGGCAGGGG', 'GTGYGTCCCCCA GGGGCAGGGGG']	['Tangier disease']
NM_005502.3(ABCA1):c.6026T>C (p.Phe2009Ser)	137854499	ABCA1	[]	['GAGTYCTTTGCC CTTTTGAGAGG']	['Familial hypoalphalipoproteinemia']
NM_015175.2(NBEAL2):c.1163T>C (p.Leu388Pro)	387907113	NBEAL2	[]	[]	['Gray platelet syndrome']
NM_000196.3(HSD11B2):c.1012T>C (p.Tyr338His)	387907117	HSD11B2	[]	['CCGCCGCYATTA CCCC GCCAGG', 'CGCCGCYATTAC CCCC GCCAGGG']	['Apparent mineralocorticoid excess']
NM_024599.5(RHBDF2):c.557T>C (p.Ile186Thr)	387907129	RHBDF2	['AGAY TGTGG ATCCG CTGGC CCGG']	['AGAYTGTGGATC CGCTGGCCCCGG']	['Howel-Evans syndrome']
NM_001077488.3(GNAS):c.299T>C (p.Leu100Pro)	137854531	GNAS	[]	[]	['Pseudohypoparathyroidism type 1A']
NM_001256714.1(DNAAF3):c.386T>C (p.Leu129Pro)	387907151	DNAAF3	[]	[]	['Kartagener syndrome', 'Ciliary dyskinesia, primary, 2']
NM_020894.2(UVSSA):c.94T>C (p.Cys32Arg)	387907164	UVSSA	['AAAA TTYGC AAGTA TGTCTT AGG']	['AAAATTYGCAA GTATGTCTTAGG', 'AAATTYGCAAGT ATGTCTTAGGG']	['UV-sensitive syndrome 3']
NM_000267.3(NF1):c.1523T>C (p.Leu508Pro)	137854558	NF1	[]	[]	['Neurofibromatosis, type 1']
NM_000267.3(NF1):c.6200T>C (p.Leu2067Pro)	137854561	NF1	[]	[]	['Neurofibromatosis, familial spinal']
NM_004453.3(ETFDH):c.1130T>C (p.Leu377Pro)	387907170	ETFDH	[]	['CCAAAACYCAC CTTTCTGGTGG']	[]

NM_000267.3(NF1):c.1070T>C (p.Leu357Pro)	137854563	NF1	[]	[]	['Neurofibromatosis, type 1', 'Neurofibromatosis, familial spinal']
NM_024306.4(FA2H):c.707T>C (p.Phe236Ser)	387907172	FA2H	[]	[]	['Spastic paraplegia 35']
NM_001004127.2(ALG11):c.1142T>C (p.Leu381Ser)	387907182	-	[]	[]	['Congenital disorder of glycosylation type 1P']
NM_021167.4(GATA D1):c.304T>C (p.Ser102Pro)	387907188	GATAD1	[]	[]	['Cardiomyopathy, dilated, 2b']
NM_033360.3(KRAS):c.211T>C (p.Tyr71His)	387907205	KRAS	[]	['GGACCAGYACA TGAGGACTGGGG', 'CCAGYACATGAG GACTGGGGAGG', 'CAGYACATGAGG ACTGGGGAGGG']	['Cardiofaciocutaneous syndrome 2']
NM_006265.2(RAD21):c.1753T>C (p.Cys585Arg)	387907213	-	[]	[]	['Cornelia de Lange syndrome 4']
NM_000222.2(KIT):c.1859T>C (p.Val620Ala)	387907217	KIT	[]	[]	[]
NM_000335.4(SCN5A):c.5380T>C (p.Tyr1794His)	137854615	SCN5A	[]	[]	['Brugada syndrome', 'Brugada syndrome 1']
NM_000076.2(CDKN1C):c.827T>C (p.Phe276Ser)	387907224	CDKN1C	[]	[]	['Intrauterine growth retardation, metaphyseal dysplasia, adrenal hypoplasia congenita, and genital anomalies']
NM_005691.3(ABCC9):c.3058T>C (p.Ser1020Pro)	387907229	ABCC9	[]	[]	['Hypertrichotic osteochondrodysplasia']
NM_012343.3(NNT):c.2930T>C (p.Leu977Pro)	387907233	NNT	[]	[]	['Glucocorticoid deficiency 4']
NM_024110.4(CARD14):c.467T>C (p.Leu156Pro)	387907240	CARD14	[]	['CAGCAGCYGCA GGAGCACCTGGG']	['Pityriasis rubra pilaris']
NM_002501.3(NFIX):c.179T>C (p.Leu60Pro)	387907254	NFIX	[]	[]	['Sotos syndrome 2']
NM_001165963.1(SCN1A):c.121A>T (p.Lys41Ter)	764444350	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_005022.3(PFN1):c.341T>C (p.Met114Thr)	387907265	PFN1	[]	[]	['Amyotrophic lateral sclerosis 18']
NM_001172567.1(MYD88):c.818T>C (p.Leu273Pro)	387907272	MYD88	[]	[]	['Macroglobulinemia, waldenstrom, somatic']
NM_152296.4(ATP1A3):c.2431T>C (p.Ser811Pro)	387907282	ATP1A3	[]	['TGCCATCYCACT GGCGTACGAGG']	['Alternating hemiplegia of childhood 2']

NM_022787.3(NMNAT1):c.838T>C (p.Ter280Gln)	387907290	NMNAT1	[]	[]	['Leber congenital amaurosis 9']
NM_005120.2(MED12):c.3493T>C (p.Ser1165Pro)	387907361	MED12	[]	['AGGACYCTGAGCCAGGGGCCCGG']	['Ohdo syndrome, X-linked']
NM_006194.3(PAX9):c.62T>C (p.Leu21Pro)	28933970	PAX9	[]	['GGCCGCGCC AACGCCATCCGG']	['Tooth agenesis, selective, 3']
NM_000492.3(CFTR):c.2780T>C (p.Leu927Pro)	397508435	CFTR	[]	[]	['Cystic fibrosis']
NM_177976.2(ARL6):c.272T>C (p.Ile91Thr)	137854907	ARL6	[]	[]	['Bardet-Biedl syndrome']

Table 8. Human gene mutations that may be corrected by changing a guanine (G) to an adenine (A). The gene name, gene symbol, and dbSNP database reference number (RS#) are indicated. Also indicated are exemplary protospacers with their PAM sequences (gRNAs and gRNAall) and the base to be edited *e.g.*, a C, indicated by a “Y”. The “gRNAs” sequences, from top to bottom, correspond to SEQ ID NOs: 3434-3601. The “gRNA all” sequences, from top to bottom, correspond to SEQ ID NOs: 3602-4266.

Name	RS# (dbSNP)	Gene Symbol	gRNAs	gRNAall	Phenotypes
NM_000138.4(FBN1):c.3128A>G (p.Lys1043Arg)	137854472	FBN1	[]	['TGCACYTGCCGTGGGTGCAGAGG']	[]
NM_000237.2(LPL):c.953A>G (p.Asn318Ser)	268	LPL	[]	[]	['Hyperlipidemia, familial combined']
NM_000257.3(MYH7):c.2708A>G (p.Glu903Gly)	727504261	MYH7	[]	['AGCGCYCCTCAGCATCTGCCAGG']	['Cardiomyopathy', 'not specified']
NM_000059.3(BRCA2):c.476-2A>G	81002853	BRCA2	[]	['ACCACYGGGGTAAAAAAGGGG', 'TACCACYGGGGTAAAAAAGGGG', 'ATACCACYGGGGTAAAAAAGG']	['Familial cancer of breast', 'Breast-ovarian cancer, familial 2', 'Hereditary cancer-predisposing syndrome']
NM_000059.3(BRCA2):c.9118-2A>G	81002862	BRCA2	[]	[]	['Familial cancer of breast', 'Breast-ovarian cancer, familial 2']
NM_000059.3(BRCA2):c.9649-2A>G	81002895	BRCA2	[]	[]	['Familial cancer of breast', 'Breast-ovarian cancer, familial 2']
NM_000387.5(SLC25A20):c.713A>G (p.Gln238Arg)	28934589	SLC25A20	[]	[]	['Carnitine acylcarnitine translocase deficiency', 'not provided']
NM_000060.3(BTD):c.755A>G (p.Asp252Gly)	28934601	BTD	[]	[]	['Biotinidase deficiency']
NM_172107.2(KCNQ2):c.851A>G	28939683	KCNQ2	[]	[]	['Benign familial neonatal seizures 1']

(p.Tyr284Cys)					
NM_006158.4(NEFL): c.293A>G (p.Asn98Ser)	58982919	NEFL	[]	[]	['Charcot-Marie-Tooth disease, type IF', 'not provided']
NM_000019.3(ACAT1): c.473A>G (p.Asn158Ser)	199524907	ACAT1	[]	[]	['Deficiency of acetyl-CoA acetyltransferase']
NM_007294.3(BRCA1): c.4987-2A>G	397509212	BRCA1	[]	[]	['Familial cancer of breast']
NM_007294.3(BRCA1): c.213-12A>G	80358163	BRCA1	[]	[]	['Familial cancer of breast', 'Hereditary breast and ovarian cancer syndrome', 'Breast-ovarian cancer, familial 1', 'Hereditary cancer-predisposing syndrome']
NM_001382.3(DPAGT1): c.509A>G (p.Tyr170Cys)	28934876	DPAGT1	['ACAYAGTACAGGATTCTGCGGG', 'GACAYAGTACAGTTCCTGCGG']	['ACAYAGTACAGGATTCTGCGGG', 'GACAYAGTACAGTTCCTGCGG']	['Congenital disorder of glycosylation type 1J']
NM_032237.4(POMK): c.773A>G (p.Gln258Arg)	397509386	POMK	[]	[]	['Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies, type A12']
NM_201647.2(STAMBP): c.125A>G (p.Glu42Gly)	397509387	STAMBP	[]	[]	['Microcephaly-capillary malformation syndrome']
NM_006876.2(B4GAT1): c.1168A>G (p.Asn390Asp)	397509397	B4GAT1	['CTGATYTTTCAGCCTCCTTTTGGG', 'GCTGATYTTTCAGCCTCCTTTTGGG']	['TGATYTTTCAGCCTCCTTTTGGG', 'CTGATYTTTCAGCCTCCTTTTGGG', 'GCTGATYTTTCAGCCTCCTTTTGGG']	['Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies, type A13']
NM_004315.4(ASAHI): c.965+4A>G	397509415	ASAHI	[]	[]	['Farber lipogranulomatosis']
NM_000257.3(MYH7): c.1477A>G (p.Met493Val)	730880875	MYH7	[]	[]	['Cardiomyopathy']
NM_021020.3(LZTS1): c.355A>G (p.Lys119Glu)	119473032	LZTS1	[]	['CCCTYCTCGGAGCCCTGTAGAGG']	[]
NM_022455.4(NSD1): c.5893-2A>G	587784163	NSD1	[]	[]	['Sotos syndrome 1']
NM_006894.5(FMO3): c.182A>G (p.Asn61Ser)	72549322	FMO3	[]	[]	['Trimethylaminuria']
NM_000403.3(GALE): c.101A>G (p.Asn34Ser)	121908046	GALE	['TGGAAGYTATCGATGACCACAGG']	['TGGAAGYTATCGATGACCACAGG']	['UDPglucose-4-epimerase deficiency']

]		
NM_000314.6(PTEN): c.527A>G (p.Tyr176Cys)	757498880	PTEN	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000053.3(ATP7B): c.2305A>G (p.Met769Val)	193922103	ATP7B	[]	[]	['Wilson disease', 'not specified', 'not provided']
NM_000173.6(GP1BA): c.763A>G (p.Met255Val)	121908064	GP1BA	[]	[]	['Pseudo von Willebrand disease']
NM_000422.2(KRT17): c.275A>G (p.Asn92Ser)	59151893	KRT17	['GTCAYTG AGGTTCTG CATGGTGG ']	['GTCAYTGAGGTT CTGCATGGTGG', 'GCGGTCAYTGAG GTTCTGCATGG']	['Pachyonychia congenita type 2', 'not provided']
NM_000288.3(PEX7): c.854A>G (p.His285Arg)	62653611	PEX7	[]	[]	['Rhizomelic chondrodysplasia punctata type 1']
NM_003742.2(ABCB11): c.890A>G (p.Glu297Gly)	11568372	ABCB11	[]	[]	['Progressive familial intrahepatic cholestasis 2', 'Benign recurrent intrahepatic cholestasis 2']
NM_012243.2(SLC35A3): c.886A>G (p.Ser296Gly)	141952252	SLC35A3	[]	[]	['Arthrogryposis, mental retardation, and seizures']
NM_000061.2(BTK): c.1082A>G (p.Tyr361Cys)	28935478	BTK	['GATGGYA GTTAATGA GCTCAGGG , 'TGATGGY AGTTAATG AGCTCAGG ']	['GATGGYAGTTA ATGAGCTCAGGG', 'TGATGGYAGTTA ATGAGCTCAGG']	[]
NM_000169.2(GLA): c.815A>G (p.Asn272Ser)	28935495	-	[]	[]	['Fabry disease']
NM_016069.9(PAM16): c.226A>G (p.Asn76Asp)	786203989	-	['TCATAGT YCTGCAGA GGAGAGG G']	['CATAGTYCTGCA GAGGAGAGGGG', 'TCATAGTYCTGC AGAGGAGAGGG']	['Chondrodysplasia, megarbane-daghermelki type']
NM_058163.1(TSR2): c.191A>G (p.Glu64Gly)	786203996	TSR2	[]	[]	['Diamond-Blackfan anemia with microtia and cleft palate', 'DIAMOND-BLACKFAN ANEMIA 14 WITH MANDIBULOFACIAL DYSOSTOSIS']
NM_001098398.1(COPA): c.728A>G (p.Asp243Gly)	794727994	-	[]	[]	[]
NM_005957.4(MTHFR): c.1114A>G (p.Lys372Glu)	786204024	MTHFR	[]	[]	['Homocysteinemia due to MTHFR deficiency']
NM_012193.3(FZD4): c.1024A>G (p.Met342Val)	80358293	-	[]	[]	['Exudative vitreoretinopathy 1']
NM_000186.3(CFH): c.	460184	CFH	['CAGYTGA	['CAGYTGAATTTG	['Atypical

.3590T>C (p.Val1197Ala)			ATTTGTGT GTAAACGG]	TGTGTAAACGG']	hemolytic-uremic syndrome 1']
NM_014846.3(KIAA0196):c.1411A>G (p.Asn471Asp)	80338865	KIAA0196	[]	[]	['Spastic paraplegia 8']
NM_014946.3(SPAST):c.1165A>G (p.Thr389Ala)	786204132	SPAST	['AGCATTG YCTTCCCA TTCCCAGG']	['ATTGYCTTCCCA TTCCAGGTGG', 'AGCATTGYCTTC CCATTCCCAGG']	['Spastic paraplegia 4, autosomal dominant']
NM_000925.3(PDHB):c.395A>G (p.Tyr132Cys)	28935769	PDHB	[]	[]	['Pyruvate dehydrogenase E1-beta deficiency']
NM_000090.3(COL3A1):c.2338-2A>G	794728050	COL3A1	[]	[]	['Thoracic aortic aneurysms and aortic dissections']
NM_000540.2(RYR1):c.97A>G (p.Lys33Glu)	193922746	RYR1	[]	[]	['King Denborough syndrome', 'not provided']
NM_000540.2(RYR1):c.7043A>G (p.Glu2348Gly)	193922801	RYR1	[]	['TTCYCCTCCACG CTCTCGCCTGG']	['not provided']
NM_000218.2(KCNQ1):c.652A>G (p.Lys218Glu)	36210419	KCNQ1	[]	['GCCCTYGGAGC CCACGCAGAGG']	['Torsades de pointes', 'Cardiac arrhythmia']
NM_000540.2(RYR1):c.14647-1449A>G	193922886	RYR1	[]	[]	['Minicore myopathy with external ophthalmoplegia', 'not provided']
NM_004429.4(EFNB1):c.472A>G (p.Met158Val)	28936071	EFNB1	[]	[]	['Craniofrontonasal dysplasia']
NM_007254.3(PNKP):c.1029+2T>C	199919568	PNKP	['CCGGYGA GGCCCTGG GGCGGGG G', 'TCCGGYG AGGCCCTG GGGCGGG G']	['CCGGYGAGGCC CTGGGGCGGGG' , 'TCCGGYGAGGCC CTGGGGCGGG', 'ATCCGGYGAGGC CCTGGGGCGGG', 'GATCCGGYGAGG CCCTGGGGCGG']	['not provided']
NM_005448.2(BMP15):c.704A>G (p.Tyr235Cys)	104894765	BMP15	['ATTGAAA YAGAGTAA CAAGAAG G']	['ATTGAAAYAGA GTAACAAGAAGG']	['Ovarian dysgenesis 2']
NM_004415.2(DSP):c.1141-2A>G	794728111	DSP	[]	[]	['not provided']
NM_016035.4(COQ4):c.155T>C (p.Leu52Ser)	786204770	COQ4	['GCTGTYG GCCGCCGG CTCCGCGG']	['GCTGTYGGCCGC CGGCTCCGCGG']	['COENZYME Q10 DEFICIENCY, PRIMARY, 7']
NM_015717.4(CD207):c.790T>C (p.Trp264Arg)	200837270	CD207	[]	[]	['Birbeck granule deficiency']
NM_004771.3(MMP20):c.611A>G (p.His204Arg)	786204826	MMP20	['CGAAAYG TGTATCTC CTCCCAGG']	['CGAAAYGTGTAT CTCCTCCCAGG']	['Amelogenesis imperfecta, hypomaturation type, IIA2']
NM_003392.4(WNT5)	786204836	WNT5A	[]	[]	['Robinow

A):c.257A>G (p.Tyr86Cys)					syndrome']
NM_000918.3(P4HB): c.1178A>G (p.Tyr393Cys)	786204843	P4HB	[]	[]	['Cole Carpenter syndrome']
NM_000314.6(PTEN): c.139A>G (p.Arg47Gly)	786204855	PTEN	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000314.6(PTEN): c.320A>G (p.Asp107Gly)	786204858	PTEN	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000447.2(PSEN2)):c.715A>G (p.Met239Val)	28936379	PSEN2	[]	[]	['Alzheimer disease, type 4', 'not provided']
NM_005263.3(GFI1):c .1145A>G (p.Asn382Ser)	28936381	GFI1	[]	[]	['Severe congenital neutropenia 2, autosomal dominant']
NM_005263.3(GFI1):c .1208A>G (p.Lys403Arg)	28936382	GFI1	[]	[]	['Neutropenia, nonimmune chronic idiopathic, of adults']
NM_000314.6(PTEN): c.512A>G (p.Gln171Arg)	786204865	PTEN	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000314.6(PTEN): c.254-2A>G	786204926	PTEN	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_001103.3(ACTN2)):c.1883A>G (p.Glu628Gly)	786204951	ACTN2	[]	[]	['Familial hypertrophic cardiomyopathy 23']
NM_003159.2(CDKL5)):c.-162-2A>G	786204973	CDKL5	[]	[]	['Early infantile epileptic encephalopathy 2']
NM_001159287.1(TPI1)):c.622A>G (p.Ile208Val)	121964849	TPI1	[]	[]	['Triosephosphate isomerase deficiency']
NM_003159.2(CDKL5)):c.2277-2A>G	786204979	CDKL5	[]	[]	['Early infantile epileptic encephalopathy 2']
NM_003159.2(CDKL5)):c.458A>G (p.Asp153Gly)	786204985	CDKL5	[]	[]	['Early infantile epileptic encephalopathy 2']
NM_003159.2(CDKL5)):c.91A>G (p.Arg31Gly)	786204991	CDKL5	[]	[]	['Early infantile epileptic encephalopathy 2']
NM_001007792.1(NTRK1): c.986A>G (p.Tyr329Cys)	121964869	NTRK1	[]	[]	['Hereditary insensitivity to pain with anhidrosis']
NM_001007792.1(NTRK1): c.1651A>G (p.Met551Val)	121964870	NTRK1	[]	[]	['Hereditary insensitivity to pain with anhidrosis']
NM_004360.3(CDH1): c.2512A>G (p.Ser838Gly)	121964872	CDH1	[]	[]	['Hereditary diffuse gastric cancer', 'Hereditary cancer-predisposing syndrome', 'Neoplasm of ovary']
NM_003122.4(SPINK1)	17107315	SPINK1	[]	[]	['Hereditary

1):c.101A>G (p.Asn34Ser)					pancreatitis', 'Tropical calcific pancreatitis', 'Pancreatitis, chronic, susceptibility to']
NM_021098.2(CACNA1H):c.4645A>G (p.Met1549Val)	786205050	CACNA1H	[]	[]	['Primary hyperaldosteronism']
NM_001385.2(DPYS): c.1001A>G (p.Gln334Arg)	121964923	DPYS	[]	[]	['Dihydropyrimidina se deficiency']
NM_003494.3(DYSF): c.3443-33A>G	786205083	DYSF	['GCCAGAG YGAGTGGC TGGAGTGG ']	['GCCAGAGYGAG TGGCTGGAGTGG']	['Limb-girdle muscular dystrophy, type 2B']
NM_003816.2(ADAM9):c.411-8A>G	786205086	ADAM9	[]	[]	[]
NM_000532.4(PCCB): c.1304A>G (p.Tyr435Cys)	121964961	PCCB	[]	[]	['Propionic acidemia']
NM_000071.2(CBS):c. 1150A>G (p.Lys384Glu)	121964967	CBS	['AACTYGG TCCTGCGG GATGGGGG ']	['AACTYGGTCCTG CGGGATGGGGG', 'GAACTYGGTCCT GCGGGATGGGG', 'GGA ACTYGGTCC TGCGGGATGGG', 'AGGAACTYGGTC CTGCGGGATGG']	['Homocystinuria, pyridoxine- responsive']
NM_000093.4(COL5A1):c.655-2A>G	786205101	COL5A1	[]	[]	['Ehlers-Danlos syndrome, classic type']
NM_000481.3(AMT): c.125A>G (p.His42Arg)	121964983	AMT	['GCCAGGY GGAAGTCA TAGAGCGG ']	['GCCAGGYGGAA GTCATAGAGCGG']	['Non-ketotic hyperglycinemia']
NM_000108.4(DLD):c. .214A>G (p.Lys72Glu)	121964987	DLD	[]	[]	['Maple syrup urine disease, type 3']
NM_000108.4(DLD):c. .1483A>G (p.Arg495Gly)	121964989	DLD	[]	['TTCTCYAAAAGC TTCTGATAAGG']	['Maple syrup urine disease, type 3']
NM_000108.4(DLD):c. .1081A>G (p.Met361Val)	121964993	DLD	[]	[]	['Maple syrup urine disease, type 3']
NM_001918.3(DBT):c. .1355A>G (p.His452Arg)	121965002	DBT	[]	[]	[]
NM_030813.5(CLPB): c.1850A>G (p.Tyr617Cys)	786205138	CLPB	[]	[]	['3- METHYLGLUTAC ONIC ACIDURIA WITH CATARACTS, NEUROLOGIC INVOLVEMENT, AND NEUTROPENIA']
NM_006610.3(MASP2):c.359A>G (p.Asp120Gly)	72550870	MASP2	[]	[]	['MASP2 deficiency']

NM_000398.6(CYB5R3):c.719A>G (p.Asp240Gly)	121965018	CYB5R3	[]	[]	['METHEMOGLOBINEMIA, TYPE I']
NM_000095.2(COMP):c.1358A>G (p.Asn453Ser)	28936668	COMP	[]	[]	[]
NM_000095.2(COMP):c.1418A>G (p.Asp473Gly)	28936669	COMP	[]	['ATTGYCGTCGTCGTCGTCGAGG']	[]
NM_003816.2(ADAM9):c.1396-2A>G	786205151	ADAM9	[]	[]	[]
NM_001204830.1(LIPT1):c.535A>G (p.Thr179Ala)	786205156	-	[]	[]	['LIPOYLTRANSFERASE 1 DEFICIENCY']
NM_000274.3(OAT):c.734A>G (p.Tyr245Cys)	121965046	OAT	[]	[]	['Ornithine aminotransferase deficiency']
NM_018488.2(TBX4):c.1592A>G (p.Gln531Arg)	28936696	TBX4	[]	['GTACYGTAAGG AAGATTCTCGGG', 'GGTACYGTAAGG AAGATTCTCGG']	['Ischiopatellar dysplasia']
NM_001110556.1(FLNA):c.1829-2A>G	786205183	FLNA	[]	[]	['X-linked periventricular heterotopia']
NM_003865.2(HESX1):c.541A>G (p.Thr181Ala)	28936704	HESX1	[]	[]	['Growth hormone deficiency with pituitary anomalies']
NM_000137.2(FAH):c.1141A>G (p.Arg381Gly)	121965077	FAH	[]	['TCCYGGTCTGACCATTCCCCAGG']	['Tyrosinemia type I']
NM_000137.2(FAH):c.836A>G (p.Gln279Arg)	121965078	FAH	[]	[]	['Tyrosinemia type I']
NM_006129.4(BMP1):c.808A>G (p.Met270Val)	786205219	BMP1	[]	[]	['Osteogenesis imperfecta type 13']
NM_001987.4(ETV6):c.1252A>G (p.Arg418Gly)	786205226	ETV6	[]	[]	['Thrombocytopenia 5']
NM_014423.3(AFF4):c.760A>G (p.Thr254Ala)	786205233	AFF4	[]	[]	['CHOPS SYNDROME']
NM_000140.3(FECH):c.68_194del	786205247	FECH	[]	[]	['Erythropoietic protoporphyria']
NM_000138.4(FBN1):c.3344A>G (p.Asp1115Gly)	794728203	FBN1	[]	['ACTCAYCAATATCTGCAAAATGG']	['Thoracic aortic aneurysms and aortic dissections']
NM_198270.3(NHS):c.853-2A>G	786205257	NHS	[]	[]	['Nance-Horan syndrome']
NM_005502.3(ABCA1):c.1790A>G (p.Gln597Arg)	2853578	ABCA1	[]	[]	['Tangier disease']
NM_003002.3(SDHD):c.275A>G (p.Asp92Gly)	786205436	SDHD	[]	['GAATAGYCCATCGCAGAGCAAGG']	['Fatal infantile mitochondrial cardiomyopathy']
NM_005259.2(MSTN):c.458A>G (p.Lys153Arg)	1805086	-	[]	[]	[]
NM_198056.2(SCN5A)	1805124	SCN5A	[]	[]	['Progressive

):c.1673A>G (p.His558Arg)					familial heart block type 1A', 'not specified', 'not provided']
NM_000784.3(CYP27A1):c.776A>G (p.Lys259Arg)	72551317	CYP27A1	[]	['AGTCCACYTGGG GAGGAAGGTGG']	['Cholestanol storage disease']
NM_000784.3(CYP27A1):c.1061A>G (p.Asp354Gly)	72551320	CYP27A1	[]	[]	['Cholestanol storage disease']
NM_000463.2(UGT1A1):c.992A>G (p.Gln331Arg)	72551348	-	[]	[]	['Crigler-Najjar syndrome, type II']
NM_000463.2(UGT1A1):c.1070A>G (p.Gln357Arg)	72551351	-	[]	[]	['Crigler Najjar syndrome, type 1']
NM_016218.2(POLK):c.1385A>G (p.Asn462Ser)	786205687	POLK	[]	['ATTACAYTCTT CAACTTAATGG']	['Malignant tumor of prostate']
NM_016218.2(POLK):c.181A>G (p.Asn61Asp)	786205689	POLK	[]	[]	['Malignant tumor of prostate']
NM_016218.2(POLK):c.1477A>G (p.Lys493Glu)	786205692	POLK	[]	[]	['Malignant tumor of prostate']
NM_000138.4(FBN1):c.7916A>G (p.Tyr2639Cys)	794728280	FBN1	[]	['TGTTTCAYACTGG AAGCCGGCGGG', 'CTGTTTCAYACTG GAAGCCGGCGG']	['Thoracic aortic aneurysms and aortic dissections']
NM_000132.3(F8):c.1331A>G (p.Lys444Arg)	28937272	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000531.5(OTC):c.268A>G (p.Ser90Gly)	72554340	OTC	[]	[]	['not provided']
NM_005502.3(ABCA1):c.2804A>G (p.Asn935Ser)	28937313	ABCA1	['TCCAYTG TGGCCAG GAAGGAG G']	['TCCAYTGTGGCC CAGGAAGGAGG', 'CGCTCCAYTGTG GCCCAGGAAGG']	['Tangier disease']
NM_004380.2(CREBBP):c.3524A>G (p.Tyr1175Cys)	28937315	CREBBP	[]	[]	['Rubinstein-Taybi syndrome']
NM_000335.4(SCN5A):c.3971A>G (p.Asn1324Ser)	28937317	SCN5A	[]	['GCAYTGACCACC ACCTCAAGTGG']	['Long QT syndrome 3', 'Congenital long QT syndrome']
NM_000891.2(KCNJ2):c.901A>G (p.Met301Val)	786205818	KCNJ2	[]	[]	['Cardiac arrhythmia']
NM_144499.2(GNAT1):c.386A>G (p.Asp129Gly)	786205854	GNAT1	[]	['CGGAGYCCTTCC ACAGCCGCTGG']	['NIGHT BLINDNESS, CONGENITAL STATIONARY, TYPE 1G']
NM_000523.3(HOXD13):c.974A>G (p.Gln325Arg)	104893635	HOXD13	[]	[]	['Syndactyly type 5']
NM_013953.3(PAX8):c.160A>G (p.Ser54Gly)	104893660	-	[]	[]	['Thyroid agenesis']
NM_014585.5(SLC40)	104893663	SLC40A1	[]	[]	['Hemochromatosis']

A1):c.470A>G (p.Asp157Gly)					type 4']
NM_003124.4(SCR):c. 448A>G (p.Arg150Gly)	104893665	SCR	[]	[]	['Sepiapterin reductase deficiency']
NM_000258.2(MYL3) :c.445A>G (p.Met149Val)	104893748	MYL3	[]	[]	['Familial hypertrophic cardiomyopathy 8', 'Cardiomyopathy', 'Hypertrophic cardiomyopathy']
NM_000539.3(RHO):c. .533A>G (p.Tyr178Cys)	104893776	RHO	[]	['GGATGYACCTG AGGACAGGCAGG']	['Retinitis pigmentosa 4']
NM_000539.3(RHO):c. .569A>G (p.Asp190Gly)	104893777	RHO	[]	[]	['Retinitis pigmentosa 4']
NM_000539.3(RHO):c. .44A>G (p.Asn15Ser)	104893786	RHO	[]	[]	['Retinitis pigmentosa', 'Retinitis pigmentosa 4']
NM_001814.4(CTSC): c.1235A>G (p.Tyr412Cys)	28937571	CTSC	[]	[]	['Periodontitis, aggressive, 1']
NM_001701.3(BAAT) :c.226A>G (p.Met76Val)	28937579	BAAT	[]	[]	['Hypercholeltemia, familial']
NM_001257342.1(BC S1L):c.232A>G (p.Ser78Gly)	28937590	BCS1L	[]	['GACACYGAGGT GCTGAGTACGGG', 'CGACACYGAGGT GCTGAGTACGG']	['GRACILE syndrome']
NM_004407.3(DMP1) :c.1A>G (p.Met1Val)	104893834	DMP1	[]	[]	['Autosomal recessive hypophosphatemic vitamin D refractory rickets']
NM_000406.2(GNRH R):c.317A>G (p.Gln106Arg)	104893836	GNRHR	[]	[]	['Hypogonadotropic hypogonadism']
NM_172250.2(MMAA (p.Tyr207Cys)	104893849	MMAA	[]	[]	['Methylmalonic aciduria cblA type']
NM_000320.2(QDPR) :c.449A>G (p.Tyr150Cys)	104893866	QDPR	[]	['TGCCGYACCCGA TCATACCTGGG', 'ATGCCGYACCCG ATCATACCTGG']	['Dihydropteridine reductase deficiency']
NM_015629.3(PRPF3 1):c.527+3A>G	587776590	PRPF31	[]	['GACAYACCCCTG GGTGGTGGAGG', 'GCGGACAYACCC CTGGGTGGTGG']	['Retinitis pigmentosa 11']
NM_000112.3(SLC26 A2):c.1273A>G (p.Asn425Asp)	104893920	SLC26A2	[]	[]	['Diastrophic dysplasia', 'Achohdrogenesis, type IB']
NM_000344.3(SMN1) :c.815A>G (p.Tyr272Cys)	104893922	SMN1	[]	[]	['Werdnig-Hoffmann disease']
NM_000344.3(SMN1) :c.784A>G	104893932	SMN1	[]	[]	['Kugelberg- Wclander disease']

(p.Ser262Gly)					
NM_000409.3(GUCA1A):c.296A>G (p.Tyr99Cys)	104893967	GUCA1A	[]	[]	['Cone dystrophy 3']
NM_182548.3(LHFPL5):c.380A>G (p.Tyr127Cys)	104893975	LHFPL5	[]	[]	['Deafness, autosomal recessive 67']
NM_001024630.3(RUNX2):c.598A>G (p.Thr200Ala)	104893993	RUNX2	[]	[]	['Cleidocranial dysostosis', 'Cleidocranial dysplasia, forme fruste, dental anomalies only']
NM_000162.3(GCK):c.641A>G (p.Tyr214Cys)	104894015	GCK	[]	['GTAGYAGCAGG AGATCATCGTGG']	['Hyperinsulinemic hypoglycemia familial 3']
NM_001002010.2(NT5C3A):c.686A>G (p.Asn229Ser)	104894028	NT5C3A	[]	[]	['Uridine 5-prime monophosphate hydrolase deficiency, hemolytic anemia due to']
NM_203288.1(RP9):c.509A>G (p.Asp170Gly)	104894039	RP9	[]	[]	['Retinitis pigmentosa 9']
NM_000474.3(TWIST1):c.466A>G (p.Ile156Val)	104894059	TWIST1	[]	[]	['Saethre-Chotzen syndrome']
NM_000532.4(PCCB):c.1606A>G (p.Asn536Asp)	202247823	PCCB	[]	['ATATYTG CATGT TTTCTCCAAGG']	['Propionic acidemia', 'not provided']
NM_021615.4(CHST6):c.521A>G (p.Lys174Arg)	28937877	CHST6	[]	[]	['Macular corneal dystrophy Type 1']
NM_178138.4(LHX3):c.332A>G (p.Tyr111Cys)	104894117	LHX3	['CAGGTGG YACACGAA GTCCTGGG']	['CAGGTGGYACA CGAAGTCCTGGG']	['Pituitary hormone deficiency, combined 3']
NM_004897.4(MINPP1):c.809A>G (p.Gln270Arg)	104894171	MINPP1	[]	[]	['Thyroid cancer, follicular']
NM_000073.2(CD3G):c.1A>G (p.Met1Val)	104894199	CD3G	[]	['CCAYGTCAGTCT CTGTCCTCCGG']	['Immunodeficiency 17']
NM_001885.2(CRYAB):c.358A>G (p.Arg120Gly)	104894201	CRYAB	[]	[]	['Alpha-B crystallinopathy']
NM_001814.4(CTSC):c.857A>G (p.Gln286Arg)	104894208	CTSC	[]	['CTCCYGAGGGCT TAGGATTGGG', 'CCTCCYGAGGGC TTAGGATTGGG', 'ACCTCCYGAGGG CTTAGGATTGG']	['Papillon-Lefevre syndrome', 'Haim-Munk syndrome']
NM_001814.4(CTSC):c.1040A>G (p.Tyr347Cys)	104894211	CTSC	[]	['TCCTACAYAGTG TACTCAGAGG']	['Papillon-Lefevre syndrome', 'Periodontitis, aggressive, 1']
NM_012193.3(FZD4):c.766A>G	104894223	-	['GAAATAY GATGGGGC']	['GAAATAYGATG GGGCGCTCAGGG']	['Retinopathy of prematurity']

(p.Ile256Val)			GCTCAGGG , 'AGAAATA YGATGGGG CGCTCAGG '	'AGAAATAYGATG GGGCGCTCAGG']	
NM_005343.2(HRAS) :c.350A>G (p.Lys117Arg)	104894227	-	[]	[]	['Costello syndrome']
NM_145014.2(HYLS1)):c.632A>G (p.Asp211Gly)	104894232	-	[]	[]	['Hydrolethalus syndrome']
NM_000525.3(KCNJ11)):c.776A>G (p.His259Arg)	104894248	KCNJ11	['GACAYGG TAGATGAT CAGCGGG '	['GACAYGGTAGA TGATCAGCGGG', 'TGACAYGGTAGA TGATCAGCGGG', 'ATGACAYGGTAG ATGATCAGCGG']	['Islet cell hyperplasia']
NM_000317.2(PTS):c. 155A>G (p.Asn52Ser)	104894275	PTS	['TAAYTGT GCCCATGG CCATTTGG' '	['TAAYTGTGCCCA TGGCCATTTGG']	['6-pyruvoyl- tetrahydropterin synthase deficiency']
NM_000317.2(PTS):c. 139A>G (p.Asn47Asp)	104894278	PTS	[]	[]	['Hyperphenylalanin emia, bh4-deficient, a, due to partial pts deficiency']
NM_000317.2(PTS):c. 347A>G (p.Asp116Gly)	104894279	PTS	[]	[]	['Hyperphenylalanin emia, bh4-deficient, a, due to partial pts deficiency']
NM_022051.2(EGLN1)):c.1121A>G (p.His374Arg)	119476045	EGLN1	[]	[]	['Erythrocytosis, familial, 3']
NM_015915.4(ATL1): c.773A>G (p.His258Arg)	119476048	ATL1	[]	[]	['Spastic paraplegia 3']
NM_000448.2(RAG1): c.2735A>G (p.Tyr912Cys)	104894290	RAG1	[]	['CTGYACTGGCAG AGGGATTCTGG']	['Histiocytic medullary reticulosis']
NM_000448.2(RAG1): c.1286A>G (p.Asp429Gly)	104894292	RAG1	[]	[]	['Histiocytic medullary reticulosis']
NM_003002.3(SDHD) :c.341A>G (p.Tyr114Cys)	104894304	SDHD	[]	[]	['Hereditary Paranglioma- Pheochromocytoma Syndromes', 'Parangliomas 1']
NM_015141.3(GPD1L)):c.370A>G (p.Ile124Val)	72552293	GPD1L	[]	[]	['Brugada syndrome 2', 'Primary familial hypertrophic cardiomyopathy', 'Long QT syndrome', 'Sudden infant death syndrome', 'Cardiomyopathy']
NM_020661.2(AICDA)):c.415A>G (p.Met139Val)	104894322	AICDA	[]	[]	['Immunodeficiency with hyper IgM type 2']

NM_014365.2(HSPB8):c.421A>G (p.Lys141Glu)	104894351	HSPB8	[]	[]	['Charcot-Marie-Tooth disease', 'Distal hereditary motor neuropathy type 2A']
NM_000217.2(KCNA1):c.676A>G (p.Thr226Ala)	104894354	KCNA1	[]	['GCGYTTCCACGATGAAGAAGGG', 'AGCGYTTCCACGATGAAGAAGGG', 'CAGCGYTTCCACGATGAAGAAGG']	['Episodic ataxia type 1']
NM_014239.3(EIF2B2):c.638A>G (p.Glu213Gly)	104894425	EIF2B2	[]	['AGTTGTCYCAATACCTGCTTTGG']	['Leukoencephalopathy with vanishing white matter', 'Ovariokodystrophy']
NM_002408.3(MGAT2):c.785A>G (p.His262Arg)	104894447	MGAT2	[]	[]	['Carbohydrate-deficient glycoprotein syndrome type II']
NM_002408.3(MGAT2):c.952A>G (p.Asn318Asp)	104894448	MGAT2	[]	[]	['Carbohydrate-deficient glycoprotein syndrome type II']
NM_000270.3(PNP):c.383A>G (p.Asp128Gly)	104894450	PNP	[]	['ATAYCTCCAACCTCAAACCTTGGG', 'GATAYCTCCAACCTCAAACCTTGG']	['Purine-nucleoside phosphorylase deficiency']
NM_000270.3(PNP):c.575A>G (p.Tyr192Cys)	104894452	PNP	[]	[]	['Purine-nucleoside phosphorylase deficiency']
NM_005982.3(SIX1):c.386A>G (p.Tyr129Cys)	104894478	SIX1	[]	[]	['Melnick-Fraser syndrome', 'Branchiootic syndrome 3']
NM_000101.3(CYBA):c.281A>G (p.His94Arg)	104894510	CYBA	[]	[]	['Granulomatous disease, chronic, autosomal recessive, cytochrome b-negative']
NM_024887.3(DHDDS):c.124A>G (p.Lys42Glu)	147394623	DHDDS	[]	['GGCACTYCTTGGCATAGCGACGG']	['Retinitis pigmentosa 59']
NM_024006.5(VKORC1):c.172A>G (p.Arg58Gly)	104894541	VKORC1	[]	[]	['Warfarin response']
NM_001128085.1(ASPA):c.692A>G (p.Tyr231Cys)	104894550	-	[]	[]	['Spongy degeneration of central nervous system']
NM_001128085.1(ASPA):c.71A>G (p.Glu24Gly)	104894551	-	[]	[]	['Spongy degeneration of central nervous system']
NM_000019.3(ACAT1):c.472A>G (p.Asn158Asp)	148639841	ACAT1	[]	[]	['Deficiency of acetyl-CoA acetyltransferase', 'not provided']
NM_014254.2(TMEM5):c.1016A>G	150736997	TMEM5	[]	[]	['Congenital muscular dystrophy-']

(p.Tyr339Cys)					dystroglycanopathy with brain and eye anomalies, type A10']
NM_005557.3(KRT16):c.374A>G (p.Asn125Ser)	60723330	KRT16	[]	['GCGGTCAYTGA GGTTCGCATGG']	['Pachyonychia congenita, type 1', 'Palmoplantar keratoderma, nonepidermolytic, focal', 'not provided']
NM_005450.4(NOG):c.665A>G (p.Tyr222Cys)	104894602	NOG	[]	[]	['Tarsal carpal coalition syndrome', 'Cushing symphalangism']
NM_030665.3(RAI1):c.4685A>G (p.Gln1562Arg)	104894634	RAI1	[]	['CTGCTGCYGTCTG TCGTCGCTTGG']	['Smith-Magenis syndrome']
NM_000346.3(SOX9):c.517A>G (p.Lys173Glu)	104894647	SOX9	[]	[]	['Acampomelic campomelic dysplasia']
NM_024301.4(FKRP):c.926A>G (p.Tyr309Cys)	104894679	FKRP	[]	[]	['Congenital muscular dystrophy-dystroglycanopathy without mental retardation, type B5']
NM_000495.4(COL4A5):c.2394A>G (p.Lys798=)	281874691	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_001604.5(PAX6):c.1075-2A>G	794726661	PAX6	[]	[]	['Congenital aniridia']
NM_000363.4(TNNI3):c.569A>G (p.Asp190Gly)	104894728	TNNI3	[]	[]	['Familial restrictive cardiomyopathy 1', 'Familial hypertrophic cardiomyopathy 7']
NM_000363.4(TNNI3):c.532A>G (p.Lys178Glu)	104894730	TNNI3	[]	['CCTYCTTCACCT GCTTGAGGTGG', 'CCTCCTYCTTCAC CTGCTTGAGG']	['Familial restrictive cardiomyopathy 1']
NM_000054.4(AVPR2):c.839A>G (p.Tyr280Cys)	104894752	AVPR2	[]	[]	['Nephrogenic diabetes insipidus, X-linked']
NM_000074.2(CD40LG):c.386A>G (p.Glu129Gly)	104894772	CD40LG	[]	[]	['Immunodeficiency with hyper IgM type 1']
NM_000495.4(COL4A5):c.4977-2A>G	281874752	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_001165963.1(SCN1A):c.5264A>G (p.Asp1755Gly)	794726722	-	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_000495.4(COL4A5):c.547-2A>G	281874756	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.610-2A>G	281874758	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_032520.4(GNPTG):c.610-2A>G	193302855	GNPTG	['CCCYGAA GGTGGAGG ATGCAGGG']	['CCCYGAAGGTG GAGGATGCAGGG', 'GCCCYGAAGGTG']	['Mucopolidosis III Gamma']

				GAGGATGCAGG']	
NM_002049.3(GATA1):c.653A>G (p.Asp218Gly)	104894816	GATA1	[]	['GTCCTGYCCCTC CGCCACAGTGG']	['GATA-1-related thrombocytopenia with dyserythropoiesis']
NM_000157.3(GBA):c.680A>G (p.Asn227Ser)	364897	GBA	['CCAYTGG TCTTGAGC CAAGTGGG , 'TCCAYTGG TCTTGAGC CAAGTGG']	['CCAYTGGTCTTG AGCCAAGTGGG', 'TCCAYTGGTCTT GAGCCAAGTGG']	['Gaucher disease', 'Subacute neuronopathic Gaucher disease', 'Gaucher disease, type 1']
NM_001097642.2(GJB1):c.194A>G (p.Tyr65Cys)	104894819	GJB1	[]	[]	['X-linked hereditary motor and sensory neuropathy']
NM_000166.5(GJB1):c.614A>G (p.Asn205Ser)	104894822	GJB1	[]	[]	['X-linked hereditary motor and sensory neuropathy']
NM_001165963.1(SCN1A):c.747T>G (p.Asp249Glu)	773407463	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_000169.2(GLA):c.886A>G (p.Met296Val)	104894830	-	[]	[]	['Fabry disease', 'Fabry disease, cardiac variant']
NM_000169.2(GLA):c.101A>G (p.Asn34Ser)	104894835	-	[]	[]	['Fabry disease']
NM_001165963.1(SCN1A):c.1662+3A>G	794726773	SCN1A	[]	['GTGCCAYACCTG GTGTGGGGAGG']	['Severe myoclonic epilepsy in infancy']
NM_000169.2(GLA):c.1228A>G (p.Thr410Ala)	104894852	-	[]	[]	['Fabry disease']
NM_000202.6(IDS):c.404A>G (p.Lys135Arg)	104894861	IDS	[]	['AAAGACTYTTCC CACCGACATGG']	['Mucopolysaccharid osis, MPS-II']
NM_001165963.1(SCN1A):c.383+1A>G	794726803	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_000266.3(NDP):c.131A>G (p.Tyr44Cys)	104894870	NDP	[]	[]	['Atrophia bulborum hereditaria']
NM_000266.3(NDP):c.125A>G (p.His42Arg)	104894874	NDP	[]	['TGGYGCCTCATG CAGCGTCGAGG']	[]
NM_001128227.2(GNE):c.604A>G (p.Met202Val)	121908634	GNE	[]	[]	['Inclusion body myopathy 2']
NM_001165963.1(SCN1A):c.3880-2A>G	794726816	-	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_001165963.1(SCN1A):c.1046A>G (p.Tyr349Cys)	794726844	SCN1A	['ACATAYA TCCCTCTG GACATTGG ']	['ACATAYATCCCT CTGGACATTGG']	['Severe myoclonic epilepsy in infancy']
NM_001015877.1(PHF6):c.700A>G (p.Lys234Glu)	104894917	PHF6	[]	[]	['Borjeson- Forssman-Lehmann syndrome']
NM_001015877.1(PHF6):c.686A>G (p.His229Arg)	104894918	PHF6	[]	[]	['Borjeson- Forssman-Lehmann syndrome']
NM_001015877.1(PHF6):c.769A>G (p.Arg257Gly)	104894919	PHF6	['CTCYTGA TGTTGTTG TGAGCTGG ']	['CTCYTGATGTTG TTGTGAGCTGG']	['Borjeson- Forssman-Lehmann syndrome']

NM_000307.4(POU3F4):c.1000A>G (p.Lys334Glu)	104894922	POU3F4	[]	[]	['Deafness, X-linked 2']
NM_000330.3(RS1):c.667T>C (p.Cys223Arg)	104894929	-	[]	[]	['Juvenile retinoschisis']
NM_003413.3(ZIC3):c.1213A>G (p.Lys405Glu)	104894962	ZIC3	['TGTGTTY GCGCAGGG AGCTCGGG', 'ATGTGTTY GCGCAGGG AGCTCGG']	['TGTGTTYGCGCAGGGAGCTCGGG', 'ATGTGTTYGCGCAGGGAGCTCGG']	['Heterotaxy, visceral, X-linked']
NM_002420.5(TRPM1):c.296T>C (p.Leu99Pro)	191205969	TRPM1	[]	['AAGCYCTTAATA TCTGTGCATGG']	['Congenital stationary night blindness, type 1C']
NM_004006.2(DMD):c.1150-2A>G	794727030	DMD	[]	[]	['Duchenne muscular dystrophy', 'Becker muscular dystrophy']
NM_203290.2(POLR1C):c.221A>G (p.Asn74Ser)	371802902	POLR1C	[]	[]	['LEUKODYSTROPHY, HYPOMYELINATING, 11']
NM_019109.4(ALG1):c.1188-2A>G	794727073	ALG1	[]	['TAAACYGCAGAGAGAACCAAGGG', 'GTAAACYGCAGAGAGAACCAAGG']	['Congenital disorder of glycosylation type 1K']
NM_004463.2(FGD1):c.2016-2A>G	794727099	FGD1	[]	[]	['Aarskog syndrome']
NM_024110.4(CARD14):c.425A>G (p.Glu142Gly)	281875213	CARD14	[]	[]	['Psoriasis susceptibility 2', 'not provided']
NM_001004334.3(GPR179):c.659A>G (p.Tyr220Cys)	281875236	GPR179	[]	['CCCACAYATCCATCTGCCTGCGG']	['Congenital stationary night blindness, type 1E', 'not provided']
NM_018965.3(TREM2):c.401A>G (p.Asp134Gly)	28939079	TREM2	['CCGGTGA YCCAGGGG GTCTATGG']	['TGAYCCAGGGGTCTATGGGAGG', 'CGGTGAYCCAGGGGTCTATGGG', 'CCGGTGAYCCAGGGGTCTATGG']	['Polycystic lipomembranous osteodysplasia with sclerosing leukoencephalopathy']
NM_015915.4(ATL1):c.1222A>G (p.Met408Val)	28939094	ATL1	[]	['CACCCAYCTTCTTCACCCCTCGG']	['Spastic paraplegia 3']
NM_002437.4(MPV17):c.186+2T>C	147952488	MPV17	['TGGYAAGTTCTCCCC TCAACAGG']	['TGGYAAGTTCTCCCCCTCAACAGG']	['Navajo neurohepatopathy', 'not provided']
NM_005359.5(SMAD4):c.1500A>G (p.Ile500Met)	281875320	SMAD4	['TGAGYATGCATAAGC GACGAAGG']	['TGAGYATGCATAAGCGACGAAGG']	['Myhre syndrome', 'not provided']
NM_005359.5(SMAD4):c.1498A>G (p.Ile500Val)	281875322	SMAD4	['TGAGTAYGCATAAGC GACGAAGG']	['TGAGTAYGCATAAGCGACGAAGG']	['Hereditary cancer-predisposing syndrome', 'Myhre syndrome', 'not provided']

NM_005359.5(SMAD4):c.989A>G (p.Glu330Gly)	281875324	SMAD4	[]	['ATCCATTYCAAA GTAAGCAATGG']	['Juvenile polyposis syndrome', 'Hereditary cancer-predisposing syndrome', 'not provided']
NM_000342.3(SLC4A1):c.166A>G (p.Lys56Glu)	5036	SLC4A1	[]	[]	[]
NM_000518.4(HBB):c.*113A>G	33985472	HBB	[]	[]	[]
NM_001127255.1(NLRP7):c.2738A>G (p.Asn913Ser)	104895503	-	['TCTGGYT GATACTCA AGTCCAGG ']	['TCTGGYTGATAC TCAAGTCCAGG']	['Hydatidiform mole']
NM_000037.3(ANK1):c.-108T>C	77173848	ANK1	[]	['GGGCCYGGCCC GCACGTACAGG']	['Spherocytosis, type 1, autosomal recessive']
NM_201631.3(TGM5):c.763T>C (p.Trp255Arg)	115677373	TGM5	['TGCGGAG YGGACGG GCAGCGTG G']	['TGCGGAGYGGA CGGGCAGCGTGG']	['Peeling skin syndrome, acral type']
NM_020435.3(GJC2):c.-167A>G	587776888	GJC2	[]	[]	['Leukodystrophy, hypomyelinating, 2']
NM_130466.3(UBE3B):c.1A>G (p.Met1Val)	672601304	UBE3B	[]	[]	['Kaufman oculocerebrofacial syndrome']
NM_022124.5(CDH23):c.146-2A>G	794727649	-	[]	[]	['Usher syndrome, type 1D']
NM_014191.3(SCN8A):c.667A>G (p.Arg223Gly)	672601319	SCN8A	[]	[]	['Early infantile epileptic encephalopathy 13']
NM_001164405.1(BHLHA9):c.211A>G (p.Asn71Asp)	672601337	BHLHA9	[]	[]	['Syndactyly type 9']
NM_021830.4(C10orf2):c.1754A>G (p.Asn585Ser)	672601360	C10orf2	[]	[]	['Perrault syndrome 5']
NM_002887.3(RARS):c.5A>G (p.Asp2Gly)	672601372	RARS	[]	[]	['Leukodystrophy, hypomyelinating, 9']
NM_002887.3(RARS):c.1A>G (p.Met1Val)	672601375	RARS	[]	[]	['Leukodystrophy, hypomyelinating, 9']
NM_001943.3(DSG2):c.1880-2A>G	397514038	DSG2	[]	[]	['Arrhythmogenic right ventricular cardiomyopathy, type 10', 'Cardiomyopathy']
NM_024422.4(DSC2):c.631-2A>G	397514042	DSC2	[]	[]	['Arrhythmogenic right ventricular cardiomyopathy, type 11', 'Cardiomyopathy']
NM_001159772.1(CANT1):c.671T>C (p.Leu224Pro)	150181226	CANT1	[]	['CGTCYGTACGTG GGCGGCCTGGG', 'GCGTCYGTACGT GGCGGCCTGG']	['Desbuquois syndrome']
NM_031418.2(ANO3):c.2053A>G (p.Ser685Gly)	587776923	ANO3	[]	[]	['Dystonia 24']

NM_014191.3(SCN8A):c.5302A>G (p.Asn1768Asp)	202151337	SCN8A	[]	[]	['Early infantile epileptic encephalopathy 13']
NM_000367.3(TPMT):c.719A>G (p.Tyr240Cys)	1142345	TPMT	[]	[]	['Thiopurine methyltransferase deficiency']
NM_003907.2(EIF2B5):c.271A>G (p.Thr91Ala)	28939717	EIF2B5	['AAATGYT TCCTGTAC ACCTGTGG']	['AAATGYTTCCTG TACACCTGTGG']	['Leukoencephalopathy with vanishing white matter']
NM_004006.2(DMD):c.10554-2A>G	794727890	DMD	[]	[]	['Duchenne muscular dystrophy', 'Becker muscular dystrophy', 'Dilated cardiomyopathy 3B']
NM_000084.4(CLCN5):c.815A>G (p.Tyr272Cys)	273585644	CLCN5	[]	[]	['Dent disease 1']
NM_000084.4(CLCN5):c.1637A>G (p.Lys546Arg)	273585649	CLCN5	[]	[]	['Dent disease 1']
NM_000041.3(APOE):c.237-2A>G	397514253	APOE	[]	['CGCCCYGCGGCC GAGAGGGCGGG', 'GCGCCCYGCGGCC GAGAGGGCGGG']	['Familial type 3 hyperlipoproteinemia']
NM_000155.3(GALT):c.940A>G (p.Asn314Asp)	2070074	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase', 'not provided']
NM_005045.3(RELN):c.2288A>G (p.Asp763Gly)	794727998	RELN	[]	[]	['EPILEPSY, FAMILIAL TEMPORAL LOBE, 7']
NM_001914.3(CYB5A):c.130-2A>G	794728010	CYB5A	[]	[]	['Methemoglobinemia type 4']
NM_001613.2(ACTA2):c.1A>G (p.Met1Val)	794728019	ACTA2	[]	[]	['Thoracic aortic aneurysms and aortic dissections']
NM_000109.3(DMD):c.1700T>C (p.Leu567Pro)	370644567	DMD	[]	[]	['Becker muscular dystrophy', 'Exertional myalgia, muscle stiffness and myoglobinuria', 'not provided']
NM_000060.3(BTD):c.194A>G (p.His65Arg)	397514341	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c.278A>G (p.Tyr93Cys)	397514348	BTD	[]	['GTTCAAYAGATGT CAAGGTTCTGG']	['Biotinidase deficiency']
NM_000060.3(BTD):c.356A>G (p.Asn119Ser)	397514353	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c.364A>G (p.Arg122Gly)	397514354	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c.515A>G (p.Asn172Ser)	397514366	BTD	[]	[]	['Biotinidase deficiency']
NM_000060.3(BTD):c	397514370	BTD	[]	[]	['Biotinidase

.583A>G (p.Asn195Asp)					deficiency']
NM_000060.3(BTD):c .584A>G (p.Asn195Ser)	397514371	BTD	[]	[]	['Biotinidase deficiency', 'not provided']
NM_000060.3(BTD):c .641A>G (p.Asn214Ser)	397514377	BTD	['AGAGGYT GTGTTTAC GGTAGCGG ']	['AGAGGYTGTGTT TACGGTAGCGG']	['Biotinidase deficiency']
NM_000061.2(BTK):c .1288A>G (p.Lys430Glu)	128620184	BTK	['TCTYGAT GGCCACGT CGTACTGG' ']	['TCTYGATGGCCA CGTCGTAAGG']	['X-linked agammaglobulinemi a']
NM_001002294.2(FM O3):c.923A>G (p.Glu308Gly)	2266780	FMO3	[]	[]	['Trimethylaminuria' ']
m.15579A>G	207460002	MT-CYB	[]	[]	['Multisystem disorder']
NM_000060.3(BTD):c .1313A>G (p.Tyr438Cys)	397514415	BTD	[]	['GGCAYACAGCT CTTTGGATAAGG']	['Biotinidase deficiency']
NM_000060.3(BTD):c .1619A>G (p.Tyr540Cys)	397514431	BTD	[]	[]	['Biotinidase deficiency']
NM_000023.2(SGCA) :c.410A>G (p.Glu137Gly)	397514451	SGCA	[]	[]	['Limb-girdle muscular dystrophy, type 2D']
NM_004813.2(PEX16) :c.992A>G (p.Tyr331Cys)	397514472	PEX16	['AAGYAGA TTTTCTGC CAGGTGGG ' ' 'GAAGYAG ATTTTCTG CCAGGTGG ' ' 'GTAGAAG YAGATTTT CTGCCAGG ' ']	['AAGYAGATTTTC TGCCAGGTGGG', 'GAAGYAGATTTT CTGCCAGGTGG', 'GTAGAAGYAGAT TTTCTGCCAGG']	['Peroxisome biogenesis disorder 8B']
NM_000933.3(PLCB4) :c.1868A>G (p.Tyr623Cys)	397514480	PLCB4	[]	[]	['Auriculocondylar syndrome 1', 'Auriculocondylar syndrome 2']
NM_005340.6(HINT1) :c.152A>G (p.His51Arg)	397514491	HINT1	['AAAAYGT GTTGGTGC TTGAGGGG ' ' 'GAAAAYG TGTTGGTG CTTGAGGG ' ' 'AGAAAAY GTGTTGGT GCTTGAGG ' ']	['AAAAYGTGTTG GTGCTTGAGGGG', 'GAAAAYGTGTTG GTGCTTGAGGG', 'AGAAAAYGTGTT GGTGCTTGAGG']	['Gamstorp-Wohlfart syndrome']
NM_007171.3(POMT 1):c.430A>G (p.Asn144Asp)	397514501	POMT1	[]	['GAGCATYCTCTG TTTCAAAGAGG']	['Limb-girdle muscular dystrophy- dystroglycanopathy, type C1']

NM_003863.3(DPM2):c.68A>G (p.Tyr23Cys)	397514503	DPM2	['TG TAGYA GGTGAAGA TGATCAGG ']	['TG TAGYAGGTG AAGATGATCAGG ']	['Congenital disorder of glycosylation type 1u']
NM_174917.4(ACSF3):c.1A>G (p.Met1Val)	370382601	ACSF3	[]	['GGCAGCAYTGC ACTGACAGGCGG ']	['not provided']
NM_183075.2(CYP2U1):c.1139A>G (p.Glu380Gly)	397514514	CYP2U1	[]	[]	['Spastic paraplegia 56, autosomal recessive']
NM_000344.3(SMN1):c.389A>G (p.Tyr130Cys)	397514517	SMN1	[]	[]	['Kugelberg-Welander disease']
NM_000138.4(FBN1):c.4337-2A>G	794728216	FBN1	[]	[]	['Thoracic aortic aneurysms and aortic dissections']
NM_012082.3(ZFPM2):c.2209A>G (p.Lys737Glu)	397514521	-	[]	[]	['Double outlet right ventricle']
NM_001168272.1(ITPR1):c.1759A>G (p.Asn587Asp)	397514536	ITPR1	[]	[]	['Spinocerebellar ataxia 29']
NM_178012.4(TUBB2B):c.767A>G (p.Asn256Ser)	397514568	TUBB2B	[]	[]	['Polymicrogyria, asymmetric']
NM_000531.5(OTC):c.155A>G (p.Glu52Gly)	72554317	OTC	[]	[]	['not provided']
NM_001866.2(COX7B):c.41-2A>G	397514584	COX7B	[]	[]	['Aplasia cutis congenita, reticuloliner, with microcephaly, facial dysmorphism, and other congenital anomalies']
NM_001099922.2(ALG13):c.339A>G (p.Ala113=)	397514587	ALG13	[]	[]	['Congenital disorder of glycosylation type 1s']
NM_000531.5(OTC):c.238A>G (p.Lys80Glu)	72554332	OTC	[]	['AAGGACTYCCCT TGCAATAAAGG']	['Ornithine carbamoyltransferase deficiency', 'not provided']
NM_001083614.1(EARS2):c.502A>G (p.Arg168Gly)	397514591	EARS2	[]	[]	['Combined oxidative phosphorylation deficiency 12']
NM_001083614.1(EARS2):c.193A>G (p.Lys65Glu)	397514595	EARS2	[]	[]	['Combined oxidative phosphorylation deficiency 12']
NM_198578.3(LRRK2):c.3364A>G (p.Ile1122Val)	34805604	LRRK2	[]	[]	['Parkinson disease 8, autosomal dominant']
NM_033109.4(PNPT1):c.1160A>G (p.Gln387Arg)	397514598	PNPT1	[]	[]	['Combined oxidative phosphorylation deficiency 13']
NM_033109.4(PNPT1):c.1424A>G (p.Glu475Gly)	397514599	PNPT1	[]	['GACTYCAGATGT AACTCTTATGG']	['Deafness, autosomal recessive 70']

NM_000531.5(OTC):c.277A>G (p.Thr93Ala)	72554344	OTC	[]	[]	['not provided']
NM_000390.2(CHM):c.1520A>G (p.His507Arg)	397514603	CHM	[]	[]	['Choroideremia']
NM_181690.2(AKT3):c.686A>G (p.Asn229Ser)	397514605	AKT3	[]	[]	['Megalencephaly-polymicrogyria-polydactyly-hydrocephalus syndrome 2']
NM_006567.3(FARS2):c.431A>G (p.Tyr144Cys)	397514610	FARS2	[]	[]	['Mitochondrial encephalomyopathy', 'Combined oxidative phosphorylation deficiency 14', 'Global developmental delay']
NM_000531.5(OTC):c.377A>G (p.Asp126Gly)	72554358	OTC	[]	[]	['not provided']
NM_005609.2(PYGM):c.152A>G (p.Asp51Gly)	397514631	PYGM	[]	[]	['Glycogen storage disease, type V']
NM_000130.4(F5):c.1601G>A (p.Arg534Gln)	6025	F5	[]	[]	['Recurrent abortion', 'Thrombophilia due to factor V Leiden']
NM_000108.4(DLD):c.1444A>G (p.Arg482Gly)	397514650	DLD	[]	['GACTCYAGCTATATCTTCACAGG']	['Maple syrup urine disease, type 3']
NM_138554.4(TLR4):c.896A>G (p.Asp299Gly)	4986790	TLR4	[]	[]	['Endotoxin hyporesponsiveness']
NM_003156.3(STIM1):c.251A>G (p.Asp84Gly)	397514675	STIM1	[]	['TTCCACAYCCACATCACCATTGG']	['Myopathy with tubular aggregates']
NM_003156.3(STIM1):c.326A>G (p.His109Arg)	397514677	STIM1	[]	[]	['Myopathy with tubular aggregates']
NM_000238.3(KCNH2):c.1900A>G (p.Thr634Ala)	794728377	KCNH2	[]	[]	['Cardiac arrhythmia']
NM_000238.3(KCNH2):c.1913A>G (p.Lys638Arg)	794728378	KCNH2	[]	['ATCYTCTCTGAGTTGGTGTGGG', 'GATCYTCTCTGAGTTGGTGTGG']	['Cardiac arrhythmia']
NM_001457.3(FLNB):c.604A>G (p.Met202Val)	121908895	FLNB	[]	[]	['Atelosteogenesis type 1']
NM_001893.4(CSNK1D):c.137A>G (p.His46Arg)	397514693	CSNK1D	[]	[]	['Advanced sleep phase syndrome, familial, 2']
NM_002163.2(IRF8):c.322A>G (p.Lys108Glu)	397514710	IRF8	[]	[]	['Monocyte and dendritic cell deficiency, autosomal recessive']

NM_002163.2(IRF8):c.238A>G (p.Thr80Ala)	397514711	IRF8	[]	['AACCTCGYCTTC CAAGTGGCTGG']	['Autosomal dominant CD11C+/CD1C+ dendritic cell deficiency']
NM_001127217.2(SMAD9):c.127A>G (p.Lys43Glu)	397514715	SMAD9	[]	[]	['Primary pulmonary hypertension 2']
NM_001035.2(RYR2):c.12290A>G (p.Asn4097Ser)	794728784	RYR2	[]	[]	['not provided']
NM_000388.3(CASR):c.85A>G (p.Lys29Glu)	397514729	CASR	[]	['CCCCCTYCTTTT GGGCTCGCTGG']	['Hypocalcemia, autosomal dominant 1, with bartter syndrome']
NM_003793.3(CTSF):c.962A>G (p.Gln321Arg)	397514731	CTSF	[]	[]	['Ceroid lipofuscinosis, neuronal, 13']
NM_173076.2(ABCA12):c.4139A>G (p.Asn1380Ser)	28940269	ABCA12	[]	[]	['Autosomal recessive congenital ichthyosis 4A']
NM_017890.4(VPS13B):c.8978A>G (p.Asn2993Ser)	28940272	VPS13B	['TCAYTGA TAAGCAGG GCCCAGGG , 'TTCAYTGA TAAGCAGG GCCCAGG']	['TCAYTGATAAGC AGGGCCCAGGG', 'TTCAYTGATAAG CAGGGCCCAGG']	['Cohen syndrome', 'not specified']
NM_022114.3(PRDM16):c.2447A>G (p.Asn816Ser)	397514743	PRDM16	[]	['GCCGCCGYTTTG GCTGGCACGGG']	['Left ventricular noncompaction 8']
NM_005689.2(ABCB6):c.508A>G (p.Ser170Gly)	397514757	ABCB6	[]	['TGGGCYGTCCA AGACACCAGGG', 'GTGGGCYGTCC AAGACACCAGG']	['Dyschromatosis universalis hereditaria 3']
NM_015335.4(MED13L):c.752A>G (p.Glu251Gly)	28940309	MED13L	[]	[]	['Transposition of great arteries']
NM_152443.2(RDH12):c.677A>G (p.Tyr226Cys)	28940313	RDH12	[]	['CACTGCGYAGGT GGTGACCCCGG']	['Leber congenital amaurosis 13']
NM_000517.4(HBA2):c.96-2A>G	41457746	HBA2	[]	[]	[]
NM_000218.2(KCNQ1):c.1787A>G (p.Glu596Gly)	794728538	KCNQ1	[]	['GTCTYCTACTCG GTTTCAGGCGGG', 'GTCTYCTACTCG GTTTCAGGCGG']	['Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.605A>G (p.Asp202Gly)	794728569	KCNQ1	[]	['AGGYCTGTGGA GTGCAGGAGAGG']	['Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.1515-2A>G	794728573	KCNQ1	[]	['GCCYGCAGTGG AGAGAGGAGAGG']	['Cardiac arrhythmia']
NM_000498.3(CYP11B2):c.1492A>G (p.Thr498Ala)	72554626	-	[]	[]	['Corticosterone methyloxidase type 2 deficiency']
NM_000169.2(GLA):c.644A>G (p.Asn215Ser)	28935197	-	[]	[]	['Fabry disease', 'not provided']

NM_000218.2(KCNQ1):c.1085A>G (p.Lys362Arg)	12720458	KCNQ1	[]	[]	['Congenital long QT syndrome', 'Cardiac arrhythmia', 'Long QT syndrome, LQT1 subtype']
NM_001035.2(RYR2):c.12533A>G (p.Asn4178Ser)	794728787	RYR2	[]	[]	['not provided']
NM_003494.3(DYSF):c.3349-2A>G	370874727	DYSF	[]	['CCGCCCYGGAG ACACGAAGCTGG']	['Limb-girdle muscular dystrophy, type 2B']
NM_001035.2(RYR2):c.568A>G (p.Arg190Gly)	794728814	RYR2	[]	[]	['not provided']
NM_198056.2(SCN5A):c.2788-2A>G	794728859	SCN5A	[]	['ACCYGTCGAGAT AATGGGTCAGG']	['not provided']
NM_198056.2(SCN5A):c.4453A>G (p.Ile1485Val)	794728886	SCN5A	[]	[]	['not provided']
NM_198056.2(SCN5A):c.4462A>G (p.Thr1488Ala)	794728887	SCN5A	[]	['CCTCTGYCATGA AGATGTCCTGG']	['not provided']
NM_001927.3(DES):c.1324A>G (p.Thr442Ala)	794728995	DES	[]	[]	['not provided']
NM_001613.2(ACTA2):c.145A>G (p.Met49Val)	397515325	ACTA2	[]	[]	['Aortic aneurysm, familial thoracic 6']
NM_000782.4(CYP24A1):c.1226T>C (p.Leu409Ser)	6068812	CYP24A1	[]	[]	['Idiopathic hypercalcemia of infancy']
NM_000372.4(TYR):c.125A>G (p.Asp42Gly)	28940878	TYR	[]	['CTCCTGYCCCCG CTCCACGGTGG']	['Tyrosinase-negative oculocutaneous albinism', 'not provided']
NM_000372.4(TYR):c.1A>G (p.Met1Val)	28940881	TYR	[]	[]	['Tyrosinase-negative oculocutaneous albinism', 'Oculocutaneous albinism type 1B', 'not provided']
NM_000403.3(GALE):c.770A>G (p.Lys257Arg)	28940884	GALE	[]	[]	['UDPglucose-4-epimerase deficiency']
NM_000529.2(MC2R):c.761A>G (p.Tyr254Cys)	28940892	MC2R	['ACATGYA GCAGGCGC AGTAGGGG', 'GACATGY AGCAGGCG CAGTAGGG']	['ACATGYAGCAG GCGCAGTAGGGG', 'GACATGYAGCAG GCGCAGTAGGG', 'AGACATGYAGCA GCGCAGTAGG']	['ACTH resistance']
NM_000061.2(BTK):c.919A>G (p.Arg307Gly)	128621195	BTK	[]	[]	['X-linked agammaglobulinemia']
NM_000061.2(BTK):c.1766A>G	128621206	BTK	[]	[]	['X-linked agammaglobulinemia']

(p.Glu589Gly)					a']
NM_018486.2(HDAC8):c.539A>G (p.His180Arg)	397515416	HDAC8	[]	[]	['Cornelia de Lange syndrome 5']
NM_018486.2(HDAC8):c.1001A>G (p.His334Arg)	397515418	HDAC8	['CTCAYGATCTGG TCTGGGATCTCAGAGG']	['CTCAYGATCTGG GATCTCAGAGG']	['Cornelia de Lange syndrome 5']
NM_172107.2(KCNQ2):c.1636A>G (p.Met546Val)	397515420	KCNQ2	[]	['GCAYGACACTG CAGGGGGGTGGG', 'CGCAYGACACTG CAGGGGGGTGG', 'AACCGCAYGACA CTGCAGGGGGG']	['Early infantile epileptic encephalopathy 7']
NM_001410.2(MEGF8):c.7099A>G (p.Ser2367Gly)	397515428	MEGF8	[]	['GACYCCCGTGA AATGATTCCCGG']	['Carpenter syndrome 2']
NM_004247.3(EFTUD2):c.623A>G (p.His208Arg)	397515431	EFTUD2	[]	[]	['Growth and mental retardation, mandibulofacial dysostosis, microcephaly, and cleft palate']
NM_004572.3(PKP2):c.1171-2A>G	794729133	PKP2	[]	[]	['not provided']
NM_018972.2(GDAP1):c.368A>G (p.His123Arg)	397515442	GDAP1	[]	[]	['Charcot-Marie-Tooth disease type 2K']
NM_014795.3(ZEB2):c.3134A>G (p.His1045Arg)	397515449	ZEB2	[]	[]	['Mowat-Wilson syndrome']
NM_002336.2(LRP6):c.1298A>G (p.Asn433Ser)	397515473	LRP6	[]	[]	['Coronary artery disease, autosomal dominant 2']
NM_001015879.1(AURKC):c.379-2A>G	397515484	AURKC	[]	[]	['Infertility associated with multi-tailed spermatozoa and excessive DNA']
NM_000495.4(COL4A5):c.3107-4A>G	397515497	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_201631.3(TGM5):c.122T>C (p.Leu41Pro)	143601447	TGM5	[]	['TCAACCYCACCC TGTACTTCAGG']	['Peeling skin syndrome, acral type']
NM_013254.3(TBK1):c.1201A>G (p.Lys401Glu)	756751089	TBK1	[]	[]	['FRONTOTEMPORAL DEMENTIA AND/OR AMYOTROPHIC LATERAL SCLEROSIS 4']
NM_000207.2(INS):c.*59A>G	397515519	-	[]	['GGGCYTTATTCC ATCTCTCTCGG']	['Permanent neonatal diabetes mellitus']
NM_000370.3(TTPA):c.191A>G (p.Asp64Gly)	397515523	TTPA	[]	['CAGGYCCAGAT CGAAATCCC GGG', 'CCAGGYCCAGAT CGAAATCCC GGG']	['Ataxia with vitamin E deficiency']
NM_001006657.1(WDR35):c.2912A>G	397515535	WDR35	[]	[]	['Cranioectodermal dysplasia 2']

(p.Tyr971Cys)					
NM_000424.3(KRT5): c.1424A>G (p.Glu475Gly)	61348633	KRT5	[]	[]	['Epidermolysis bullosa herpetiformis, Dowling-Meara', 'not provided']
NM_004595.4(SMS):c .443A>G (p.Gln148Arg)	397515551	SMS	[]	[]	['Snyder Robinson syndrome']
NM_001256850.1(TTN):c.45629-2A>G	794729266	-	[]	[]	['not provided']
NM_000404.2(GLB1): c.947A>G (p.Tyr316Cys)	72555361	GLB1	[]	[]	['Infantile GM1 gangliosidosis']
NM_000404.2(GLB1): c.1498A>G (p.Thr500Ala)	72555368	GLB1	[]	[]	['Mucopolysaccharidosis, MPS-IV-B']
NM_000404.2(GLB1): c.1772A>G (p.Tyr591Cys)	72555371	GLB1	[]	[]	['GM1-GANGLIOSIDOSIS, TYPE I, WITH CARDIAC INVOLVEMENT']
NM_000487.5(ARSA) :c.1055A>G (p.Asn352Ser)	2071421	ARSA	[]	[]	['Metachromatic leukodystrophy', 'not provided']
NM_001037811.2(HSD17B10):c.713A>G (p.Asn238Ser)	122461163	HSD17B10	[]	[]	['2-methyl-3-hydroxybutyric aciduria']
NM_000138.4(FBN1): c.1148-2A>G	397515756	FBN1	[]	[]	['Marfan syndrome']
NM_001875.4(CPS1): c.1010A>G (p.His337Arg)	28940283	CPS1	[]	[]	['Congenital hyperammonemia, type I']
NM_000169.2(GLA):c .1153A>G (p.Thr385Ala)	397515869	-	['AGCTGTG YGATGAAG CAGGCAGG']	['AGCTGTGYGATG AAGCAGGCAGG']	['not specified', 'not provided']
NM_000256.3(MYBPC3):c.1224-2A>G	397515891	MYBPC3	[]	['TACTTGTCYGTAG AACAGAAGGGG']	['Familial hypertrophic cardiomyopathy 4', 'Cardiomyopathy']
NM_000048.3(ASL):c. 857A>G (p.Gln286Arg)	28941472	ASL	[]	[]	['Argininosuccinate lyase deficiency', 'not provided']
NM_000256.3(MYBPC3):c.1928-2A>G	397515937	MYBPC3	[]	[]	['Primary familial hypertrophic cardiomyopathy', 'Familial hypertrophic cardiomyopathy 4', 'Cardiomyopathy']
NM_002693.2(POLG): c.1283T>C (p.Leu428Pro)	774610098	POLG	[]	[]	['not provided']
NM_004628.4(XPC):c .413-24A>G	794729657	XPC	[]	[]	['Xeroderma pigmentosum, group C']
NM_030973.3(MED25)):c.116A>G	794729668	MED25	[]	[]	['BASEL-VANAGAITE-

(p.Tyr39Cys)					SMIRIN-YOSEF SYNDROME']
NM_001955.4(EDN1): c.271A>G (p.Lys91Glu)	587777231	EDN1	[]	[]	['Auriculocondylar syndrome 3']
NM_003002.3(SDHD) :c.149A>G (p.His50Arg)	11214077	SDHD	[]	[]	['Pheochromocytoma', 'Merkel cell carcinoma', 'Hereditary cancer-predisposing syndrome', 'Carcinoid tumor of intestine', 'Cowden syndrome 3', 'not specified', 'not provided']
NM_001003722.1(GLE1): c.433-10A>G	386833693	GLE1	[]	[]	['Lethal arthrogyrosis with anterior horn cell disease']
NM_000371.3(TTR): c.185A>G (p.Glu62Gly)	11541796	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis']
NM_000256.3(MYBPC3): c.927-2A>G	397516082	MYBPC3	[]	['GTCCCYGTGTCCCGCAGTCTAGG']	['Familial hypertrophic cardiomyopathy 4', 'Cardiomyopathy']
NM_001148.4(ANK2) :c.4373A>G (p.Glu1458Gly)	72544141	ANK2	[]	[]	['Long QT syndrome', 'Congenital long QT syndrome', 'Long QT syndrome 4', 'Cardiac arrhythmia, ankyrin B-related', 'Cardiac arrhythmia']
NM_000257.3(MYH7) :c.2206A>G (p.Ile736Val)	397516138	MYH7	[]	['TATCAAYGAACTGTCCCTCAGGG', 'CTATCAAYGAAC TGTCCCTCAGG']	['Familial hypertrophic cardiomyopathy 1', 'Cardiomyopathy', 'not specified']
NM_000356.3(TCOF1) :c.149A>G (p.Tyr50Cys)	28941769	TCOF1	['GTGTGTA YAGATGTC CAGAAGG G']	['GTGTGTAYAGAT GTCCAGAAGGG']	['Treacher collins syndrome 1']
NM_002150.2(HPD): c.97G>A (p.Ala33Thr)	1154510	HPD	[]	['ATGACGYGGCCTGAATCACAGGG', 'AATGACGYGGCC TGAATCACAGG']	['4-Alpha-hydroxyphenylpyruvate hydroxylase deficiency']
NM_000375.2(UROS) :c.184A>G (p.Thr62Ala)	28941775	UROS	[]	[]	['Congenital erythropoietic porphyria']
NM_001008216.1(GALE) :c.308A>G (p.Asp103Gly)	28940883	GALE	[]	[]	['UDPglucose-4-epimerase deficiency']
NM_000260.3(MYO7A): c.6439-2A>G	397516330	MYO7A	[]	['ATATCCYGGGG GAGCAGAAAGGG', 'GATATCCYGGGG GAGCAGAAAGG']	['Usher syndrome, type 1']

NM_033071.3(SYNE1):c.15705-12A>G	606231134	SYNE1	[]	[]	['Spinocerebellar ataxia, autosomal recessive 8']
NM_000187.3(HGD):c.1112A>G (p.His371Arg)	120074172	HGD	[]	[]	['Alkaptonuria']
NM_000053.3(ATP7B):c.3443T>C (p.Ile1148Thr)	60431989	ATP7B	['TGCTGAY TGGAAACC GTGAGTGG']	['TGCTGAYTGGAA ACCGTGAGTGG']	['Wilson disease']
NM_000441.1(SLC26A4):c.-3-2A>G	397516411	-	[]	[]	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_003041.3(SLC5A2):c.1961A>G (p.Asn654Ser)	61742739	-	[]	[]	['Familial renal glucosuria']
NM_000551.3(VHL):c.467A>G (p.Tyr156Cys)	397516441	VHL	[]	[]	['Von Hippel-Lindau syndrome']
NM_000531.5(OTC):c.481A>G (p.Asn161Asp)	72556270	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.482A>G (p.Asn161Ser)	72556271	OTC	[]	['CAGCCCAYTGAT AATTGGGATGG']	['not provided']
NM_000531.5(OTC):c.541-2A>G	72556289	OTC	['TCCYAAA AGGCACGG GATGAAGG']	['TCCYAAAAGGC ACGGGATGAAGG']	['not provided']
NM_000531.5(OTC):c.542A>G (p.Glu181Gly)	72556290	OTC	['ATAGTGT YCCTAAAA GGCACGGG']	['ATAGTGTTCCTA AAAGGCACGGG']	['not provided']
NM_000527.4(LDLR):c.2483A>G (p.Tyr828Cys)	28942085	LDLR	[]	[]	['Familial hypercholesterolemia', 'not provided']
NM_000271.4(NPC1):c.3467A>G (p.Asn1156Ser)	28942105	NPC1	[]	[]	['Niemann-Pick disease type C1']
NM_000271.4(NPC1):c.3263A>G (p.Tyr1088Cys)	28942106	NPC1	[]	[]	['NIEMANN-PICK DISEASE, TYPE C1, JUVENILE FORM']
NM_020247.4(ADCK3):c.1541A>G (p.Tyr514Cys)	119468008	ADCK3	[]	[]	['Coenzyme Q10 deficiency, primary, 4']
NM_001063.3(TF):c.956A>G (p.His319Arg)	41295774	TF	[]	[]	[]
NM_172201.1(KCNE2):c.281A>G (p.Glu94Gly)	74424227	KCNE2	[]	[]	['Congenital long QT syndrome']
NM_002294.2(LAMP2):c.65-2A>G	397516743	LAMP2	[]	[]	['Danon disease']
NM_002880.3(RAF1):c.524A>G (p.His175Arg)	397516822	RAF1	[]	[]	['Noonan syndrome 5']
NM_033360.3(KRAS):c.13A>G (p.Lys5Glu)	193929331	KRAS	[]	[]	['Noonan syndrome 3', 'Rasopathy']
NM_000525.3(KCNJ1)	193929337	KCNJ11	[]	[]	['Permanent neonatal']

1):c.155A>G (p.Gln52Arg)					diabetes mellitus']
NM_000525.3(KCNJ11):c.544A>G (p.Ile182Val)	193929348	KCNJ11	['AGAYGAGGGTCTCAGCCCTGCGG']	['AGAYGAGGGTCTCAGCCCTGCGG']	['Permanent neonatal diabetes mellitus']
NM_000525.3(KCNJ11):c.989A>G (p.Tyr330Cys)	193929356	KCNJ11	[]	[]	['Permanent neonatal diabetes mellitus', 'Neonatal insulin-dependent diabetes mellitus']
NM_001288953.1(TTC7A):c.1715A>G (p.Lys572Arg)	139010200	TTC7A	[]	[]	['Multiple gastrointestinal atresias']
NM_024809.4(TCTN2):c.1506-2A>G	374349989	TCTN2	[]	[]	['Meckel syndrome type 8']
NM_012275.2(IL36RN):c.104A>G (p.Lys35Arg)	187015338	IL36RN	[]	[]	['Pustular psoriasis, generalized']
NM_178517.3(PIGW):c.499A>G (p.Met167Val)	200024253	PIGW	[]	[]	['Hyperphosphatasia with mental retardation syndrome 5']
NM_015662.2(IFT172):c.5179T>C (p.Cys1727Arg)	149614625	-	[]	[]	['Short-rib thoracic dysplasia 10 with or without polydactyly']
NM_000226.3(KRT9):c.469A>G (p.Met157Val)	58597584	KRT9	[]	[]	['Epidermolytic palmoplantar keratoderma', 'not provided']
NM_023073.3(C5orf42):c.3290-2A>G	606231260	C5orf42	[]	['ATCYATCAAATCAAAAATTTGG']	['Orofaciodigital syndrome 6']
NM_005633.3(SOS1):c.508A>G (p.Lys170Glu)	397517172	SOS1	[]	[]	['Noonan syndrome 4', 'Rasopathy', 'not provided']
NM_006306.3(SMC1A):c.3254A>G (p.Tyr1085Cys)	587784418	SMC1A	['CTTAYAGATCTCATCAATGTTGG']	['CTTAYAGATCTCATCAATGTTGG']	['Congenital muscular hypertrophy-cerebral syndrome']
NM_006218.2(PIK3CA):c.1637A>G (p.Gln546Arg)	397517201	PIK3CA	[]	[]	['Neoplasm of ovary']
NM_006218.2(PIK3CA):c.3073A>G (p.Thr1025Ala)	397517202	PIK3CA	[]	[]	['Non-small cell lung cancer']
NM_002354.2(EPCAM):c.492-2A>G	606231281	EPCAM	[]	[]	['Diarrhea 5, with tufting enteropathy, congenital']
NM_033056.3(PCDH15):c.1998-2A>G	397517452	PCDH15	[]	[]	['Usher syndrome, type 1F']
NM_000301.3(PLG):c.112A>G (p.Lys38Glu)	73015965	PLG	[]	[]	['Plasminogen deficiency, type 1']
NM_000155.3(GALT):c.424A>G (p.Met142Val)	111033692	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000096.3(CP):c.2953A>G	386134132	CP	[]	[]	['Deficiency of ferroxidase']

(p.Met985Val)					
NM_000197.1(HSD17B3):c.703A>G (p.Met235Val)	119481074	HSD17B3	[]	[]	['Testosterone 17-beta-dehydrogenase deficiency']
NM_000197.1(HSD17B3):c.389A>G (p.Asn130Ser)	119481079	HSD17B3	[]	[]	['Testosterone 17-beta-dehydrogenase deficiency']
NM_015474.3(SAMHD1):c.1106T>C (p.Leu369Ser)	515726139	SAMHD1	[]	[]	['Aicardi Goutieres syndrome 5']
NM_004817.3(TJP2):c.1992-2A>G	587777521	TJP2	[]	['CAGCTCYGAGAAGAAACCACGGG', 'TCAGCTCYGAGAAGAAACCACGGG']	['Progressive familial intrahepatic cholestasis 4']
NM_000257.3(MYH7):c.617A>G (p.Lys206Arg)	730880846	MYH7	[]	['CTTCYTGCTGCGGTCCCAATGG']	['Cardiomyopathy']
NM_020919.3(ALS2):c.2980-2A>G	386134184	ALS2	[]	[]	['Juvenile primary lateral sclerosis']
m.10044A>G	41362547	MT-TG	[]	[]	['Sudden death']
NM_002977.3(SCN9A):c.406A>G (p.Ile136Val)	80356468	SCN9A	[]	[]	['Primary erythromelalgia']
NM_001128425.1(MUTYH):c.536A>G (p.Tyr179Cys)	34612342	MUTYH	[]	[]	['MYH-associated polyposis', 'Hereditary cancer-predisposing syndrome', 'Endometrial carcinoma', 'Carcinoma of colon', 'not specified', 'not provided']
NM_206933.2(USH2A):c.12067-2A>G	397517978	USH2A	[]	['TTCCCYGTAAGA AAATTAACAGG']	['Usher syndrome, type 2A', 'Retinitis pigmentosa 39']
NM_000216.2(ANOS1):c.1A>G (p.Met1Val)	606231409	ANOS1	[]	['GCACCAYGGCTGCGGGTCGAGGG', 'GGCACCAYGGCTGCGGGTCGAGG']	['Kallmann syndrome 1']
NM_206933.2(USH2A):c.1841-2A>G	397518003	USH2A	[]	[]	['Usher syndrome, type 2A']
NM_000368.4(TSC1):c.1760A>G (p.Lys587Arg)	118203576	TSC1	[]	[]	['Tuberous sclerosis syndrome', 'Tuberous sclerosis 1', 'Hereditary cancer-predisposing syndrome', 'not specified']
NM_021830.4(C10orf2):c.1523A>G (p.Tyr508Cys)	80356540	C10orf2	[]	[]	['Mitochondrial DNA depletion syndrome 7 (hepatocerebral type)', 'not provided']
NM_003334.3(UBA1):	80356546	UBA1	[]	['TGGCYTGTCACC	['Arthrogryposis

c.1639A>G (p.Ser547Gly)				CGGATATGTGG']	multiplex congenita, distal, X-linked']
NM_206933.2(USH2A):c.8559-2A>G	397518039	USH2A	['ATCYAAA GCAAAAAG ACAAGCAG G']	['ATCYAAAAGCAA AAGACAAGCAGG']	['Retinitis pigmentosa', 'Usher syndrome, type 2A']
NM_000038.5(APC):c.1744-2A>G	587783035	APC	['TCCYAGT AAGAAAC AGAATATG G']	['TCCYAGTAAGA AACAGAATATGG']	['Familial adenomatous polyposis 1']
NM_194248.2(OTOF):c.766-2A>G	80356584	OTOF	[]	['GACCYGCAGGC AGGAGAAGGGGG , 'TGACCYGCAGGC AGGAGAAGGGG', 'CTGACCYGCAGG CAGGAGAAGGG', 'GCTGACCYGCAG GCAGGAGAAGG']	['Deafness, autosomal recessive 9']
NM_000525.3(KCNJ11):c.509A>G (p.Lys170Arg)	80356621	KCNJ11	[]	[]	['Permanent neonatal diabetes mellitus']
NM_000352.4(ABCC8):c.215A>G (p.Asn72Ser)	80356634	ABCC8	[]	[]	['Permanent neonatal diabetes mellitus']
NM_000352.4(ABCC8):c.4270A>G (p.Ile1424Val)	80356653	ABCC8	[]	[]	['Permanent neonatal diabetes mellitus']
NM_000207.2(INS):c.323A>G (p.Tyr108Cys)	80356672	-	[]	[]	['Permanent neonatal diabetes mellitus']
NM_000083.2(CLCN1):c.382A>G (p.Met128Val)	80356699	CLCN1	[]	[]	['Myotonia congenita', 'Congenital myotonia, autosomal dominant form']
NM_001008211.1(OPTN):c.1433A>G (p.Glu478Gly)	267606929	OPTN	[]	[]	['Amyotrophic lateral sclerosis type 12']
NM_007375.3(TARDBP):c.506A>G (p.Asp169Gly)	80356717	TARDBP	[]	[]	['Amyotrophic lateral sclerosis type 10']
NM_007375.3(TARDBP):c.1009A>G (p.Met337Val)	80356730	TARDBP	[]	[]	['Amyotrophic lateral sclerosis type 10']
NM_007375.3(TARDBP):c.1028A>G (p.Gln343Arg)	80356731	TARDBP	[]	[]	['Amyotrophic lateral sclerosis type 10']
NM_001701.3(BAAT):c.967A>G (p.Ile323Val)	80356747	BAAT	['CAAYGAA GAGGAATT GCCCTGG']	['CAAYGAAGAGG AATTGCCCTGG']	['Atypical hemolytic-uremic syndrome 1']
NM_012463.3(ATP6V0A2):c.732-2A>G	80356753	ATP6V0A2	[]	[]	['Cutis laxa with osteodystrophy']
NM_001876.3(CPT1A):c.1361A>G (p.Asp454Gly)	80356778	CPT1A	[]	[]	['Carnitine palmitoyltransferase I deficiency']
NM_001876.3(CPT1A):c.1079A>G	80356787	CPT1A	[]	[]	['Carnitine palmitoyltransferase

(p.Glu360Gly)					I deficiency']
NM_001876.3(CPT1A):c.1493A>G (p.Tyr498Cys)	80356791	CPT1A	[]	[]	['Carnitine palmitoyltransferase I deficiency']
NM_003159.2(CDKL5):c.211A>G (p.Asn71Asp)	587783072	CDKL5	[]	[]	['Atypical Rett syndrome', 'not provided']
NM_000257.3(MYH7):c.1615A>G (p.Met539Val)	730880930	MYH7	[]	['GGAACAYGCAC TCCTCTCCAGG']	['Cardiomyopathy']
m.5843A>G	118203894	MT-TY	[]	[]	[]
NM_000130.4(F5):c.1000A>G (p.Arg334Gly)	118203905	F5	[]	[]	[]
NM_000130.4(F5):c.5189A>G (p.Tyr1730Cys)	118203907	F5	['GTAGYAG GCCCAAGC CCGACAGG']	['GTAGYAGGCCCC AAGCCCGACAGG']	['Factor V deficiency']
NM_000052.6(ATP7A):c.3911A>G (p.Asn1304Ser)	151340632	ATP7A	[]	[]	['Menkes kinky-hair syndrome', 'Cutis laxa, X-linked']
NM_007294.3(BRCA1):c.5053A>G (p.Thr1685Ala)	80356890	BRCA1	[]	[]	['Familial cancer of breast', 'Breast-ovarian cancer, familial 1', 'Hereditary cancer-predisposing syndrome']
NM_000046.3(ARSB):c.629A>G (p.Tyr210Cys)	118203943	ARSB	[]	[]	['Mucopolysaccharidosis type VI', 'not provided']
NM_013319.2(UBIAD1):c.305A>G (p.Asn102Ser)	118203945	UBIAD1	['GTAAGTG YTGACCAA ATTACCGG']	['GTAAGTG YTGACCAA ATTACCGG']	['Schnyder crystalline corneal dystrophy']
NM_013319.2(UBIAD1):c.355A>G (p.Arg119Gly)	118203947	UBIAD1	[]	['TCCYGT CATCAC TCTTTTGTGG']	['Schnyder crystalline corneal dystrophy']
NM_013319.2(UBIAD1):c.695A>G (p.Asn232Ser)	118203949	UBIAD1	['GGTGTG YTGGAATG GAGAATGG']	['GGTGTG YTGGAATG ATGGAGAATGG']	['Schnyder crystalline corneal dystrophy']
NM_013319.2(UBIAD1):c.335A>G (p.Asp112Gly)	118203950	UBIAD1	[]	[]	['Schnyder crystalline corneal dystrophy']
NM_024334.2(TMEM43):c.271A>G (p.Ile91Val)	144811578	TMEM43	[]	[]	['Emery-Dreifuss muscular dystrophy 7, autosomal dominant', 'not provided']
NM_012073.4(CCT5):c.440A>G (p.His147Arg)	118203986	CCT5	[]	[]	['Neuropathy, hereditary sensory, with spastic paraplegia, autosomal recessive']
NM_000033.3(ABCD1):c.443A>G (p.Asn148Ser)	128624216	ABCD1	['CACTGYT GACGAAG GTAGCAGG']	['CACTGYT GACGAAG GTAGCAGG', 'GCACTGYT GACG']	['Adrenoleukodystrophy']

			G']	AAGGTAGCAGG']	
NM_000146.3(FTL):c.1A>G (p.Met1Val)	139732572	FTL	['CTCAYGG TTGGTTGG CAAGAAG G']	['CTCAYGGTTGGT TGGCAAGAAGG']	['L-ferritin deficiency']
NM_000785.3(CYP27B1):c.566A>G (p.Glu189Gly)	118204012	CYP27B1	[]	[]	['Vitamin D-dependent rickets, type 1']
NM_000252.2(MTM1):c.1261-10A>G	397518445	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_139281.2(WDR36):c.1064A>G (p.Asn355Ser)	118204022	WDR36	[]	[]	['Glaucoma 1, open angle, G']
NM_000392.4(ABCC2):c.4145A>G (p.Gln1382Arg)	72558202	ABCC2	[]	[]	['Dubin-Johnson syndrome']
NM_000165.4(GJA1):c.617A>G (p.Lys206Arg)	397518464	GJA1	[]	[]	['Oculodentodigital dysplasia']
NM_000833.4(GRIN2A):c.1123-2A>G	397518469	GRIN2A	[]	[]	['Focal epilepsy with speech disorder with or without mental retardation']
NM_015702.2(MMADHC):c.746A>G (p.Tyr249Cys)	118204046	MMADHC	[]	[]	['Homocystinuria, cblD type, variant 1']
NM_000526.4(KRT14):c.368A>G (p.Asn123Ser)	60171927	KRT14	[]	['GCGGTCAYTGA GGTCTGCATGG']	['Epidermolysis bullosa herpetiformis, Dowling-Meara', 'not provided']
NM_000237.2(LPL):c.548A>G (p.Asp183Gly)	118204064	LPL	['AGCTGGA YCGAGGCC TAAAAGG']	['GCTGGAYCGAG GCCTTAAAAGG', 'AGCTGGAYCGAG GCCTTAAAAGG']	['Hyperlipoproteinaemia, type I']
NM_016247.3(IMPG2):c.370T>C (p.Phe124Leu)	201893545	IMPG2	['ACTYTTT GGGATCGA CTTCCTGG']	['ACTYTTTGGGAT CGACTTCCTGG']	['Macular dystrophy, vitelliform, 5']
NM_004035.6(ACOX1):c.832A>G (p.Met278Val)	118204090	ACOX1	[]	[]	['Pseudoneonatal adrenoleukodystrophy']
NM_004035.6(ACOX1):c.926A>G (p.Gln309Arg)	118204092	ACOX1	[]	[]	['Pseudoneonatal adrenoleukodystrophy']
NM_000190.3(HMBS):c.1A>G (p.Met1Val)	118204118	HMBS	[]	[]	['Porphyria, acute intermittent, nonerythroid variant']
NM_001363.4(DKC1):c.941A>G (p.Lys314Arg)	199422248	DKC1	[]	['AATCYTGGCCCC ATAGCAGATGG']	['Dyskeratosis congenita X-linked']
NM_000078.2(CETP):c.1376A>G (p.Asp459Gly)	2303790	CETP	[]	[]	['Hyperalphalipoproteinemia']
NM_000531.5(OTC):c.595A>G (p.Asn199Asp)	72558405	OTC	[]	[]	['not provided']

NM_000531.5(OTC):c.596A>G (p.Asn199Ser)	72558406	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.613A>G (p.Met205Val)	72558411	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.717+3A>G	72558432	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.718-2A>G	72558433	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.788A>G (p.Asp263Gly)	72558443	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.790A>G (p.Thr264Ala)	72558444	OTC	[]	[]	['not provided']
NM_000531.5(OTC):c.929A>G (p.Glu310Gly)	72558467	OTC	[]	['TCCACTYCTTCTGGCTTTCTGGG', 'ATCCACTYCTTCTGGCTTTCTGG']	['not provided']
NM_000531.5(OTC):c.988A>G (p.Arg330Gly)	72558478	OTC	[]	['ACTTTCYGT TTTCTGCCTCTGGG', 'CACTTTCYGT TTTCTGCCTCTGG']	['not provided']
NM_000488.3(SERPINC1):c.655A>G (p.Asn219Asp)	121909571	SERPINC1	[]	[]	['Antithrombin III deficiency']
NM_007294.3(BRCA1):c.122A>G (p.His41Arg)	80357276	BRCA1	['AAATATGYGGTCACTTTGTGG']	['AAATATGYGGTCACTTTGTGG']	['Familial cancer of breast', 'Breast-ovarian cancer, familial 1']
NM_007294.3(BRCA1):c.1A>G (p.Met1Val)	80357287	BRCA1	[]	[]	['Familial cancer of breast', 'Breast-ovarian cancer, familial 1', 'Hereditary cancer-predisposing syndrome']
NM_198056.2(SCN5A):c.1134T>A (p.Tyr378Ter)	373172185	SCN5A	[]	[]	['not provided']
m.7526A>G	121434454	MT-TD	[]	[]	[]
NM_007294.3(BRCA1):c.211A>G (p.Arg71Gly)	80357382	BRCA1	[]	[]	['Familial cancer of breast', 'Hereditary breast and ovarian cancer syndrome', 'Breast-ovarian cancer, familial 1', 'Hereditary cancer-predisposing syndrome']
NM_000512.4(GALNS):c.1460A>G (p.Asn487Ser)	118204440	GALNS	['ACGYTGAGCTGGGCTGCGCGG', 'CACGYTGAGCTGGGCTGCGCGG']	['ACGYTGAGCTGGGCTGCGCGG', 'CACGYTGAGCTGGGCTGCGCGG']	['Mucopolysaccharidosis, MPS-IV-A']

NM_000505.3(F12):c.158A>G (p.Tyr53Cys)	118204455	F12	[]	['GGTGGYACTGG AAGGGGAAGTGG']	[]
NM_007294.3(BRCA1):c.5453A>G (p.Asp1818Gly)	80357477	BRCA1	[]	['TTGYCCTCTGTC CAGGCATCTGG']	['Familial cancer of breast', 'Breast-ovarian cancer, familial 1']
NM_032492.3(JAGN1):c.485A>G (p.Gln162Arg)	587777730	JAGN1	[]	[]	['Severe congenital neutropenia', 'Severe congenital neutropenia 6, autosomal recessive']
NM_000257.3(MYH7):c.2087A>G (p.Asn696Ser)	730880732	MYH7	[]	[]	['Cardiomyopathy']
NM_000430.3(PAFAH1B1):c.446A>G (p.His149Arg)	121434482	PAFAH1B1	[]	[]	['Lissencephaly 1']
NM_000363.4(TNNI3):c.547A>G (p.Lys183Glu)	730881077	TNNI3	[]	[]	['Cardiomyopathy']
NM_018105.2(THAP1):c.266A>G (p.Lys89Arg)	267607111	THAP1	[]	[]	['Dystonia 6, torsion']
NM_016599.4(MYOZ2):c.738A>G (p.Ile246Met)	140126678	MYOZ2	[]	[]	['Familial hypertrophic cardiomyopathy 16', 'not specified', 'not provided']
NM_000161.2(GCH1):c.671A>G (p.Lys224Arg)	41298442	GCH1	[]	[]	['Dystonia 5, Dopa-responsive type', 'Dystonia, dopa-responsive, with or without hyperphenylalaninemia, autosomal recessive']
NM_017415.2(KLHL3):c.926A>G (p.Gln309Arg)	199469627	KLHL3			['Pseudohypoadosteronism, type 2']
NM_017415.2(KLHL3):c.1670A>G (p.Tyr557Cys)	199469645	KLHL3			['Pseudohypoadosteronism, type 2', 'Pseudohypoadosteronism type 2D']
NM_003590.4(CUL3):c.1207-26A>G	199469650	CUL3			['Pseudohypoadosteronism, type 2']
NM_003590.4(CUL3):c.1238A>G (p.Asp413Gly)	199469656	CUL3			['Pseudohypoadosteronism, type 2', 'Pseudohypoadosteronism type 2E']
NM_003590.4(CUL3):c.1376A>G (p.Lys459Arg)	199469658	CUL3			['Pseudohypoadosteronism, type 2']
NM_003590.4(CUL3):c.1377+3A>G	199469661	CUL3			['Pseudohypoadosteronism, type 2']
NM_007294.3(BRCA1):c.4096+3A>G	80358015	BRCA1	[]	[]	['Hereditary breast and ovarian cancer syndrome', 'Breast-

					ovarian cancer, familial 1', 'Hereditary cancer-predisposing syndrome']
NM_007294.3(BRCA1):c.135-2A>G	80358065	BRCA1	[]	[]	['Breast-ovarian cancer, familial 1']
NM_024426.4(WT1):c.1391A>G (p.Asp464Gly)	121907902	WT1	[]	[]	['Drash syndrome']
NM_007294.3(BRCA1):c.212+3A>G	80358083	BRCA1	[]	[]	['Familial cancer of breast', 'Breast-ovarian cancer, familial 1']
NM_024426.4(WT1):c.1021A>G (p.Ser341Gly)	121907908	WT1	[]	['CGCYCTCGTACCCTGTGCTGTGG']	['Mesothelioma']
NM_007294.3(BRCA1):c.4676-2A>G	80358096	BRCA1	[]	[]	['Breast-ovarian cancer, familial 1', 'Hereditary cancer-predisposing syndrome']
NM_000280.4(PAX6):c.1171A>G (p.Thr391Ala)	121907926	PAX6	[]	['GTGGYGCCCGA GGTGCCCATTTGG']	['Optic nerve aplasia, bilateral']
NM_000520.4(HEXA):c.611A>G (p.His204Arg)	121907976	HEXA	[]	[]	['Tay-Sachs disease']
NM_000159.3(GCDH):c.1213A>G (p.Met405Val)	141437721	GCDH	[]	[]	['Glutaric aciduria, type 1']
NM_024740.2(ALG9):c.860A>G (p.Tyr287Cys)	121908023	ALG9	[]	['TTAYACAAAAC AATGTTGAGTGG']	['Congenital disorder of glycosylation type 1L']
NM_003051.3(SLC16A1):c.610A>G (p.Lys204Glu)	80358222	SLC16A1	[]	[]	['Erythrocyte lactate transporter defect']
NM_000229.1(LCAT):c.463A>G (p.Asn155Asp)	121908057	LCAT	[]	[]	['Fish-eye disease']
NM_000639.2(FASLG):c.466A>G (p.Arg156Gly)	80358238	FASLG	[]	[]	['Autoimmune lymphoproliferative syndrome']
NM_001369.2(DNAH5):c.1121T>C (p.Ile374Thr)	147499872	DNAH5	[]	[]	['Ciliary dyskinesia, primary, 3']
NM_138691.2(TMC1):c.1960A>G (p.Met654Val)	121908074	TMC1	[]	[]	['Deafness, autosomal recessive 7']
NM_024301.4(FKRP):c.1387A>G (p.Asn463Asp)	121908110	FKRP	[]	[]	['Congenital muscular dystrophy-dystroglycanopathy (with or without mental retardation) type B5', 'Limb-girdle muscular dystrophy-dystroglycanopathy, type C5', 'Muscular

					dystrophy', 'Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies type A5', 'Congenital muscular dystrophy-dystroglycanopathy without mental retardation, type B5', 'not provided']
NM_175073.2(APTX): c.602A>G (p.His201Arg)	121908133	APTX	['GCCAAYG GTAACGGG CCTTTGGG']	['GCCAAYGGTAA CGGGCCTTTGGG', 'AGCCAAYGGTAA CGGGCCTTTGG']	['Adult onset ataxia with oculomotor apraxia']
NM_001243133.1(NLRP3): c.1880A>G (p.Glu627Gly)	121908148	NLRP3	[]	['ACAATYCCAGCT GGCTGGGCTGG']	['Familial cold urticaria']
NM_006492.2(ALX3): c.608A>G (p.Asn203Ser)	121908166	ALX3	[]	['CGGYTCTGGAAC CAGACCTGGGG', 'GCGGYTCTGGAA CCAGACCTGGG', 'TGCGGYTCTGGA ACCAGACCTGG']	['Frontonasal dysplasia 1']
NM_020451.2(SEPN1): c.1A>G (p.Met1Val)	121908184	SEPN1	[]	['CCCAYGGCTGCG GCTGGCGGCGG', 'CGGCCAYGGCT GCGGCTGGCGG']	['Eichsfeld type congenital muscular dystrophy']
NM_001127221.1(CACNA1A): c.4151A>G (p.Tyr1384Cys)	121908219	CACNA1A	[]	[]	['Familial hemiplegic migraine type 1']
NM_005249.4(FOXG1): c.686T>A (p.Ile229Asn)	199502880	FOXG1	[]	[]	['not provided']
NM_130468.3(CHST14): c.878A>G (p.Tyr293Cys)	121908258	CHST14	[]	['AAGTCAYAGTG CACGGCACAAGG']	['Ehlers-Danlos syndrome, musculocontractural type']
NM_013391.3(DMGDH): c.326A>G (p.His109Arg)	121908331	DMGDH	[]	[]	['Dimethylglycine dehydrogenase deficiency']
NM_015166.3(MLC1): c.422A>G (p.Asn141Ser)	121908344	MLC1	[]	[]	['Megalencephalic leukoencephalopathy with subcortical cysts 1']
NM_000441.1(SLC26A4): c.1105A>G (p.Lys369Glu)	121908361	SLC26A4	[]	[]	['Enlarged vestibular aqueduct syndrome']
NM_000441.1(SLC26A4): c.2168A>G (p.His723Arg)	121908362	SLC26A4	[]	[]	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_015560.2(OPA1): c.1745A>G (p.Tyr582Cys)	121908376	OPA1	[]	[]	['Optic Atrophy Type 1']
NM_001128425.1(MUTYH): c.1241A>G (p.Gln414Arg)	121908383	MUTYH	[]	['AAGCYGCTCTGA GGGCTCCCAGG']	['Neoplasm of stomach']
NM_015247.2(CYLD): c.2240A>G	121908389	CYLD	[]	[]	['Familial multiple trichoepitheliomata',

(p.Glu747Gly)					'Spiegler-Brooke syndrome']
NM_021102.3(SPINT2):c.488A>G (p.Tyr163Cys)	121908403	SPINT2	['TCCAYAGATGAAGTTATTGCAGG']	['TCCAYAGATGAAGTTATTGCAGG']	['Diarrhea 3, secretory sodium, congenital, syndromic']
NM_004924.4(ACTN4):c.763A>G (p.Lys255Glu)	121908415	ACTN4	[]	[]	['Focal segmental glomerulosclerosis 1']
NM_004795.3(KL):c.578A>G (p.His193Arg)	121908423	KL	['CAGYGGTACAGGGTACAGGGG', 'CCAGYGGTACAGGGTACAGGGG']	['CAGYGGTACAGGGTACAGGGG', 'CCAGYGGTACAGGGTACAGGGG']	[]
NM_005682.6(ADGRG1):c.263A>G (p.Tyr88Cys)	121908466	ADGRG1	['TGGYAGAGGGCCCTGGGGTCAGG']	['TGGYAGAGGGCCCTGGGGTCAGG']	['Polymicrogyria, bilateral frontoparietal']
NM_139025.4(ADAMTS13):c.1582A>G (p.Arg528Gly)	121908473	ADAMTS13	[]	[]	['Upshaw-Schulman syndrome']
NM_014270.4(SLC7A9):c.695A>G (p.Tyr232Cys)	121908487	SLC7A9	[]	[]	['Cystinuria']
NM_004211.3(SLC6A5):c.1472A>G (p.Tyr491Cys)	121908494	SLC6A5	[]	[]	['Hyperekplexia 3']
NM_004211.3(SLC6A5):c.1526A>G (p.Asn509Ser)	121908497	SLC6A5	[]	[]	['Hyperekplexia 3']
NM_182643.2(DLC1):c.2875A>G (p.Thr959Ala)	121908500	DLC1	[]	[]	['Carcinoma of colon']
NM_014946.3(SPAST):c.1322A>G (p.Asp441Gly)	121908512	SPAST	[]	[]	['Spastic paraplegia 4, autosomal dominant']
NM_014946.3(SPAST):c.1157A>G (p.Asn386Ser)	121908514	SPAST	[]	[]	['Spastic paraplegia 4, autosomal dominant']
NM_000026.2(ADSL):c.736A>G (p.Lys246Glu)	119450944	ADSL	[]	[]	['Adenylosuccinate lyase deficiency']
NM_000334.4(SCN4A):c.3478A>G (p.Ile1160Val)	121908549	SCN4A	['TGAYGGAGGGGATGGCGCCTAGG']	['TGAYGGAGGGGATGGCGCCTAGG']	[]
NM_000334.4(SCN4A):c.421A>G (p.Ile141Val)	121908561	SCN4A	[]	[]	['Paramyotonia congenita of von Eulenburg']
NM_004328.4(BCS1L):c.148A>G (p.Thr50Ala)	121908580	BCS1L	[]	['GTGYGATCATGTAATGGCGCCGG']	['Mitochondrial complex III deficiency']
NM_152384.2(BBS5):c.547A>G (p.Thr183Ala)	121908582	BBS5	[]	[]	['Bardet-Biedl syndrome 5']
NM_016417.2(GLRX5)	121908584	GLRX5	[]	['CCTGACCYTGTC']	['Anemia,

:c.294A>G (p.Gln98=)				GGAGCTCCGGG']	sideroblastic, pyridoxine- refractory, autosomal recessive']
NM_006206.4(PDGFR A):c.1664A>G (p.Tyr555Cys)	121908589	PDGFRA	[]	[]	[]
NM_002755.3(MAP2 K1):c.389A>G (p.Tyr130Cys)	121908595	MAP2K1	['CCAYAGA AGCCCACG ATGTACGG ']	['CCAYAGAAGCC CACGATGTACGG']	['Cardiofaciocutaneo us syndrome 3', 'Rasopathy']
NM_012082.3(ZFPM2 (p.Glu30Gly)	121908601	ZFPM2	[]	[]	['Double outlet right ventricle', 'Tetralogy of Fallot', 'Diaphragmatic hernia 3']
NM_012082.3(ZFPM2 (p.Thr843Ala)	121908604	-	[]	[]	['Diaphragmatic hernia 3']
NM_022817.2(PER2): c.1984A>G (p.Ser662Gly)	121908635	PER2	[]	['GCCACACYCTCT GCCTTGCCCGG']	['Advanced sleep phase syndrome, familial']
NM_030761.4(WNT4) :c.647A>G (p.Glu216Gly)	121908650	WNT4	[]	[]	['Mullerian aplasia and hyperandrogenism']
NM_003839.3(TNFRS F11A):c.508A>G (p.Arg170Gly)	121908655	TNFRSF1 1A	[]	['GGGTCYGCATTT GTCCGTGGAGG']	['Osteopetrosis autosomal recessive 7']
NM_000539.3(RHO):c .886A>G (p.Lys296Glu)	29001653	RHO	[]	['CGCTCTYGGCAA AGAACGCTGGG', 'GCGCTCTYGGCA AAGAACGCTGG']	['Retinitis pigmentosa 4']
NM_004006.2(DMD): c.2317A>G (p.Lys773Glu)	128626244	DMD	[]	[]	['Duchenne muscular dystrophy']
NM_003722.4(TP63):c .697A>G (p.Lys233Glu)	121908838	TP63	['AGCTTYT TTGTAGAC AGGCATGG ']	['AGCTTYTTTGT GACAGGCATGG']	['Split-hand/foot malformation 4']
NM_003722.4(TP63):c .1052A>G (p.Asp351Gly)	121908844	TP63	[]	[]	['Ectrodactyly, ectodermal dysplasia, and cleft lip/palate syndrome 3']
NM_003722.4(TP63):c .1054A>G (p.Arg352Gly)	121908847	TP63	[]	[]	['ADULT syndrome', 'Orofacial cleft 8']
NM_000369.2(TSHR): c.1856A>G (p.Asp619Gly)	121908859	TSHR	[]	[]	['Thyroid adenoma, hyperfunctioning']
NM_000369.2(TSHR): c.548A>G (p.Lys183Arg)	121908879	TSHR	[]	[]	['Hyperthyroidism, familial gestational']
NM_003060.3(SLC22 A5):c.632A>G (p.Tyr211Cys)	121908888	SLC22A5	[]	[]	['Renal carnitine transport defect', 'not provided']
NM_006502.2(POLH): c.1603A>G	56307355	POLH	[]	['AGACTTTYCTGC TTAAAGAAGGG']	['Xeroderma pigmentosum,

(p.Lys535Glu)					variant type']
NM_002977.3(SCN9A):c.1964A>G (p.Lys655Arg)	121908919	-	[]	['CCTTTTCYTGTG TATTTGATTGG']	['Generalized epilepsy with febrile seizures plus, type 7', 'not specified']
NM_002977.3(SCN9A):c.184A>G (p.Ile62Val)	121908920	SCN9A	[]	[]	['Febrile seizures, familial, 3b']
NM_006892.3(DNMT3B):c.2450A>G (p.Asp817Gly)	121908939	DNMT3B	[]	['GACACGYCTGTG TAGTGCACAGG']	['Centromeric instability of chromosomes 1,9 and 16 and immunodeficiency']
NM_001130978.1(DYSF):c.3892A>G (p.Ile1298Val)	121908954	DYSF	[]	[]	['Miyoshi muscular dystrophy 1', 'Limb-girdle muscular dystrophy, type 2B', 'not specified']
NM_001130978.1(DYSF):c.5264A>G (p.Glu1755Gly)	121908961	DYSF	[]	[]	['Limb-girdle muscular dystrophy, type 2B']
NM_016203.3(PRKAG2):c.1148A>G (p.His383Arg)	121908988	PRKAG2	[]	[]	['Familial hypertrophic cardiomyopathy 6']
NM_000492.3(CFTR):c.2738A>G (p.Tyr913Cys)	121909008	CFTR	['CACATAA YACGAAC TGGTCTGG']	['CACATAAYACG AACTGGTGTGG']	['Cystic fibrosis']
NM_000492.3(CFTR):c.326A>G (p.Tyr109Cys)	121909031	CFTR	[]	[]	['Cystic fibrosis']
NM_000492.3(CFTR):c.650A>G (p.Glu217Gly)	121909046	CFTR	[]	[]	['Cystic fibrosis']
NM_001040667.2(HSF4):c.256A>G (p.Ile86Val)	121909050	HSF4	[]	[]	['Cataract, zonular']
NM_005025.4(SERPINI1):c.1013A>G (p.His338Arg)	121909052	SERPINI1	[]	[]	['Familial encephalopathy with neuroserpin inclusion bodies']
NM_005422.2(TECTA):c.5609A>G (p.Tyr1870Cys)	121909058	TECTA	[]	[]	['Deafness, autosomal dominant 12']
NM_170695.3(TGIF1):c.838A>G (p.Thr280Ala)	121909068	TGIF1	[]	[]	['Holoprosencephaly 4']
NM_001005360.2(DNM2):c.1684A>G (p.Lys562Glu)	121909088	DNM2	[]	['ACTYCTTCTCTT TCTCCTGAGGG', 'TACTYCTTCTCTT TCTCCTGAGG']	['Charcot-Marie-Tooth disease, dominant intermediate b, with neutropenia']
NM_000483.4(APOC2):c.1A>G (p.Met1Val)	120074112	-	[]	['GCCAYAGTGTC CAGAGACCTGG']	['Apolipoprotein C2 deficiency']
NM_000543.4(SMPD1):c.1154A>G (p.Asn385Ser)	120074123	SMPD1	[]	[]	['Niemann-Pick disease, type B']
NM_000019.3(ACAT1):c.278A>G	120074145	ACAT1	[]	[]	['Deficiency of acetyl-CoA

(p.Asn93Ser)					acetyltransferase']
NM_138477.2(CDAN1):c.1796A>G (p.Asn599Ser)	120074166	CDAN1	[]	[]	['Congenital dyserythropoietic anemia, type I']
NM_000187.3(HGD):c.1102A>G (p.Met368Val)	120074173	HGD	[]	[]	['Alkaptonuria']
NM_001089.2(ABCA3):c.1702A>G (p.Asn568Asp)	121909184	ABCA3	['ACCGTYGTGGCC TGGCCCAG CAGGACGG ']	['ACCGTYGTGGCC CAGCAGGACGG']	['Surfactant metabolism dysfunction, pulmonary, 3']
NM_000503.5(EYA1):c.1639A>G (p.Arg547Gly)	121909197	EYA1	[]	[]	[]
NM_000218.2(KCNQ1):c.418A>G (p.Ser140Gly)	120074192	KCNQ1	['CGCYGAAGATG GATGAGGC AGACCAGG ']	['CGCYGAAGATG AGGCAGACCAGG ']	['Atrial fibrillation, familial, 3', 'Atrial fibrillation']
NM_000314.6(PTEN):c.368A>G (p.His123Arg)	121909222	PTEN	[]	[]	['Cowden syndrome 1']
NM_000314.6(PTEN):c.278A>G (p.His93Arg)	121909238	PTEN	[]	[]	['Hereditary cancer-predisposing syndrome', 'Macrocephaly/autism syndrome']
NM_000314.6(PTEN):c.755A>G (p.Asp252Gly)	121909239	PTEN	[]	['ATAYCACCACAC ACAGGTAACGG']	['Macrocephaly/autism syndrome']
NM_198217.2(ING1):c.515A>G (p.Asn172Ser)	121909251	ING1	[]	['TGGYTGCACAG ACAGTACGTGGG', 'CTGGYTGCACAG ACAGTACGTGG']	['Squamous cell carcinoma of the head and neck']
NM_012338.3(TSPAN12):c.734T>C (p.Leu245Pro)	200519776	TSPAN12	[]	[]	['Exudative vitreoretinopathy 5']
NM_001001557.2(GDF6):c.1271A>G (p.Lys424Arg)	121909353	GDF6	[]	[]	['Klippel-Feil syndrome 1, autosomal dominant']
NM_000163.4(GHR):c.594A>G (p.Glu198=)	121909360	GHR	[]	[]	['Laron-type isolated somatotropin defect']
NM_000256.3(MYBPC3):c.175A>G (p.Thr59Ala)	121909375	MYBPC3	[]	[]	['Familial hypertrophic cardiomyopathy 4']
NM_001174089.1(SLC4A11):c.2518A>G (p.Met840Val)	121909396	SLC4A11	[]	['GATCAYCTTCAT GTAGGGCAGGG', 'AGATCAYCTTCA TGTAGGGCAGG']	['Corneal dystrophy and perceptive deafness']
NM_001100.3(ACTA1):c.350A>G (p.Asn117Ser)	121909520	ACTA1	['GCGGYTGGCCTTGG GCCTTGGG ATTGAGGG , 'CGCGGYTGGCCTTGG GGCCTTGG GATTGAGG ']	['CGGYTGGCCTTGG GGATTGAGGG', 'GCGGYTGGCCTTGG GGATTGAGGG', 'CGCGGYTGGCCTTGG GGATTGAGG']	['Nemaline myopathy 3']
NM_000034.3(ALDO)	121909533	ALDOA	[]	['CCAYCCAACCCT	['HNSHA due to

A):c.386A>G (p.Asp129Gly)				AAGAGAAGAGG']	aldolase A deficiency']
NM_000495.4(COL4A5):c.3455-9A>G	104886388	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_001145.4(ANG):c.208A>G (p.Ile70Val)	121909541	-	[]	[]	['Amyotrophic lateral sclerosis type 9']
NM_000051.3(ATM):c.3118A>G (p.Met1040Val)	3092857	ATM	[]	[]	['Hereditary cancer-predisposing syndrome', 'not specified']
NM_023110.2(FGFR1):c.1121A>G (p.Tyr374Cys)	121909631	FGFR1	[]	[]	['Osteoglophonic dysplasia']
NM_182925.4(FLT4):c.3104A>G (p.His1035Arg)	121909653	FLT4	['CTGYGGA TGC ACTGG GGTGCGGG', 'TCTGYGG ATGCACTG GGGTGC GG']	['CTGYGGATGCAC TGGG GTGCGGG', 'TCTGYGGATGCA CTGGG GTGCGG']	[]
NM_000145.3(FSHR):c.1345A>G (p.Thr449Ala)	121909663	FSHR	[]	[]	['Ovarian hyperstimulation syndrome']
NM_001017420.2(ESCO2):c.1132-7A>G	80359862	ESCO2	[]	[]	['Roberts-SC phocomelia syndrome']
NM_001017420.2(ESCO2):c.1674-2A>G	80359869	ESCO2	[]	[]	['Roberts-SC phocomelia syndrome']
NM_024577.3(SH3TC2):c.505T>C (p.Tyr169His)	80359890	SH3TC2	[]	[]	['Charcot-Marie-Tooth disease, type 4C', 'Charcot-Marie-Tooth disease, type IV', 'Mononeuropathy of the median nerve, mild']
NM_032119.3(ADGRV1):c.18131A>G (p.Tyr6044Cys)	121909763	ADGRV1	[]	[]	['Usher syndrome, type 2C']
NM_001360.2(DHCR7):c.839A>G (p.Tyr280Cys)	121909766	DHCR7	[]	[]	['Smith-Lemli-Opitz syndrome']
NM_000517.4(HBA2):c.1A>G (p.Met1Val)	121909803	HBA2	[]	[]	['Hemoglobin H disease, nondeletional']
NM_004006.2(DMD):c.835A>G (p.Thr279Ala)	128627255	DMD	[]	['TGACCGYGATCT GCAGAGAAGGG', 'CTGACCGYGATC TGCAGAGAAGG']	['Dilated cardiomyopathy 3B']
NM_015896.3(ZMYND10):c.797T>C (p.Leu266Pro)	200913791	ZMYND10	[]	[]	['Kartagener syndrome', 'Ciliary dyskinesia, primary, 22']
NM_001085.4(SERPINA3):c.1240A>G (p.Met414Val)	116929575	SERPINA3	[]	['GCTCAYGAAGA AGATGTTCTGGG', 'TGCTCAYGAAGA	[]

				AGATGTTCTGG']	
NM_058216.2(RAD51C):c.1027-2A>G	587780835	RAD51C	[]	[]	['Fanconi anemia, complementation group O']
NM_006231.3(POLE):c.4444+3A>G	398122515	POLE	[]	[]	['Facial dysmorphism, immunodeficiency, livedo, and short stature']
NM_002769.4(PRSS1):c.161A>G (p.Asn54Ser)	144422014	-	[]	[]	['Hereditary pancreatitis']
NM_001204316.1(PRLR):c.635A>G (p.His212Arg)	398122522	PRLR	[]	[]	['Hyperprolactinemia']
NM_004992.3(MECP2):c.410A>G (p.Glu137Gly)	61748392	MECP2	[]	['CAACYCCACTTTAGAGCGAAAGG']	['Mental retardation, X-linked, syndromic 13']
NM_020366.3(RPGRIP1):c.3749-2A>G	376517859	RPGRIP1	[]	[]	['Cone-rod dystrophy 13']
NM_000552.3(VWF):c.2384A>G (p.Tyr795Cys)	61748478	VWF	['GTCAYAGTTCTGGCACGTTTTGG']	['GTCAYAGTTCTGGCACGTTTTGG']	['von Willebrand disease type 2N', 'not provided']
NM_001040613.2(TMEM70):c.*7-2A>G	183973249	TMEM70	[]	[]	['Nuclearly-encoded mitochondrial complex V (ATP synthase) deficiency 2']
NM_001005741.2(GBA):c.1049A>G (p.His350Arg)	78198234	GBA	[]	[]	['Gaucher disease, perinatal lethal']
NM_000218.2(KCNQ1):c.332A>G (p.Tyr111Cys)	199472678	KCNQ1	[]	[]	['Congenital long QT syndrome', 'Cardiac arrhythmia', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.344A>G (p.Glu115Gly)	199472679	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.440A>G (p.Gln147Arg)	199472689	KCNQ1			['Atrial fibrillation']
NM_000218.2(KCNQ1):c.592A>G (p.Ile198Val)	199472700	KCNQ1	[]	[]	['Congenital long QT syndrome']
NM_001943.3(DSG2):c.880A>G (p.Lys294Glu)	752432726	DSG2	[]	[]	['Cardiomyopathy']
NM_000218.2(KCNQ1):c.820A>G (p.Ile274Val)	199472728	KCNQ1	[]	[]	['Sudden infant death syndrome', 'Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.842A>G (p.Tyr281Cys)	199472732	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.842A>G (p.Tyr281Cys)	199472750	KCNQ1			['Congenital long

1):c.950A>G (p.Asp317Gly)					QT syndrome']
NM_000218.2(KCNQ1):c.964A>G (p.Thr322Ala)	199472754	KCNQ1			['Congenital long QT syndrome', 'Cardiac arrhythmia', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.1138A>G (p.Arg380Gly)	199472770	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.1193A>G (p.Lys398Arg)	199472777	KCNQ1			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000218.2(KCNQ1):c.1640A>G (p.Gln547Arg)	199472798	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.1669A>G (p.Lys557Glu)	199472801	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1):c.1705A>G (p.Lys569Glu)	199472808	KCNQ1			['Congenital long QT syndrome']
NM_001005741.2(GBA):c.667T>C (p.Trp223Arg)	61748906	GBA	[]	['CCCACTYGGCTC AAGACCAATGG']	['Gaucher disease, type 1', 'not provided']
NM_000218.2(KCNQ1):c.1756A>G (p.Asn586Asp)	199472812	KCNQ1			['Congenital long QT syndrome', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1):c.1793A>G (p.Lys598Arg)	199472817	KCNQ1			['Sudden infant death syndrome']
NM_000238.3(KCNH2):c.82A>G (p.Lys28Glu)	199472829	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.128A>G (p.Tyr43Cys)	199472836	KCNH2			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2):c.301A>G (p.Lys101Glu)	199472856	KCNH2	[]	[]	['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2):c.652A>G (p.Met218Val)	199472869	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1424A>G (p.Tyr475Cys)	199472907	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1777A>G (p.Ile593Val)	199472930	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1783A>G (p.Lys595Glu)	199472932	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1790A>G (p.Tyr597Cys)	199472934	KCNH2			['Long QT syndrome', 'Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1826A>G	199472940	KCNH2			['Congenital long QT syndrome']

(p.Asp609Gly)					
NM_000238.3(KCNH2):c.1847A>G (p.Tyr616Cys)	199472946	KCNH2			['Long QT syndrome', 'Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2):c.1885A>G (p.Asn629Asp)	199472956	KCNH2			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2):c.1897A>G (p.Asn633Asp)	199472960	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1903A>G (p.Asn635Asp)	199472963	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.1910A>G (p.Glu637Gly)	199472967	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.2510A>G (p.Asp837Gly)	199473004	KCNH2			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2):c.2591A>G (p.Asp864Gly)	199473008	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2):c.3118A>G (p.Ser1040Gly)	199473024	KCNH2	[]	['CTGTCYCTCCACG TCGCCCCGGG', 'CCTGTCYCTCCAC GTCGCCCCGGG', 'GCCTGTCYCTCCA CGTCGCCCCGG']	['Sudden infant death syndrome']
NM_000238.3(KCNH2):c.3233A>G (p.Tyr1078Cys)	199473029	KCNH2			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.343A>G (p.Ser115Gly)	199473057	SCN5A			['Congenital long QT syndrome']
m.827A>G	28358569	MT-RNR1	[]	[]	['Aminoglycoside-induced deafness', 'Deafness, nonsyndromic sensorineural, mitochondrial']
NM_000335.4(SCN5A):c.688A>G (p.Ile230Val)	199473074	SCN5A	['ATAYAGT TTTTCAGGG CCCGGAGG', 'CTGATAYAGTTT TCAGGGCCCCGG']	['ATAYAGTTTTCA GGGCCCCGGAGG', 'CTGATAYAGTTT TCAGGGCCCCGG']	['Brugada syndrome']
NM_000335.4(SCN5A):c.715A>G (p.Ile239Val)	199473075	SCN5A			['Congenital long QT syndrome']
NM_000252.2(MTM1):c.575A>G (p.Tyr192Cys)	587783838	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_004572.3(PKP2):c.275T>A (p.Leu92Ter)	763639737	PKP2	[]	[]	['not provided']

NM_000335.4(SCN5A):c.1502A>G (p.Asp501Gly)	199473117	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.2249A>G (p.Gln750Arg)	199473152	SCN5A			['Congenital long QT syndrome']
NM_198056.2(SCN5A):c.2527A>G (p.Thr843Ala)	199473165	SCN5A			['Congenital long QT syndrome', 'not provided']
NM_001165963.1(SCN1A):c.1277A>G (p.Tyr426Cys)	796052973	SCN1A	[]	[]	['not provided']
NM_000335.4(SCN5A):c.3755A>G (p.Glu1252Gly)	199473214	SCN5A			['Brugada syndrome']
NM_198056.2(SCN5A):c.4000A>G (p.Ile1334Val)	199473226	SCN5A	[]	[]	['Congenital long QT syndrome', 'not provided']
NM_000335.4(SCN5A):c.4252A>G (p.Lys1418Glu)	199473242	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.4291A>G (p.Arg1431Gly)	199473245	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.4412A>G (p.Asn1471Ser)	199473255	SCN5A			['Congenital long QT syndrome']
NM_198056.2(SCN5A):c.4478A>G (p.Lys1493Arg)	199473260	SCN5A	[]	[]	['Atrial fibrillation', 'Congenital long QT syndrome', 'not provided']
NM_000335.4(SCN5A):c.4489A>G (p.Met1497Val)	199473264	SCN5A			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.4577A>G (p.Lys1526Arg)	199473270	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.5161A>G (p.Asn1721Asp)	199473299	SCN5A			['Brugada syndrome']
NM_198056.2(SCN5A):c.5302A>G (p.Ile1768Val)	199473311	SCN5A			['Congenital long QT syndrome', 'not provided']
NM_000335.4(SCN5A):c.5318A>G (p.Asn1773Ser)	199473313	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.5366A>G (p.Asp1789Gly)	199473317	SCN5A			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.5402A>G (p.Asp1801Gly)	199473318	SCN5A			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.5513A>G (p.Asp1838Gly)	199473321	SCN5A			['Congenital long QT syndrome']
NM_198056.2(SCN5A):c.5726A>G (p.Gln1909Arg)	199473326	SCN5A			['Congenital long QT syndrome', 'not provided']
NM_172201.1(KCNE2)	199473366	KCNE2			['Atrial fibrillation']

):c.269A>G (p.Glu90Gly)					
NM_000891.2(KCNJ2))c.223A>G (p.Thr75Ala)	199473370	KCNJ2			['Congenital long QT syndrome']
NM_000891.2(KCNJ2))c.233A>G (p.Asp78Gly)	199473371	KCNJ2			['Andersen Tawil syndrome', 'Congenital long QT syndrome']
NM_000891.2(KCNJ2))c.574A>G (p.Thr192Ala)	199473382	KCNJ2			['Congenital long QT syndrome']
NM_000218.2(KCNQ1))c.548A>G (p.Lys183Arg)	199473396	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1))c.1061A>G (p.Lys354Arg)	199473404	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1))c.1070A>G (p.Gln357Arg)	199473405	KCNQ1			['Congenital long QT syndrome', 'Cardiac arrhythmia', 'Long QT syndrome, LQT1 subtype']
NM_000218.2(KCNQ1))c.1078A>G (p.Arg360Gly)	199473406	KCNQ1	[]	[]	['Congenital long QT syndrome']
NM_000238.3(KCNH2))c.209A>G (p.His70Arg)	199473419	KCNH2			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2))c.1724A>G (p.Glu575Gly)	199473424	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2))c.1747A>G (p.Ile583Val)	199473427	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2))c.1762A>G (p.Asn588Asp)	199473431	KCNH2			['Congenital long QT syndrome']
NM_000218.2(KCNQ1))c.430A>G (p.Thr144Ala)	199473451	KCNQ1			['Congenital long QT syndrome']
NM_000218.2(KCNQ1))c.931A>G (p.Thr311Ala)	199473469	KCNQ1			['Congenital long QT syndrome']
NM_000238.3(KCNH2))c.286A>G (p.Ile96Val)	199473496	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2))c.1205A>G (p.His402Arg)	199473506	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2))c.1259A>G (p.Tyr420Cys)	199473507	KCNH2			['Congenital long QT syndrome']
NM_000238.3(KCNH2))c.1502A>G (p.Asp501Gly)	199473513	KCNH2			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2))c.1912A>G (p.Lys638Glu)	199473528	KCNH2			['Congenital long QT syndrome']

NM_000238.3(KCNH2):c.2131A>G (p.Ile711Val)	199473532	KCNH2	[]	[]	['Long QT syndrome', 'Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000238.3(KCNH2):c.2266A>G (p.Met756Val)	199473534	KCNH2			['Acquired long QT syndrome']
NM_000238.3(KCNH2):c.3343A>G (p.Met1115Val)	199473546	KCNH2			['Congenital long QT syndrome', 'Cardiac arrhythmia']
NM_000335.4(SCN5A):c.89A>G (p.Glu30Gly)	199473551	SCN5A			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.1217A>G (p.Asn406Ser)	199473568	SCN5A			['Brugada syndrome']
NM_000249.3(MLH1):c.791-2A>G	267607794	MLH1	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms', 'Hereditary cancer-predisposing syndrome']
NM_000335.4(SCN5A):c.2780A>G (p.Asn927Ser)	199473589	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.3164A>G (p.Asp1055Gly)	199473593	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.4223A>G (p.Tyr1408Cys)	199473610	SCN5A			['Brugada syndrome']
NM_198056.2(SCN5A):c.4346A>G (p.Tyr1449Cys)	199473613	SCN5A			['Brugada syndrome', 'not provided']
NM_198056.2(SCN5A):c.4978A>G (p.Ile1660Val)	199473625	SCN5A	['CGAYGTTGAAGAGG GCAGGCAGG']	['CGAYGTTGAAGAGGCCAGGCAGG', 'AGCCCGAYGTTG AAGAGGGCAGG']	['Brugada syndrome', 'not provided']
NM_000335.4(SCN5A):c.5138A>G (p.Asp1713Gly)	199473628	SCN5A			['Brugada syndrome']
NM_000335.4(SCN5A):c.5297A>G (p.Tyr1766Cys)	199473632	SCN5A			['Congenital long QT syndrome']
NM_000335.4(SCN5A):c.5317A>G (p.Asn1773Asp)	199473633	SCN5A			['Congenital long QT syndrome']
NM_000531.5(OTC):c.527A>G (p.Tyr176Cys)	72556283	OTC	['TGAGGYAATCAGCCAGGATCTGG']	['TGAGGYAATCAGCCAGGATCTGG']	['not provided']
NM_001130823.1(DNMT1):c.1532A>G (p.Tyr511Cys)	199473690	DNMT1			['Hereditary sensory neuropathy type IE']
NM_000303.2(PMM2):c.563A>G	80338704	PMM2	[]	[]	['Carbohydrate-deficient']

(p.Asp188Gly)					glycoprotein syndrome type I', 'not provided']
NM_000051.3(ATM): c.8030A>G (p.Tyr2677Cys)	28942103	-	[]	[]	['Ataxia-telangiectasia variant']
NM_175053.3(KRT74): c.821T>C (p.Phe274Ser)	147962513	KRT74	[]	[]	['Ectodermal dysplasia, 'pure' hair-nail type', 'Ectodermal dysplasia 7, hair/nail type']
NM_000059.3(BRCA2): c.426-2A>G	398122779	BRCA2	[]	[]	['Familial cancer of breast', 'Breast-ovarian cancer, familial 2']
NM_133433.3(NIPBL): c.5428-2A>G	587783974	NIPBL	[]	[]	['Cornelia de Lange syndrome 1']
NM_000238.3(KCNH2): c.1129-2A>G	794728365	KCNH2	[]	['GGACCYGCACC CGGGGAAGGCGG']	['Cardiac arrhythmia']
NM_000260.3(MYO7A): c.6029A>G (p.Asp2010Gly)	111033175	MYO7A	[]	[]	['Usher syndrome, type 1']
NM_000531.5(OTC): c.548A>G (p.Tyr183Cys)	72556293	OTC	[]	['AGAGCTAYAGT GTTCTAAAAGG']	['not provided']
NM_000441.1(SLC26A4): c.1151A>G (p.Glu384Gly)	111033244	SLC26A4	[]	['TGAATYCCTAAG GAAGAGACTGG']	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_004004.5(GJB2): c.617A>G (p.Asn206Ser)	111033294	GJB2	[]	[]	['Deafness, autosomal recessive 1A', 'Hearing impairment']
NM_000441.1(SLC26A4): c.919-2A>G	111033313	SLC26A4	[]	[]	['Pendred syndrome', 'Enlarged vestibular aqueduct syndrome']
NM_001363.4(DKC1): c.115A>G (p.Lys39Glu)	121912296	DKC1	[]	[]	['Dyskeratosis congenita X-linked']
NM_001363.4(DKC1): c.196A>G (p.Thr66Ala)	121912297	DKC1	[]	[]	['Dyskeratosis congenita X-linked']
NM_001363.4(DKC1): c.361A>G (p.Ser121Gly)	121912305	DKC1	[]	[]	['Dyskeratosis congenita X-linked', 'Hoyeraal Hreidarsson syndrome']
NM_178151.2(DCX): c.413A>G (p.Tyr138Cys)	587783552	DCX	[]	[]	['Heteropia']
NM_024675.3(PALB2): c.212-2A>G	730881879	PALB2	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000260.3(MYO7A): c.1344-2A>G	111033415	MYO7A	[]	['AGCYGCAGGGG CACAGGGATGGG' ; 'AAGCYGCAGGGG CACAGGGATGG']	['Usher syndrome, type 1']

NM_000454.4(SOD1): c.131A>G (p.His44Arg)	121912435	SOD1	[]	[]	['Amyotrophic lateral sclerosis type 1']
NM_000454.4(SOD1): c.302A>G (p.Glu101Gly)	121912439	SOD1	[]	['AGAATCTYCAAT AGACACATCGG']	['Amyotrophic lateral sclerosis type 1']
NM_000454.4(SOD1): c.140A>G (p.His47Arg)	121912443	SOD1	[]	[]	['Amyotrophic lateral sclerosis type 1']
NM_133433.3(NIPBL) :c.737A>G (p.Asp246Gly)	587784042	NIPBL	[]	[]	['Cornelia de Lange syndrome 1']
NM_000454.4(SOD1): c.242A>G (p.His81Arg)	121912458	SOD1	[]	[]	['Amyotrophic lateral sclerosis type 1']
NM_001754.4(RUNX1): c.328A>G (p.Lys110Glu)	121912498	RUNX1	[]	[]	['Familial platelet disorder with associated myeloid malignancy']
NM_000238.3(KCNH2): c.1408A>G (p.Asn470Asp)	121912505	KCNH2	[]	[]	['Long QT syndrome 2', 'Congenital long QT syndrome']
NM_000233.3(LHCR): c.1733A>G (p.Asp578Gly)	121912518	-	[]	[]	['Gonadotropin-independent familial sexual precocity']
NM_000493.3(COL10A1): c.1790A>G (p.Tyr597Cys)	111033554	-	[]	[]	['Metaphyseal chondrodysplasia, Schmid type']
NM_000233.3(LHCR): c.1691A>G (p.Asp564Gly)	121912540	-	[]	[]	['Gonadotropin-independent familial sexual precocity']
NM_002769.4(PRSS1) :c.68A>G (p.Lys23Arg)	111033567	-	[]	['ATCYTGTCATCA TCATCAAAGGG', 'GATCYTGTCATC ATCATCAAAGG']	['Hereditary pancreatitis']
NM_004999.3(MYO6) :c.737A>G (p.His246Arg)	121912560	MYO6	[]	[]	['Sensorineural deafness with hypertrophic cardiomyopathy']
NM_000901.4(NR3C2) :c.2327A>G (p.Gln776Arg)	121912565	NR3C2	[]	['TCATCYGTTTGC CTGCTAAGCGG']	['Pseudohypoadosteronism type 1 autosomal dominant']
NM_000901.4(NR3C2) :c.2915A>G (p.Glu972Gly)	121912574	NR3C2	[]	['CCGACYCCACCT TGGGCAGCTGG']	['Pseudohypoadosteronism type 1 autosomal dominant']
NM_001173464.1(KIF21A): c.2839A>G (p.Met947Val)	121912589	KIF21A	[]	['ATTCAyatCTGC CTCCATGTTGG']	['Fibrosis of extraocular muscles, congenital, 1']
NM_000155.3(GALT) :c.67A>G (p.Thr23Ala)	111033635	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_001041.3(SI): c.350A>G (p.Gln117Arg)	121912612	SI	[]	[]	['Sucrase-isomaltase deficiency']
NM_000155.3(GALT) :c.1A>G (p.Met1Val)	111033639	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate

					uridylyltransferase']
NM_021625.4(TRPV4):c.998A>G (p.Asp333Gly)	121912634	TRPV4	[]	[]	['Spondylometaphyseal dysplasia, Kozlowski type']
NM_000155.3(GALT):c.253-2A>G	111033661	GALT	[]	['ATTCACCYACCGACAAGGATAGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase', 'not provided']
NM_000155.3(GALT):c.290A>G (p.Asn97Ser)	111033669	GALT	[]	['GAAGTCGYTGTCAAACAGGAAGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.379A>G (p.Lys127Glu)	111033682	GALT	[]	['TGACCTYACTGGGTGGTGACGGG', 'ATGACCTYACTGGGTGGTGACGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000343.3(SLC5A1):c.83A>G (p.Asp28Gly)	121912669	SLC5A1	[]	[]	['Congenital glucose-galactose malabsorption']
NM_005159.4(ACTC1):c.1088A>G (p.Glu363Gly)	121912674	-	[]	[]	['Dilated cardiomyopathy 1R']
NM_005159.4(ACTC1):c.373A>G (p.Met125Val)	121912677	-	[]	[]	['Atrial septal defect 5']
NM_000155.3(GALT):c.565-2A>G	111033731	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_001681.3(ATP2A2):c.2300A>G (p.Asn767Ser)	121912732	ATP2A2	[]	[]	['Darier disease, acral hemorrhagic type']
NM_000155.3(GALT):c.821-2A>G	111033767	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000342.3(SLC4A1):c.2509A>G (p.Thr837Ala)	121912750	SLC4A1	[]	[]	['Spherocytosis type 4']
NM_000155.3(GALT):c.950A>G (p.Gln317Arg)	111033786	GALT	[]	['CAGCYGCCAATGGTCCAGTTGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_001202.3(BMP4):c.278A>G (p.Glu93Gly)	121912765	BMP4	[]	['CCTCCYCCCCAGACTGAAGCCGG']	['Microphthalmia syndromic 6']
NM_000155.3(GALT):c.1001A>G (p.Lys334Arg)	111033809	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.1048A>G (p.Thr350Ala)	111033817	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.1132A>G (p.Ile378Val)	111033819	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']

m.3243A>G	199474657	MT-TL1	[]	[]	['Leigh disease', 'Cyclical vomiting syndrome', 'Juvenile myopathy, encephalopathy, lactic acidosis AND stroke', 'Myoclonus with epilepsy with ragged red fibers', 'Cytochrome-c oxidase deficiency', 'Diabetes-deafness syndrome maternally transmitted', '3-Methylglutaconic aciduria', 'Age-related macular degeneration 2', 'MERRF/MELAS overlap syndrome']
m.3252A>G	199474661	MT-TL1	[]	[]	['Mitochondrial encephalomyopathy']
m.3251A>G	199474662	MT-TL1	[]	[]	[]
NM_000258.2(MYL3):c.517A>G (p.Met173Val)	199474708	MYL3	[]	[]	['Cardiomyopathy', 'not specified', 'not provided']
NM_000094.3(COL7A1):c.425A>G (p.Lys142Arg)	121912856	COL7A1	[]	['CACCYTGGGGA CACCAGGTCGGG', 'TCACCYTGGGGA CACCAGGTCGG']	['Epidermolysis bullosa dystrophica inversa, autosomal recessive']
NM_152263.3(TPM3):c.505A>G (p.Lys169Glu)	199474715	TPM3	[]	['CCAACTYACGA GCCACCTACAGG']	['Congenital myopathy with fiber type disproportion', 'not provided']
NM_152263.3(TPM3):c.733A>G (p.Arg245Gly)	199474718	TPM3	[]	['ATCYCTCAGCAA ACTCAGCACGG']	['Congenital myopathy with fiber type disproportion', 'not provided']
NM_001844.4(COL2A1):c.4172A>G (p.Tyr1391Cys)	121912889	COL2A1	['GCAGTGG YAGGTGAT GTTCTGGG']	['GCAGTGGYAGG TGATGTTCTGGG']	['Spondyloperipheral dysplasia', 'Platyspondylic lethal skeletal dysplasia Torrance type']
NM_001844.4(COL2A1):c.2974A>G (p.Arg992Gly)	121912895	COL2A1	[]	['CCTCYCTACCA CGTTGCCAGG']	['Spondyloepimetaphyseal dysplasia Strudwick type']
NM_001848.2(COL6A1):c.362A>G (p.Lys121Arg)	121912936	COL6A1	[]	[]	['Ullrich congenital muscular dystrophy', 'Bethlem myopathy', 'not provided']
NM_004004.5(GJB2):c.218A>G (p.His73Arg)	121912968	GJB2	[]	[]	['Keratoderma palmoplantar deafness']
NM_000941.2(POR):c.1733A>G (p.Tyr578Cys)	121912975	POR	[]	[]	['Antley-Bixler syndrome with genital anomalies']

					and disordered steroidogenesis']
NM_001943.3(DSG2): c.797A>G (p.Asn266Ser)	121913011	DSG2	[]	[]	['Arrhythmogenic right ventricular cardiomyopathy, type 10']
NM_000129.3(F13A1) :c.851A>G (p.Tyr284Cys)	121913074	F13A1	[]	['ATAGGCAYAGA TATTGTCCCAGG']	['Factor xiii, a subunit, deficiency of']
NM_000043.4(FAS):c. 695A>G (p.Tyr232Cys)	121913079	FAS	[]	[]	['Autoimmune lymphoproliferative syndrome, type 1a']
NM_000043.4(FAS):c. 763A>G (p.Asn255Asp)	121913082	FAS	[]	[]	[]
NM_000043.4(FAS):c. 353A>G (p.Asn118Ser)	121913083	FAS	[]	[]	[]
NM_206933.2(USH2A)):c.14020A>G (p.Arg4674Gly)	80338904	USH2A	[]	[]	['Retinitis pigmentosa', 'Retinitis pigmentosa 39']
NM_000142.4(FGFR3)):c.833A>G (p.Tyr278Cys)	121913115	FGFR3	[]	[]	['Hypochondroplasia']
NM_000183.2(HADH B):c. 788A>G (p.Asp263Gly)	121913131	HADHB	[]	[]	['Mitochondrial trifunctional protein deficiency']
NM_001079817.1(INS R):c. 1459A>G (p.Lys487Glu)	121913136	INSR	[]	[]	['Leprechaunism syndrome']
NM_000208.2(INSR): c.707A>G (p.His236Arg)	121913145	INSR	[]	['GCTGYGGCAAC AGAGGCCTTCGG']	['Leprechaunism syndrome']
NM_000208.2(INSR): c.1466A>G (p.Asn489Ser)	121913147	INSR	[]	[]	['Insulin-resistant diabetes mellitus AND acanthosis nigricans']
NM_000208.2(INSR): c.1372A>G (p.Asn458Asp)	121913160	INSR	[]	[]	['Leprechaunism syndrome']
NM_000016.5(ACADM):c. 797A>G (p.Asp266Gly)	201375579	ACADM	[]	[]	['not provided']
NM_024577.3(SH3TC 2):c. 530-2A>G	80338920	SH3TC2	[]	[]	['Charcot-Marie-Tooth disease, type 4C']
NM_001127500.1(MET):c. 3743A>G (p.Tyr1248Cys)	121913246	MET	[]	[]	['Renal cell carcinoma, papillary, 1']
NM_000517.4(HBA2): c.*92A>G	63750067	HBA2	['ACTTYAT TCAAAGAC CAGGAAG G']	['CTTYATTCAAAG ACCAGGAAGGG', 'ACTTYATTCAA GACCAGGAAGG']	['Hemoglobin H disease, nondeletional']
NM_199440.1(HSPD1):c. 86A>G (p.Asp29Gly)	72466451	HSPD1	[]	[]	['Leukodystrophy, hypomyelinating, 4']
NM_000249.3(MLH1) :c.544A>G	63750211	MLH1	[]	[]	['Hereditary Nonpolyposis

(p.Arg182Gly)					Colorectal Neoplasms']
NM_025137.3(SPG11):c.1457-2A>G	312262726	SPG11	[]	[]	['Spastic paraplegia 11, autosomal recessive']
NM_025137.3(SPG11):c.2608A>G (p.Ile870Val)	312262745	SPG11	[]	['ACTTAYCCTGGG GAGAAGGATGG']	['Spastic paraplegia 11, autosomal recessive']
NM_025137.3(SPG11):c.2833A>G (p.Arg945Gly)	312262748	SPG11	[]	[]	['Spastic paraplegia 11, autosomal recessive']
NM_003867.3(FGF17):c.560A>G (p.Asn187Ser)	398123026	FGF17	['CGTGGYT GGGGAAG GGCAGCTG G']	['CGTGGYTGGGG AAGGGCAGCTGG']	['Hypogonadotropic hypogonadism 20 with or without anosmia']
NM_025137.3(SPG11):c.6477+4A>G	312262780	SPG11	[]	[]	['Spastic paraplegia 11, autosomal recessive']
NM_000141.4(FGFR2):c.1124A>G (p.Tyr375Cys)	121913478	FGFR2	[]	[]	['Cutis Gyrate syndrome of Beare and Stevenson', 'Endometrial carcinoma']
NM_000142.4(FGFR3):c.1118A>G (p.Tyr373Cys)	121913485	FGFR3	[]	[]	['Thanatophoric dysplasia type 1']
NM_003611.2(OFD1):c.290A>G (p.Glu97Gly)	312262820	OFD1	[]	[]	['Oral-facial-digital syndrome']
NM_000222.2(KIT):c.1924A>G (p.Lys642Glu)	121913512	KIT	['GACTTYG AGTTCAGA CATGAGGG']	['GACTTYGAGTTC AGACATGAGGG', 'GGACTTYGAGTT CAGACATGAGG']	[]
NM_003611.2(OFD1):c.382-2A>G	312262829	OFD1	[]	[]	['Oral-facial-digital syndrome']
NM_000391.3(TPP1):c.857A>G (p.Asn286Ser)	119455958	TPP1	[]	[]	['Ceroid lipofuscinosis, neuronal, 2']
NM_005912.2(MC4R):c.508A>G (p.Ile170Val)	121913560	MC4R	[]	[]	['Obesity']
NM_005912.2(MC4R):c.821A>G (p.Asn274Ser)	121913561	MC4R	[]	[]	['Obesity']
NM_005912.2(MC4R):c.289A>G (p.Asn97Asp)	121913565	MC4R	[]	[]	['Obesity']
NM_005912.2(MC4R):c.185A>G (p.Asn62Ser)	121913566	MC4R	[]	[]	['Obesity']
NM_000530.6(MPZ):c.286A>G (p.Lys96Glu)	121913583	MPZ	[]	[]	['Charcot-Marie-Tooth disease type 1B']
NM_000095.2(COMP):c.1760A>G (p.His587Arg)	312262901	COMP	[]	[]	['Pseudoachondroplastic spondyloepiphyseal dysplasia syndrome']
NM_000530.6(MPZ):c.242A>G (p.His81Arg)	121913594	MPZ	['GGCATAG YGGAAGAT']	['GGCATAGYGGA AGATCTATGAGG']	['Charcot-Marie-Tooth disease type

			CTATGAGG 		1B']
NM_000484.3(APP):c.2146A>G (p.Ile716Val)	63750399	APP	[]	[]	['Alzheimer disease, type 1', 'not provided']
NM_000329.2(RPE65):c.1292A>G (p.Tyr431Cys)	62636300	RPE65	[]	[]	['Leber congenital amaurosis 2', 'not provided']
NM_002470.3(MYH3):c.1385A>G (p.Asp462Gly)	121913622	MYH3	[]	[]	['Distal arthrogyriposis type 2B']
NM_000257.3(MYH7):c.2333A>G (p.Asp778Gly)	121913634	MYH7	[]	[]	['Familial hypertrophic cardiomyopathy 1', 'not specified']
NM_001127500.1(MET):c.3785A>G (p.Lys1262Arg)	121913677	MET	[]	[]	['Childhood hepatocellular carcinoma']
NM_000222.2(KIT):c.2459A>G (p.Asp820Gly)	121913682	KIT	[]	['AGAAYCATTCCTGATGTCTCTGG']	['Mast cell disease, systemic']
NM_000222.2(KIT):c.2386A>G (p.Arg796Gly)	121913684	KIT	[]	[]	[]
NM_006005.3(WFS1):c.1385A>G (p.Glu462Gly)	398123066	WFS1	[]	[]	['Cataract, nuclear total']
NM_000495.4(COL4A5):c.3925-2A>G	587776400	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NC_012920.1:m.13514A>G	587776440	MT-ND5	[]	[]	['Leigh disease']
NM_000021.3(PSEN1):c.488A>G (p.His163Arg)	63750590	PSEN1	[]	[]	['Alzheimer disease, type 3', 'not provided']
NM_000484.3(APP):c.2140A>G (p.Thr714Ala)	63750643	APP	[]	[]	['Alzheimer disease, type 1', 'not provided']
NM_173560.3(RFX6):c.224-12A>G	587776515	RFX6	[]	[]	['Mitchell-Riley syndrome']
NM_014043.3(CHMP2B):c.85A>G (p.Ile29Val)	63750818	CHMP2B	[]	[]	['Frontotemporal Dementia, Chromosome 3-Linked', 'Amyotrophic lateral sclerosis 17', 'not provided']
NM_000057.3(BLM):c.1088-2A>G	367543015	BLM	[]	[]	['Bloom syndrome']
NM_001011658.3(TRAPPC2):c.238+4T>C	587776753	-	[]	[]	['Spondyloepiphyseal dysplasia tarda']
NM_000151.3(G6PC):c.230+4A>G	587776757	G6PC	[]	['GTTCYTACCACTTAAAGACGAGG']	['Glycogen storage disease type 1A']
NM_000463.2(UGT1A1):c.1085-2A>G	587776766	-	['ACCYGAGATGCAAAA TAGGGAGG']	['ACCYGAGATGC AAAATAGGGAGG', 'GTGACCYGAGATGCAAAAATAGGG', 'GGTGACCYGAGATGCAAAAATAGG']	['Crigler Najjar syndrome, type 1']

NM_000330.3(RS1):c.286T>C (p.Trp96Arg)	61752063	-	[]	['TTCTTCGYGGAC TGCAAACAAGG']	['Juvenile retinoschisis', 'not provided']
NM_001024847.2(TGFBR2):c.1472-2A>G	587776770	TGFBR2	[]	[]	['Loeys-Dietz syndrome 2']
NM_000257.3(MYH7):c.5807A>G (p.Ter1936Trp)	367543053	MYH7	[]	[]	['Congenital myopathy with fiber type disproportion']
NM_000321.2(RB1):c.2490-1398A>G	587776791	RB1	[]	[]	['Retinoblastoma']
NM_024549.5(TCTN1):c.221-2A>G	367543065	TCTN1	[]	['AGCAACYGCAG AAAAAAGAGGGG', 'CAGCAACYGCAG AAAAAAGAGGGG']	['Joubert syndrome 13']
NM_000228.2(LAMB3):c.565-3T>C	587776813	LAMB3	[]	[]	['Adult junctional epidermolysis bullosa']
NM_015884.3(MBTPS2):c.1523A>G (p.Asn508Ser)	587776867	MBTPS2	[]	[]	['Keratosis pilaris decalvans']
NM_000174.4(GP9):c.182A>G (p.Asn61Ser)	5030764	GP9	['GGCTGYT GTTGCCA GCAGAAG G']	['GGCTGYTGTGG CCAGCAGAAGG']	['Bernard-Soulier syndrome type C']
NM_000894.2(LHB):c.221A>G (p.Gln74Arg)	5030773	LHB	[]	['CCACCYGAGGC AGGGGCGGCAGG']	['Isolated lutropin deficiency']
NM_000264.3(PTCH1):c.2479A>G (p.Ser827Gly)	199476092	-	[]	['CGTTACYGAAAC TCCTGTGTAGG']	['Gorlin syndrome', 'Holoprosencephaly 7', 'not specified', 'not provided']
NM_000021.3(PSEN1):c.415A>G (p.Met139Val)	63751037	PSEN1	[]	[]	['Alzheimer disease, type 3', 'not provided']
NM_000484.3(APP):c.2078A>G (p.Glu693Gly)	63751039	APP	[]	[]	['Alzheimer disease', 'Alzheimer disease, type 1', 'Cerebral amyloid angiopathy, APP-related', 'not provided']
NM_000117.2(EMD):c.450-2A>G	398123158	EMD	[]	['CGTTCCCYGAGG CAAAAGAGGGG']	['not provided']
RMRP:n.71A>G	199476103	RMRP	[]	['ACTTYCCCCTAG GCGGAAAGGG', 'GACTTYCCCCTA GGCGGAAAGGG', 'GGACTTYCCCCT AGGCGGAAAGG']	['Metaphyseal chondrodysplasia, McKusick type', 'Metaphyseal dysplasia without hypotrichosis']
m.14495A>G	199476106	MT-ND6	[]	[]	['Leber optic atrophy']
m.11084A>G	199476113	MT-ND4	[]	[]	['Juvenile myopathy, encephalopathy, lactic acidosis AND stroke']
NM_000551.3(VHL):c.233A>G (p.Asn78Ser)	5030804	VHL	['TGCGAYT GCAGAAG ATGACCTG G']	['GCGAYTGCAGA AGATGACCTGGG', 'TGCGAYTGCAGA AGATGACCTGG']	['Von Hippel-Lindau syndrome']

m.3397A>G	199476120	MT-ND1	[]	[]	['Alzheimer disease', 'Parkinson disease, late-onset']
m.4136A>G	199476121	MT-ND1	[]	[]	['Leber optic atrophy']
NM_003094.3(SNRPE):c.1A>G (p.Met1Val)	587776924	SNRPE	[]	[]	['Hypotrichosis 11']
NM_001310338.1(MGME1):c.743A>G (p.Tyr248Cys)	587776944	MGME1	[]	[]	['Mitochondrial DNA depletion syndrome 11']
NM_000249.3(MLH1):c.122A>G (p.Asp41Gly)	63751094	MLH1	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_138425.3(C12orf57):c.1A>G (p.Met1Val)	587776954	C12orf57	[]	[]	['Temtamy syndrome', 'Seizures', 'Corpus callosum abnormalities', 'Colobomatous microphthalmia', 'Global developmental delay']
NM_000277.1(PAH):c.1169A>G (p.Glu390Gly)	5030856	PAH	[]	['CTCYCTGCCACG TAATACAGGGG', 'ACTCYCTGCCAC GTAATACAGGG', 'AACTCYCTGCCA CGTAATACAGG']	['Phenylketonuria', 'Hyperphenylalaninemia, non-pku', 'not provided']
NM_000277.1(PAH):c.1241A>G (p.Tyr414Cys)	5030860	PAH	[]	['GGGTCGYAGCG AACTGAGAAGGG', 'TGGGTCGYAGCG AACTGAGAAGG']	['Phenylketonuria', 'Hyperphenylalaninemia, non-pku', 'not provided']
NM_000155.3(GALT):c.308A>G (p.Gln103Arg)	367543252	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_207352.3(CYP4V2):c.1091-2A>G	199476183	CYP4V2	[]	[]	['Bietti crystalline corneoretinal dystrophy']
NM_000518.4(HBB):c.*111A>G	63751128	HBB	[]	[]	[]
NM_000155.3(GALT):c.857A>G (p.Tyr286Cys)	367543262	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_000155.3(GALT):c.854A>G (p.Lys285Arg)	367543263	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_207352.3(CYP4V2):c.761A>G (p.His254Arg)	199476193	CYP4V2	[]	[]	['Bietti crystalline corneoretinal dystrophy']
NM_000155.3(GALT):c.968A>G (p.Tyr323Cys)	367543267	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']

NM_001142519.1(FAM111A):c.1012A>G (p.Thr338Ala)	587777014	FAM111A	[]	[]	['Gracile bone dysplasia']
NM_000132.3(F8):c.1660A>G (p.Ser554Gly)	137852419	F8	['AACYAGA GTAATAGC GGGTCAGG']	['AACYAGAGTAA TAGCGGGTCAGG']	['Hereditary factor VIII deficiency disease']
NM_020988.2(GNAO1):c.521A>G (p.Asp174Gly)	587777055	GNAO1	[]	['GGATGYCCTGCT CCGTGGGCTGG']	['Early infantile epileptic encephalopathy 17']
NM_000155.3(GALT):c.905-2A>G	398123187	GALT	[]	[]	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_015662.2(IFT172):c.4607T>C (p.Leu1536Pro)	587777080	IFT172	[]	[]	[]
NM_014754.2(PTDSS1):c.1058A>G (p.Gln353Arg)	587777088	PTDSS1	[]	[]	['Lenz-Majewski hyperostosis syndrome']
NM_003859.1(DPM1):c.742T>C (p.Ser248Pro)	587777114	-	[]	[]	['Congenital disorder of glycosylation type 1E']
NM_001018005.1(TPM1):c.742A>G (p.Lys248Glu)	199476319	TPM1	[]	[]	['Left ventricular noncompaction 9', 'not provided']
NM_004826.3(ECEL1):c.2278T>C (p.Cys760Arg)	587777129	ECEL1	[]	[]	['Arthrogyposis, distal, type 5d']
NM_014908.3(DOLK):c.2T>C (p.Met1Thr)	587777137	DOLK	[]	[]	['Congenital disorder of glycosylation type 1M']
NM_000350.2(ABCA4):c.4540-2A>G	61752435	ABCA4	[]	[]	['Stargardt disease 1', 'not provided']
NM_001128085.1(ASPA):c.433-2A>G	63751297	-	[]	[]	['Spongy degeneration of central nervous system']
NM_176787.4(PIGN):c.808T>C (p.Ser270Pro)	587777186	PIGN	[]	[]	['Multiple congenital anomalies-hypotonia-seizures syndrome 1']
NM_001165899.1(PDE4D):c.1850T>C (p.Ile617Thr)	587777188	PDE4D	['CTATAYT GTTCATCC CCTCTGGG'] , 'ACTATAYT GTTCATCC CCTCTGG']	['CTATAYTGTTCA TCCCCTCTGGG', 'ACTATAYTGTTT ATCCCCTCTGG']	['Acrodysostosis 2, with or without hormone resistance']
NM_005017.3(PCYT1A):c.571T>C (p.Phe191Leu)	587777195	PCYT1A	['GCATGYT TGCTCAA CACAGAGG']	['GCATGYTTGCTC CAACACAGAGG']	['Spondylometaphyseal dysplasia with cone-rod dystrophy']
NM_024301.4(FKRP):c.1A>G (p.Met1Val)	587777223	FKRP	[]	['CCGCAYGGGGC CGAAGTCTGGG', 'GCCGCAYGGGGC CGAAGTCTGGG', 'AGCCGCAYGGGG CCGAAGTCTGG']	['Congenital muscular dystrophy-dystroglycanopathy with brain and eye anomalies type A5']

NM_198947.3(FAM111B):c.1879A>G (p.Arg627Gly)	587777237	FAM111B	[]	[]	['Poikiloderma, hereditary fibrosing, with tendon contractures, myopathy, and pulmonary fibrosis']
NM_003638.2(ITGA8):c.2982+2T>C	587777279	ITGA8	[]	[]	['Renal adysplasia']
NM_199189.2(MATR3):c.1864A>G (p.Thr622Ala)	587777301	MATR3	['CGGYTGA ACTCTCAG TCTTCTGG']	['CGGYTGA ACTCTCT CAGTCTTCTGG']	['Myopathy, distal, 2']
NM_001739.1(CA5A):c.697T>C (p.Ser233Pro)	587777316	CA5A	[]	[]	['Carbonic anhydrase VA deficiency, hyperammonemia due to']
NM_005051.2(QARS):c.169T>C (p.Tyr57His)	587777333	QARS	[]	[]	['Microcephaly, progressive, with seizures and cerebral and cerebellar atrophy']
NM_002234.3(KCNA5):c.143A>G (p.Glu48Gly)	587777336	KCNA5	[]	[]	['Atrial fibrillation, familial, 7']
NM_021803.3(IL21):c.146T>C (p.Leu49Pro)	587777338	IL21	[]	[]	['Common variable immunodeficiency 11']
NM_000132.3(F8):c.6794A>G (p.Gln2265Arg)	137852470	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_178014.3(TUBB):c.895A>G (p.Met299Val)	587777355	TUBB	[]	[]	['Cortical dysplasia, complex, with other brain malformations 6']
NM_005957.4(MTHFR):c.1969T>C (p.Ter657Arg)	768434408	MTHFR	[]	[]	['Homocysteinemia due to MTHFR deficiency']
NM_005359.5(SMAD4):c.425-6A>G	377767327	SMAD4	[]	[]	['Juvenile polyposis syndrome']
NM_022068.3(PIEZO2):c.8215T>C (p.Ser2739Pro)	587777454	PIEZO2	[]	[]	['Oculomeelic amyoplasia']
NM_003108.3(SOX11):c.347A>G (p.Tyr116Cys)	587777479	SOX11	[]	['GTACTTGYAGTC GGGGTAGTCGG']	['Mental retardation, autosomal dominant 27']
NM_021072.3(HCN1):c.814T>C (p.Ser272Pro)	587777493	HCN1	[]	[]	['Epileptic encephalopathy, early infantile, 24']
NM_020435.3(GJC2):c.-170A>G	587777496	GJC2	[]	['TTGYTCCCCCT CGGCCTCAGG', 'ATTGYTCCCCCT CGGCCTCAGG']	['Leukodystrophy, hypomyelinating, 2']
NM_022552.4(DNMT3A):c.1943T>C (p.Leu648Pro)	587777507	DNMT3A	[]	['CTCCYGGTGCTG AAGGACTTGGG', 'GCTCCYGGTGCT GAAGGACTTGG']	['Tatton-Brown-rahman syndrome']
NM_022552.4(DNMT3A):c.2705T>C	587777510	DNMT3A	[]	[]	['Tatton-Brown-rahman syndrome']

(p.Phe902Ser)					
NM_000223.3(KRT12):c.403A>G (p.Arg135Gly)	58410481	KRT12	[]	[]	['Meesman corneal dystrophy', 'not provided']
NM_000232.4(SGCB):c.1A>G (p.Met1Val)	398123262	SGCB	[]	[]	['Limb-girdle muscular dystrophy, type 2E', 'not provided']
NM_020630.4(RET):c.2342A>G (p.Gln781Arg)	377767416	RET	[]	[]	['MEN2 phenotype: Unclassified']
NM_018400.3(SCN3B):c.482T>C (p.Met161Thr)	587777557	SCN3B	[]	['AATCAYGATGTA CATCCTTCTGG']	['Atrial fibrillation, familial, 16']
NM_001030001.2(RPS29):c.149T>C (p.Ile50Thr)	587777569	RPS29	[]	['GATAYCGGTTTC ATTAAGGTAGG']	['Diamond-Blackfan anemia 13']
NM_177550.4(SLC13A5):c.1463T>C (p.Leu488Pro)	587777578	SLC13A5	[]	[]	['Epileptic encephalopathy, early infantile, 25']
NM_002880.3(RAF1):c.1808T>C (p.Leu603Pro)	587777586	RAF1	[]	[]	['Cardiomyopathy, dilated, 1NN']
NM_025150.4(TARS2):c.695+3A>G	587777594	TARS2	[]	[]	['Combined oxidative phosphorylation deficiency 21']
NM_001759.3(CCND2):c.838A>G (p.Thr280Ala)	587777618	CCND2	[]	[]	['Megalencephaly-polymicrogyria-polydactyly-hydrocephalus syndrome 3']
NM_153334.6(SCARF2):c.190T>C (p.Cys64Arg)	587777657	SCARF2	[]	['CCACGYGCTGCG CTGGCTGGAGG']	['Marden Walker like syndrome']
NM_005726.5(TSFM):c.57+4A>G	587777689	TSFM	[]	['ACTTCYCACCGG GTAGCTCCCGG']	['Combined oxidative phosphorylation deficiency 3']
NM_000255.3(MUT):c.329A>G (p.Tyr110Cys)	796052005	MUT	[]	['GCAYACTGGCG GATGGTCCAGGG', 'AGCAYACTGGCG GATGGTCCAGG']	['not provided']
NM_021870.2(FGG):c.1210T>C (p.Ser404Pro)	587777720	FGG	[]	[]	['Hypodysfibrinogenemia']
NM_017617.3(NOTCH1):c.1285T>C (p.Cys429Arg)	587777736	NOTCH1	['GGCAAGY GCATCAAC ACGCTGGG']	['GGCAAGYGCAT CAACACGCTGGG', 'GGGCAAGYGCAT CAACACGCTGG']	['Adams-Oliver syndrome 1', 'Adams-Oliver syndrome 5']
NM_014946.3(SPAST):c.1688-2A>G	587777752	SPAST	['TTCYGTA AAACATAA AAGTCAGG']	['TTCYGTA AAAACA TAAAAGTCAGG']	['Spastic paraplegia 4, autosomal dominant']
NM_014946.3(SPAST):c.1245+4A>G	587777755	SPAST	[]	[]	['Spastic paraplegia 4, autosomal dominant']
NM_014946.3(SPAST):c.1216A>G	587777757	SPAST	[]	[]	['Spastic paraplegia 4, autosomal

(p.Ile406Val)					dominant']
NM_144596.3(TTC8): c.115-2A>G	587777809	TTC8	[]	['GTTCCYGGAAA GCATTAAGAAGG']	['Retinitis pigmentosa 51']
NM_170784.2(MKKS) :c.110A>G (p.Tyr37Cys)	74315396	MKKS	[]	[]	['Bardet-Biedl syndrome 6', 'McKusick Kaufman syndrome']
NM_000252.2(MTM1) :c.566A>G (p.Asn189Ser)	132630302	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1) :c.1190A>G (p.Tyr397Cys)	132630303	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_152384.2(BBS5): c.522+3A>G	587777828	BBS5	[]	[]	['Bardet-Biedl syndrome 5']
NM_001205019.1(GK)):c.880A>G (p.Asn294Asp)	132630331	GK	[]	[]	['Deficiency of glycerol kinase']
NM_000166.5(GJB1): c.580A>G (p.Met194Val)	587777878	GJB1	[]	['TAGCAYGAAGA CGGTGAAGACGG']	['X-linked hereditary motor and sensory neuropathy']
NM_000311.3(PRNP): c.547A>G (p.Thr183Ala)	74315411	PRNP	[]	[]	['Genetic prion diseases', 'Spongiform encephalopathy with neuropsychiatric features']
NM_144773.2(PROKR2): c.629A>G (p.Gln210Arg)	74315417	PROKR2	[]	[]	['Kallmann syndrome 3']
NM_000531.5(OTC):c .919A>G (p.Lys307Glu)	796052013	OTC	[]	[]	['not provided']
NM_001029871.3(RSPO4): c.194A>G (p.Gln65Arg)	74315420	RSPO4	[]	['CGTACYGGCGG ATGCCTTCCCGG']	['Anonychia']
NM_004333.4(BRAF): c.770A>G (p.Gln257Arg)	180177035	BRAF	[]	[]	['Noonan syndrome 7', 'Cardiofaciocutaneous syndrome', 'Rasopathy', 'not provided']
NM_004333.4(BRAF): c.1495A>G (p.Lys499Glu)	180177037	BRAF	[]	[]	['Cardiofaciocutaneous syndrome', 'Rasopathy']
NM_198056.2(SCN5A)):c.5297T>A (p.Met1766Lys)	752476527	SCN5A	[]	[]	['not provided']
NM_000030.2(AGXT) :c.248A>G (p.His83Arg)	180177186	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_000030.2(AGXT) :c.424-2A>G (p.Gly_142Gln145del)	180177219	AGXT	[]	['AGGCCCYGAGG AAGCAGGGACGG']	['Primary hyperoxaluria, type I']
NM_198578.3(LRRK2)):c.5096A>G (p.Tyr1699Cys)	35801418	LRRK2	[]	[]	['Parkinson disease 8, autosomal dominant']
NM_002693.2(POLG):	367610201	POLG	[]	['CTCAYGGCACTT	['not provided']

c.1808T>C (p.Met603Thr)				ACCTGGGATGG']	
NM_000030.2(AGXT) :c.596-2A>G	180177245	AGXT	[]	[]	['Primary hyperoxaluria, type I']
NM_020223.3(FAM20C):c.1364-2A>G	796051853	FAM20C	[]	[]	['Raine syndrome']
NM_012203.1(GRHP R):c.84-2A>G	180177319	GRHPR	[]	['TCACAGCYGCG GGGAAAGGGAGG']	['Primary hyperoxaluria, type II']
NM_006017.2(PROM1):c.2077-521A>G	796051882	PROM1	[]	[]	['Cone-rod dystrophy 2']
NM_012203.1(GRHP R):c.934A>G (p.Asn312Asp)	180177324	GRHPR	['CAAGTYG TTAGCTGC CAACAAGG']	['CAAGTYGTTAGC TGCCAACAAGG']	['Primary hyperoxaluria, type II']
NM_000016.5(ACADM):c.329A>G (p.Glu110Gly)	796051900	ACADM	[]	[]	['not provided']
NM_004453.3(ETFDH):c.929A>G (p.Tyr310Cys)	796051958	ETFDH	[]	[]	['not provided']
NM_000255.3(MUT):c.1885A>G (p.Arg629Gly)	796052004	MUT	[]	[]	['not provided']
NM_012434.4(SLC17A5):c.548A>G (p.His183Arg)	119491109	SLC17A5	[]	[]	['Sialic acid storage disease, severe infantile type']
NM_000328.2(RPGR):c.155-2A>G	62638632	RPGR	[]	[]	['Retinitis pigmentosa 15', 'not provided']
NM_005557.3(KRT16):c.373A>G (p.Asn125Asp)	58608173	KRT16	[]	[]	['Pachyonychia congenita, type 1', 'not provided']
NM_000532.4(PCCB):c.655-2A>G	796052020	PCCB	[]	[]	['not provided']
NM_000030.2(AGXT):c.777-2A>G	796052068	AGXT	[]	['GGTACCYGGAA GACACGAGGGGG', 'TGGTACCYGGAA GACACGAGGGG']	['Primary hyperoxaluria, type I']
NM_000121.3(EPOR):c.1460A>G (p.Asn487Ser)	62638745	EPOR	['AGGGYTG GAGTAGGG GCCATCGG']	['AGGGYTGGAGT AGGGGCCATCGG']	['Acute myeloid leukemia, M6 type', 'Familial erythrocytosis, 1']
NM_000552.3(VWF):c.1583A>G (p.Asn528Ser)	61754010	VWF	[]	['TGCCAYTGTAAT TCCCACACAGG']	['von Willebrand disease, type 2a', 'not provided']
NM_001918.3(DBT):c.1017_1018insNC_000001.11:g.100207187_100207312	796052135	DBT	[]	[]	['Intermediate maple syrup urine disease type 2']
NM_001243473.1(B9D1):c.400+2T>C	143149764	B9D1	[]	[]	['Meckel syndrome, type 9', 'not provided']
NM_001165963.1(SCN1A):c.4766T>G (p.Val1589Gly)	764037830	-	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_000321.2(RB1):c.	587778866	RB1	[]	['ATTYCAATGGCT	['Retinoblastoma']

1927A>G (p.Lys643Glu)				TCTGGGTCTGG']	
NM_006331.7(EMG1) :c.257A>G (p.Asp86Gly)	74435397	EMG1	[]	['ATAYCTGGCCGC GCTTCCCCAGG']	['Bowen-Conradi syndrome']
NM_000249.3(MLH1) :c.113A>G (p.Asn38Ser)	587778888	MLH1	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_017777.3(MKS1) :c.1382A>G (p.Tyr461Cys)	730882120	MKS1	[]	[]	['Bardet-Biedl syndrome 13']
NM_000261.1(MYOC) :c.1010A>G (p.Gln337Arg)	74315335	MYOC	[]	[]	['Primary open angle glaucoma juvenile onset 1']
NM_152515.4(CKAP2 L):c.2T>C (p.Met1Thr)	548949031	CKAP2L	[]	[]	['Filippi syndrome']
NM_000156.5(GAMT) :c.1A>G (p.Met1Val)	796052527	GAMT	[]	['CGCTCAYGCTGC AGGCTGGACGG']	['not provided']
NM_000833.4(GRIN2 A):c.1930A>G (p.Ser644Gly)	796052544	GRIN2A	[]	[]	['not provided']
NM_000144.4(FXN):c .385-2A>G	140987490	FXN	[]	[]	['Friedreich ataxia']
NM_172107.2(KCNQ 2):c.297-2A>G	796052615	KCNQ2	[]	[]	['not provided']
NM_172107.2(KCNQ 2):c.611A>G (p.Gln204Arg)	796052624	KCNQ2	[]	[]	['not provided']
NM_172107.2(KCNQ 2):c.848A>G (p.Lys283Arg)	796052637	KCNQ2	[]	['GTACYTGTCCCC GTAGCCAATGG']	['not provided']
NM_052859.3(RFT1): c.887T>A (p.Ile296Lys)	772820136	RFT1	[]	[]	['Congenital disorder of glycosylation type 1N']
NM_000553.4(WRN): c.561A>G (p.Lys187=)	775802030	WRN	[]	[]	['Werner syndrome']
NM_194277.2(FRMD 7):c.556A>G (p.Met186Val)	786205896	FRMD7	[]	[]	['Infantile nystagmus, X- linked']
NM_000535.5(PMS2): c.989-2A>G	587779347	PMS2	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms', 'Hereditary cancer- predisposing syndrome']
NM_000203.4(IDUA): c.1874A>G (p.Tyr625Cys)	587779401	IDUA	[]	[]	['Hurler syndrome']
NM_001105243.1(PC DH19):c.1019A>G (p.Asn340Ser)	796052839	PCDH19	[]	[]	['not provided']
NM_002693.2(POLG): c.2840A>G (p.Lys947Arg)	796052891	POLG	[]	[]	['not provided']
NM_032228.5(FAR1): c.1094A>G	724159963	FAR1	[]	['GATAYCATACA GGAATGCTGGGG']	['Peroxisomal fatty acyl-coa reductase 1

(p.Asp365Gly)				'AGATAYCATACA GGAATGCTGGG', 'TAGATAYCATAC AGGAATGCTGG']	disorder']
NM_000090.3(COL3A1):c.2284-2A>G (p.Gly762_Lys779del)	587779558	COL3A1	[]	[]	['Ehlers-Danlos syndrome, type 4']
NM_014305.3(TGDS):c.269A>G (p.Glu90Gly)	724160004	TGDS	[]	[]	['Catel Manzke syndrome']
NM_014305.3(TGDS):c.892A>G (p.Asn298Asp)	724160005	TGDS	[]	[]	['Catel Manzke syndrome']
NM_000090.3(COL3A1):c.997-2A>G (p.Gly333_Lys350del+)	587779602	COL3A1	[]	[]	['Ehlers-Danlos syndrome, type 4']
NM_002185.3(IL7R):c.197T>C (p.Ile66Thr)	1494558	IL7R	[]	[]	['Severe combined immunodeficiency, autosomal recessive, T cell-negative, B cell-positive, NK cell-positive', 'not specified']
NM_000277.1(PAH):c.974A>G (p.Tyr325Cys)	62508578	PAH	[]	[]	['Phenylketonuria', 'not provided']
NM_000090.3(COL3A1):c.997-10A>G (p.Pro332_Gly333insFQ)	587779670	COL3A1	[]	[]	['Ehlers-Danlos syndrome, type 4']
NM_000090.3(COL3A1):c.3202-2A>G (p.Gly1068_Pro1085del)	587779682	COL3A1	[]	[]	['Ehlers-Danlos syndrome, type 4']
NM_000090.3(COL3A1):c.1762-2A>G (p.Gly588_Gln605del)	587779722	COL3A1	[]	['CACCCYAAAGA AGAAGTGGTCGG']	['Ehlers-Danlos syndrome, type 4']
NM_021007.2(SCN2A):c.4036A>G (p.Ile1346Val)	796053135	SCN2A	[]	[]	['not provided']
m.13637A>G	200855215	MT-ND5	[]	[]	['Leber optic atrophy']
NM_021007.2(SCN2A):c.387-2A>G	796053169	SCN2A	['AATAAAG YAGAATAT CGTCAAGG']	['AATAAAGYAGA ATATCGTCAAGG']	['not provided']
NM_021007.2(SCN2A):c.851A>G (p.Asp284Gly)	796053173	SCN2A	[]	[]	['not provided']
NM_006516.2(SLC2A1):c.848A>G (p.Gln283Arg)	796053251	SLC2A1	[]	[]	['not provided']
NM_006516.2(SLC2A1):c.19-2A>G	796053272	SLC2A1	[]	[]	['not provided']
NM_000136.2(FANCC):c.-78-2A>G	587779898	FANCC	[]	[]	['Hereditary cancer-predisposing syndrome']
m.8296A>G	118192102	MT-TK	[]	['TTTACAGYGGGC']	['Diabetes-deafness']

				TCTAGAGGGGG']	syndrome maternally transmitted']
NM_005360.4(MAF):c.172A>G (p.Thr58Ala)	727502767	MAF	[]	[]	['Cataracts, congenital, with sensorineural deafness, down syndrome-like facial appearance, short stature, and mental retardation']
NM_001145901.1(SARS2):c.1175A>G (p.Asp392Gly)	727502784	SARS2	[]	[]	['Hyperuricemia, pulmonary hypertension, renal failure, and alkalosis']
NM_001077494.3(NFKB2):c.2594A>G (p.Asp865Gly)	727502787	NFKB2	[]	['CTGYCTTCCTTC ACCTCTGCTGG']	['Common variable immunodeficiency 10']
NM_002238.3(KCNH1):c.1399A>G (p.Ile467Val)	727502819	KCNH1	[]	[]	['Zimmermann-Laband syndrome', 'Temple-Baraitser syndrome']
NM_172362.2(KCNH1):c.1508A>G (p.Gln503Arg)	727502821	KCNH1	[]	[]	['Temple-Baraitser syndrome']
NM_000546.5(TP53):c.701A>G (p.Tyr234Cys)	587780073	TP53	[]	[]	['Li-Fraumeni syndrome', 'Hereditary cancer-predisposing syndrome']
NM_003060.3(SLC22A5):c.694A>C (p.Thr232Pro)	188698686	SLC22A5	[]	[]	['not provided']
NM_000540.2(RYR1):c.14591A>G (p.Tyr4864Cys)	118192146	RYR1	[]	[]	['Central core disease', 'not provided']
NM_058216.2(RAD51C):c.706-2A>G	587780259	RAD51C	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000501.3(ELN):c.800-2A>G	727503027	ELN	[]	[]	['Supravalvar aortic stenosis', 'not provided']
NM_000117.2(EMD):c.266-2A>G	727503036	EMD	[]	['AGCCYTGGGAA GGGGGCAGCGG']	['Emery-Dreifuss muscular dystrophy 1, X-linked']
NM_003242.5(TGFBR2):c.1273A>G (p.Met425Val)	104893817	TGFBR2	[]	[]	['Loeys-Dietz syndrome 2']
NM_153638.2(PANK2):c.700A>G (p.Thr234Ala)	137852965	PANK2	[]	[]	[]
NM_005861.3(STUB1):c.194A>G (p.Asn65Ser)	690016544	STUB1	[]	['GGCCCGGYTGGT GTAATACACGG']	['Spinocerebellar ataxia, autosomal recessive 16']
NM_005360.4(MAF):c.890A>G (p.Lys297Arg)	121917736	MAF	[]	[]	['Cataract, pulverulent, juvenile-onset']
NM_005211.3(CSF1R)	690016554	CSF1R	[]	['GTATCYGGGAG	['Hereditary diffuse

:c.2655-2A>G				ATAGGACAGAGG']	leukoencephalopathy with spheroids']
NM_003361.3(UMOD):c.383A>G (p.Asn128Ser)	121917770	UMOD	['CACAYTGACACATGTGGCCAGGG']	['CACAYTGACACATGTGGCCAGGG', 'CCACAYTGACACATGTGGCCAGG']	['Familial juvenile gout']
NM_000256.3(MYBP C3):c.2234A>G (p.Asp745Gly)	727503190	MYBPC3	[]	[]	['Familial hypertrophic cardiomyopathy 4', 'Familial hypertrophic cardiomyopathy 1', 'not specified']
NM_172107.2(KCNQ2):c.1A>G (p.Met1Val)	118192185	KCNQ2	[]	['GCACCAYGGTG CCTGGCGGGAGG']	['Benign familial neonatal seizures 1']
NM_000256.3(MYBP C3):c.1213A>G (p.Met405Val)	727503207	MYBPC3	[]	[]	['Cardiomyopathy', 'not specified']
NM_000021.3(PSEN1):c.998A>G (p.Asp333Gly)	121917809	PSEN1	[]	[]	['Primary dilated cardiomyopathy', 'Cardiomyopathy, dilated, 1u', 'Heart failure']
NM_021954.3(GJA3):c.188A>G (p.Asn63Ser)	121917823	GJA3	[]	[]	['Zonular pulverulent cataract 3']
NM_000322.4(PRP H2):c.422A>G (p.Tyr141Cys)	61755781	PRPH2	[]	[]	['Macular dystrophy, vitelliform, adult-onset', 'Patterned dystrophy of retinal pigment epithelium', 'not provided']
NM_007035.3(KERA):c.740A>G (p.Asn247Ser)	121917858	KERA	[]	[]	['Cornea plana 2']
NM_002181.3(IHH):c.284A>G (p.Glu95Gly)	121917859	IHH	[]	[]	['Brachydactyly type A1']
NM_000257.3(MYH7):c.1157A>G (p.Tyr386Cys)	727503269	MYH7	[]	[]	['Primary familial hypertrophic cardiomyopathy']
NM_000097.5(CPOX):c.1210A>G (p.Lys404Glu)	121917868	CPOX	[]	[]	['Harderoporphyria']
NM_012064.3(MIP):c.401A>G (p.Glu134Gly)	121917869	MIP	[]	['AGATCYCCACTG TGGTTGCCTGG']	['Cataract 15, multiple types']
NM_025243.3(SLC19A3):c.1264A>G (p.Thr422Ala)	121917884	SLC19A3	[]	[]	['Basal ganglia disease, biotin-responsive']
NM_000373.3(UMPS):c.286A>G (p.Arg96Gly)	121917890	UMPS	[]	[]	['Orotic aciduria']
NM_000536.3(RAG2):c.115A>G (p.Arg39Gly)	121917897	RAG2	[]	[]	['Histiocytic medullary reticulosis']
NM_130838.1(UBE3A):c.1694-2A>G	587780579	UBE3A	[]	[]	['Angelman syndrome']
NM_016335.4(PROD)	2904551	PRODH	[]	[]	['Proline

H):c.1322T>C (p.Leu441Pro)					dehydrogenase deficiency', 'Schizophrenia 4']
NM_006920.4(SCN1A):c.4352A>G (p.Tyr1451Cys)	121917962	-	[]	[]	['Severe myoclonic epilepsy in infancy', 'not provided']
NM_006920.4(SCN1A):c.1876A>G (p.Ser626Gly)	121917990	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy', 'Generalized epilepsy']
NM_000478.4(ALPL):c.1250A>G (p.Asn417Ser)	121918014	ALPL	[]	['AGGCCCA YTGCC ATACAGGATGG']	['Infantile hypophosphatasia']
NM_000174.4(GP9):c.110A>G (p.Asp37Gly)	121918036	GP9	[]	['GCAGYCCACCC ACAGCCCCATGG']	['Bernard-Soulier syndrome type C']
NM_002693.2(POLG):c.2591A>G (p.Asn864Ser)	121918050	POLG	[]	[]	['Mitochondrial DNA depletion syndrome 4B, MNGIE type']
NM_000374.4(UROD):c.932A>G (p.Tyr311Cys)	121918061	UROD	[]	[]	['Hepatoerythropoietic porphyria']
NM_000217.2(KCNA1):c.763A>G (p.Asn255Asp)	121918067	KCNA1	[]	[]	[]
NM_000371.3(TTR):c.238A>G (p.Thr80Ala)	121918070	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis', 'Cardiomyopathy']
NM_000371.3(TTR):c.401A>G (p.Tyr134Cys)	121918075	TTR	['GGAGYAG GGGCTCAG CAGGGCGG', 'ATAGGAG YAGGGGCT CAGCAGGG']	['GGAGYAGGGGC TCAGCAGGGCGG', 'ATAGGAGYAGGG GCTCAGCAGGG']	['Amyloidogenic transthyretin amyloidosis']
NM_000371.3(TTR):c.205A>G (p.Thr69Ala)	121918081	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis']
NM_000371.3(TTR):c.379A>G (p.Ile127Val)	121918089	TTR	[]	['CGGCAAYGGTG TAGCGGCGGGG', 'GCGGCAAYGGTG TAGCGGCGGGG']	['Amyloidogenic transthyretin amyloidosis']
NM_000371.3(TTR):c.113A>G (p.Asp38Gly)	121918098	TTR	[]	[]	['Amyloidogenic transthyretin amyloidosis', 'AMYLOIDOSIS, LEPTOMENINGEAL, TRANSTHYRETIN-RELATED']
NM_000823.3(GHRHR):c.985A>G (p.Lys329Glu)	121918121	GHRHR	[]	['CGACTYGGAGA GACGCCTGCAGG']	['Isolated growth hormone deficiency type 1B']
NM_000275.2(OCA2):c.1465A>G (p.Asn489Asp)	121918170	OCA2	['GACATYT GGAGGGTC CCCGATGG']	['GACATYTGGAG GGTCCCCGATGG']	['Tyrosinase-positive oculocutaneous albinism']

NM_000181.3(GUSB):c.1484A>G (p.Tyr495Cys)	121918178	GUSB	[]	[]	['Mucopolysaccharidosis type VII']
NM_018122.4(DARS2):c.133A>G (p.Ser45Gly)	121918209	DARS2	[]	[]	['Leukoencephalopathy with Brainstem and Spinal Cord Involvement and Lactate Elevation']
NM_015697.7(COQ2):c.890A>G (p.Tyr297Cys)	121918230	COQ2	[]	[]	['Coenzyme Q10 deficiency, primary 1']
NM_015697.7(COQ2):c.683A>G (p.Asn228Ser)	121918232	COQ2	[]	[]	['Coenzyme Q10 deficiency, primary 1']
NM_015384.4(NIPBL):c.7289A>G (p.Tyr2430Cys)	121918265	NIPBL	[]	[]	['Cornelia de Lange syndrome 1']
NM_004183.3(BEST1):c.707A>G (p.Tyr236Cys)	121918291	BEST1	[]	[]	['Vitreoretinopathy dominant']
NM_014362.3(HIBCH):c.365A>G (p.Tyr122Cys)	121918329	HIBCH	[]	[]	['Beta-hydroxyisobutyryl-CoA deacylase deficiency']
NM_015335.4(MED13L):c.6068A>G (p.Asp2023Gly)	121918333	MED13L	[]	['ATATCAYCTAGAGGGAAGGGGG', 'CATATCAYCTAGAGGGAAGGGGG']	['Transposition of great arteries']
NM_015040.3(PIKFYVE):c.3308A>G (p.Lys1103Arg)	121918336	PIKFYVE	[]	[]	['Fleck corneal dystrophy']
NM_006306.3(SMC1A):c.2974-2A>G	727503774	SMC1A	[]	[]	['Congenital muscular hypertrophy-cerebral syndrome']
NM_002633.2(PGM1):c.343A>G (p.Thr115Ala)	121918371	PGM1	[]	[]	['Congenital disorder of glycosylation type 1t']
NM_000040.1(APOC3):c.280A>G (p.Thr94Ala)	121918381	APOC3	['CTGAAGYTGCTGACCTCAGGG', 'GCTGAAGYTGCTGACCTCAGG']	['CTGAAGYTGCTGACCTCAGGG', 'GCTGAAGYTGCTGACCTCAGG']	[]
NM_000040.1(APOC3):c.232A>G (p.Lys78Glu)	121918382	APOC3	[]	[]	['Hyperalphalipoproteinemia 2']
NM_001146040.1(GLRA1):c.910A>G (p.Lys304Glu)	121918412	GLRA1	[]	[]	['Hyperkplexia hereditary']
NM_000171.3(GLRA1):c.523A>G (p.Met175Val)	121918414	GLRA1	[]	[]	['Hyperkplexia hereditary']
NM_021957.3(GYS2):c.116A>G (p.Asn39Ser)	121918423	GYS2	[]	[]	['Hypoglycemia with deficiency of glycogen synthetase in the liver']

NM_002834.3(PTPN11):c.188A>G (p.Tyr63Cys)	121918459	PTPN11	[]	[]	['Noonan syndrome', 'Noonan syndrome 1', 'Rasopathy', 'not provided']
NM_013382.5(POMT2):c.1726-2A>G	727503873	POMT2	[]	[]	['not provided']
NM_002834.3(PTPN11):c.236A>G (p.Gln79Arg)	121918466	PTPN11	[]	[]	['Noonan syndrome', 'Noonan syndrome 1', 'Rasopathy', 'not provided']
NM_000313.3(PROS1):c.773A>G (p.Asn258Ser)	121918473	PROS1	[]	[]	['Protein S deficiency']
NM_000313.3(PROS1):c.586A>G (p.Lys196Glu)	121918474	PROS1	[]	[]	['Protein S deficiency']
NM_000141.4(FGFR2):c.983A>G (p.Tyr328Cys)	121918493	FGFR2	[]	[]	['Crouzon syndrome']
NM_000141.4(FGFR2):c.874A>G (p.Lys292Glu)	121918500	FGFR2	['TGCTYGA TCCACTGG ATGTGGGG ']	['TGCTYGATCCAC TGGATGTGGGG', 'GTGCTYGATCCA CTGGATGTGGG', 'CGTGCTYGATCC ACTGGATGTGG']	['Crouzon syndrome']
NM_000141.4(FGFR2):c.1576A>G (p.Lys526Glu)	121918507	FGFR2	[]	[]	['Crouzon syndrome', 'Scaphocephaly, maxillary retrusion, and mental retardation']
NM_002739.3(PRKCG):c.380A>G (p.Gln127Arg)	121918515	PRKCG	[]	[]	['Spinocerebellar ataxia 14']
NM_002739.3(PRKCG):c.1081A>G (p.Ser361Gly)	121918517	PRKCG	[]	[]	['Spinocerebellar ataxia 14']
NM_000098.2(CPT2):c.359A>G (p.Tyr120Cys)	121918528	CPT2	['GATAGGY ACATATCA AACCAGGG , 'AGATAGG YACATATC AAACCAGG ']	['GATAGGYACAT ATCAAACCAGGG', 'AGATAGGYACAT ATCAAACCAGG']	['Carnitine palmitoyltransferase II deficiency, infantile']
NM_005587.2(MEF2A):c.788A>G (p.Asn263Ser)	121918530	MEF2A	['CCAAGAY TACCACCA CCTGGTGG ']	['AGAYTACCACC ACCTGGTGGAGG', 'CCAAGAYTACCA CCACCTGGTGG']	[]
NM_006204.3(PDE6C):c.1363A>G (p.Met455Val)	121918539	PDE6C	[]	[]	['Achromatopsia 5']
NM_017654.3(SAMD9):c.4483A>G (p.Lys1495Glu)	121918554	SAMD9	[]	[]	['Tumoral calcinosis, familial, normophosphatemic']
NM_000191.2(HMGCL):c.698A>G (p.His233Arg)	727503963	HMGCL	[]	[]	['not provided']

NM_020166.4(MCCC1):c.137-2A>G	727504006	MCCC1	[]	[]	['3 Methylcrotonyl-CoA carboxylase 1 deficiency', 'not provided']
NM_001035.2(RYR2):c.12602A>G (p.Gln4201Arg)	121918605	RYR2	[]	['CGCCAGCYGCATTTCAAAGATGG']	['Catecholaminergic polymorphic ventricular tachycardia']
NM_002764.3(PRPS1):c.343A>G (p.Met115Val)	587781262	PRPS1	[]	['TAGCAYATTTGCAACAAGCTTGG']	['Charcot-Marie-Tooth disease, X-linked recessive, type 5', 'Deafness, high-frequency sensorineural, X-linked']
NM_001161766.1(AH1CY):c.344A>G (p.Tyr115Cys)	121918608	AH1CY	[]	['GCGGGYACTTGTGTGGATGAGG']	['Hypermethioninemia with s-adenosylhomocysteine hydrolase deficiency']
NM_000702.3(ATP1A2):c.1033A>G (p.Thr345Ala)	121918613	ATP1A2	[]	['CTGYCAGGGTCAAGCACACCTGG']	['Familial hemiplegic migraine type 2']
NM_003126.2(SPTA1):c.143A>G (p.Lys48Arg)	121918644	SPTA1	[]	[]	['Hereditary pyropoikilocytosis']
NM_001024858.2(SPTB):c.1A>G (p.Met1Val)	121918651	SPTB	[]	[]	['Elliptocytosis 3']
NM_000899.4(KITLG):c.107A>G (p.Asn36Ser)	121918653	KITLG	[]	[]	['Familial progressive hyperpigmentation with or without hypopigmentation']
NM_198253.2(TERT):c.2315A>G (p.Tyr772Cys)	121918663	TERT	[]	[]	['Aplastic anemia', 'PULMONARY FIBROSIS AND/OR BONE MARROW FAILURE, TELOMERE-RELATED, 1']
NM_001063.3(TF):c.1936A>G (p.Lys646Glu)	121918678	TF	[]	[]	[]
NM_000535.5(PMS2):c.904-2A>G	587781339	PMS2	[]	['GCAGACCYGCAAAAATACAAGG']	['Hereditary cancer-predisposing syndrome']
NM_001128177.1(THRB):c.1324A>G (p.Met442Val)	121918691	THRB	[]	['CTTCAYGTGCAGGAAGCGGCTGG']	['Thyroid hormone resistance, generalized, autosomal dominant']
NM_001128177.1(THRB):c.1327A>G (p.Lys443Glu)	121918692	THRB	[]	['CCACCTYCATGTGCAGGAAGCGG']	['Thyroid hormone resistance, generalized, autosomal dominant']
NM_001128177.1(THRB):c.1009A>G	121918709	THRB	[]	[]	['Thyroid hormone resistance, selective']

(p.Thr337Ala)					pituitary']
NM_004612.3(TGFBR1):c.1199A>G (p.Asp400Gly)	121918711	TGFBR1	['ATAGATG YCAGCACG TTTGAAGG']	['ATAGATGYCAG CACGTTTGAAGG']	['Loeys-Dietz syndrome 1']
NM_000359.2(TGM1) :c.1469A>G (p.Asp490Gly)	121918724	TGM1	[]	[]	['Autosomal recessive congenital ichthyosis 1']
NM_000257.3(MYH7) :c.1727A>G (p.His576Arg)	727504238	MYH7	[]	[]	['Familial hypertrophic cardiomyopathy 1', 'Cardiomyopathy', 'not specified']
NM_000257.3(MYH7) :c.1954A>G (p.Arg652Gly)	727504239	MYH7	[]	[]	['Primary familial hypertrophic cardiomyopathy', 'Familial hypertrophic cardiomyopathy 1']
NM_000257.3(MYH7) :c.1496A>G (p.Glu499Gly)	727504270	MYH7	[]	[]	['Cardiomyopathy', 'not specified']
NM_000257.3(MYH7) :c.2539A>G (p.Lys847Glu)	727504310	MYH7	[]	[]	['Familial hypertrophic cardiomyopathy 1', 'Cardiomyopathy', 'not specified']
NM_000256.3(MYBPC3):c.2906-2A>G	727504333	MYBPC3	[]	['CCGTTTCYGTGGG TATAGAGTGGG', 'GCCGTTTCYGTGG GTATAGAGTGG']	['Familial hypertrophic cardiomyopathy 4']
NM_001128425.1(MUTYH):c.1187-2A>G	587781628	MUTYH	['ACCYGAG AGGGAGG GCAGCCAG G']	['ACCYGAGAGGG AGGGCAGCCAGG']	['Hereditary cancer- predisposing syndrome', 'Carcinoma of colon']
NM_005188.3(CBL):c .1228-2A>G	727504426	CBL	[]	[]	['Juvenile myelomonocytic leukemia', 'Noonan syndrome-like disorder with or without juvenile myelomonocytic leukemia', 'Rasopathy']
NM_000501.3(ELN):c .890-2A>G	727504434	ELN	['GCCYGAA AACACAGC CACAGAGG']	['GCCYGAAAACA CAGCCACAGAGG']	['Supravalvar aortic stenosis']
NM_001165963.1(SCN1A):c.2877T>A (p.Cys959Ter)	775214722	SCN1A	[]	[]	['not provided']
NM_000833.4(GRIN2A):c.2449A>G (p.Met817Val)	796052551	GRIN2A	['CCAYGTT GTCAATGT CCAGCTGG']	['CCAYGTTGTCAA TGTCCAGCTGG']	['not provided']
NM_000314.6(PTEN): c.493-2A>G	587781784	PTEN	[]	[]	['Hereditary cancer- predisposing syndrome']

NM_000498.3(CYP11B2):c.1157T>C (p.Val386Ala)	61757294	-	[]	[]	['Corticosterone methyl oxidase type 2 deficiency', 'Corticosterone methyl oxidase type 1 deficiency']
NM_006204.3(PDE6C):c.1483-2A>G	786200910	PDE6C	[]	['CTTTCYGTGAAATAAGGATGGG', 'TCTTTCYGTGAAATAAGGATGGG']	['Achromatopsia 5']
NM_003588.3(CUL4B):c.901-2A>G	786200913	CUL4B	[]	[]	['Syndromic X-linked mental retardation, Cabezas type']
NM_000397.3(CYBB):c.302A>G (p.His101Arg)	137854591	CYBB	[]	[]	['Granulomatous disease, chronic, X-linked, variant', 'not provided']
NM_000311.3(PRNP):c.385A>G (p.Met129Val)	1799990	PRNP	[]	[]	['Jakob-Creutzfeldt disease', 'Genetic prion diseases', 'Fatal familial insomnia', 'not specified']
NM_000051.3(ATM):c.3994-2A>G	587782276	ATM	[]	[]	['Ataxia-telangiectasia syndrome', 'Hereditary cancer-predisposing syndrome']
NM_005211.3(CSF1R):c.1754-2A>G	281860267	CSF1R	[]	[]	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_004646.3(NPHS1):c.1756A>G (p.Arg586Gly)	730880174	NPHS1	[]	[]	['Finnish congenital nephrotic syndrome']
NM_005211.3(CSF1R):c.2320-2A>G	281860272	CSF1R	['CACYGAGGGAAAGC ACTGCAGG']	['CACYGAGGGAAAGCACTGCAGG', 'GCACYGAGGGAAAGCACTGCAGG']	['Hereditary diffuse leukoencephalopathy with spheroids']
NM_000256.3(MYBP3):c.3392T>C (p.Ile1131Thr)	370890951	MYBPC3	[]	[]	['Cardiomyopathy', 'Cardiac arrest', 'not specified']
NM_000551.3(VHL):c.586A>T (p.Lys196Ter)	281860296	VHL	[]	['GGTCTTYCTGCACATTTGGGTGG']	['Von Hippel-Lindau syndrome']
NM_005247.2(FGF3):c.146A>G (p.Tyr49Cys)	281860300	FGF3	['CAGYAGAGCTTGCGGCGCCGGG', 'GCAGYAGAGCTTGCGGCGCCGGG', 'CGCAGYAGAGCTTGCGGCGCCGG']	['CAGYAGAGCTTGCGGCGCCGGG', 'GCAGYAGAGCTTGCGGCGCCGG']	['Deafness with labyrinthine aplasia microtia and microdontia (LAMM)']
NM_005247.2(FGF3):c.317A>G	281860306	FGF3	[]	[]	['Deafness with labyrinthine aplasia']

(p.Tyr106Cys)					microtia and microdontia (LAMM)']
NM_000314.6(PTEN):c.403A>G (p.Ile135Val)	587782360	PTEN	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_004990.3(MARS):c.1031A>G (p.Tyr344Cys)	766466297	MARS	[]	[]	['Pulmonary alveolar proteinosis', 'Interstitial lung and liver disease']
NM_006343.2(MERTK):c.1605-2A>G	730880273	MERTK	[]	[]	['Retinitis pigmentosa 38']
NM_003611.2(OFD1):c.935+706A>G	730880283	OFD1	[]	[]	['Retinitis Pigmentosa 23']
NM_004793.3(LONP1):c.2353A>G (p.Arg785Gly)	730880293	LONP1	[]	[]	['CODAS syndrome']
NM_001698.2(AUH):c.263-2A>G	730880311	AUH	[]	[]	['3-Methylglutaconic aciduria']
NM_001698.2(AUH):c.943-2A>G	730880312	AUH	[]	[]	['3-Methylglutaconic aciduria']
NM_000169.2(GLA):c.370-2A>G	730880444	-	[]	['GTGAACCYGAAATGAGAGGGAGG']	['not provided']
NM_001110792.1(MECP2):c.520A>G (p.Arg174Gly)	727505391	MECP2	[]	[]	['Rett disorder']
NM_030662.3(MAP2K2):c.181A>G (p.Lys61Glu)	730880517	MAP2K2	[]	[]	['Cardiofaciocutaneous syndrome', 'Rasopathy']
NM_000256.3(MYBP C3):c.1227-2A>G	730880531	MYBPC3	[]	['GTACCYGGGTG GGGGCCGCAGGG', 'TGTACCYGGGTG GGGGCCGCAGG']	['Familial hypertrophic cardiomyopathy 4', 'Cardiomyopathy']
NM_000642.2(AGL):c.4260-12A>G	369973784	AGL	[]	[]	['Glycogen storage disease type III', 'Glycogen storage disease IIIa', 'Glycogen storage disease IIIb']
NM_000267.3(NF1):c.1642-8A>G	267606602	NF1	[]	[]	['Neurofibromatosis, type 1', 'Juvenile myelomonocytic leukemia']
NM_000267.3(NF1):c.5944-5A>G	267606604	NF1	[]	[]	['Neurofibromatosis, type 1', 'Neurofibromatosis, familial spinal']
m.1555A>G	267606617	MT-RNR1	[]	[]	['Aminoglycoside-induced deafness', 'Cardiomyopathy, restrictive', 'Deafness, nonsyndromic sensorineural, mitochondrial']
NM_022458.3(LMBR1):c.423+5252A>G	606231150	LMBR1	[]	[]	['Triphalangeal thumb', 'Preaxial

					polydactyly 2']
NM_000642.2(AGL):c.3439A>G (p.Arg1147Gly)	267606639	AGL	[]	[]	['Glycogen storage disease IIIc']
NM_013411.4(AK2):c.494A>G (p.Asp165Gly)	267606643	AK2	[]	['TCAYCTTTCATG GGCTCTTTTGG']	['Reticular dysgenesis']
NM_001142800.1(EYS):c.9209T>C (p.Ile3070Thr)	183589498	EYS	[]	[]	['Retinitis pigmentosa']
NM_004183.3(BEST1):c.680A>G (p.Tyr227Cys)	267606677	BEST1	[]	[]	['Vitelliform dystrophy', 'Retinitis pigmentosa, concentric', 'not provided']
NM_005188.3(CBL):c.1144A>G (p.Lys382Glu)	267606705	CBL	[]	['TATTTYACATAG TTGGAATGTGG']	['Noonan syndrome-like disorder with or without juvenile myelomonocytic leukemia']
NM_001017361.2(KHDC3L):c.1A>G (p.Met1Val)	606231235	KHDC3L	[]	[]	['Hydatidiform mole, recurrent, 2']
NM_144577.3(CCDC114):c.487-2A>G	606231239	CCDC114	[]	[]	['Ciliary dyskinesia, primary, 20']
NM_000277.1(PAH):c.916A>G (p.Ile306Val)	62642934	PAH	[]	['GGCCAAYTTCCT GTAATTGGGGG', 'AGGCCAAyttcc TGTAATTGGGG']	['Phenylketonuria', 'Hyperphenylalaninemia, non-pku', 'not provided']
NM_000257.3(MYH7):c.2792A>G (p.Glu931Gly)	730880760	MYH7	[]	[]	['Cardiomyopathy']
NM_207034.2(EDN3):c.335A>G (p.His112Arg)	267606778	EDN3	[]	[]	['Waardenburg syndrome type 4B']
NM_000117.2(EMD):c.1A>G (p.Met1Val)	267606782	EMD	[]	['TCCAYGGCGGGT GCGGGCTCAGG']	['Emery-Dreifuss muscular dystrophy, X-linked']
NM_003937.2(KYNU):c.592A>G (p.Thr198Ala)	606231307	KYNU	[]	[]	['Hydroxykynureninuria']
NM_004387.3(NKX2-5):c.461A>G (p.Glu154Gly)	587782928	NKX2-5	[]	[]	['Atrial septal defect 7 with or without atrioventricular conduction defects']
NM_000142.4(FGFR3):c.1454A>G (p.Gln485Arg)	267606808	FGFR3	[]	[]	['Thanatophoric dysplasia type 1']
NM_014053.3(FLVCR1):c.361A>G (p.Asn121Asp)	267606820	FLVCR1	[]	['AGGCGTYGACC AGCGAGTACAGG']	['Posterior column ataxia with retinitis pigmentosa']
NM_000257.3(MYH7):c.4664A>G (p.Glu1555Gly)	730880805	-	[]	['GCCCYCCTCGTG CTCCAGGGAGG', 'CTTGCCCYCCTC GTGCTCCAGGG']	['Cardiomyopathy']
NM_138387.3(G6PC3):c.346A>G (p.Met116Val)	267606834	G6PC3	[]	['TGATCAYGCAGT GTCCAGAAGGG', 'GTGATCAYGCAG TGTCCAGAAGG']	['Dursun syndrome']

NM_020347.3(LZTFL1):c.260T>C (p.Leu87Pro)	515726135	LZTFL1	[]	[]	['Bardet-Biedl syndrome', 'Bardet-Biedl syndrome 17']
NM_000175.3(GPI):c.1028A>G (p.Gln343Arg)	267606851	GPI	[]	['GTACYGGTCATAGGCAGCATGG']	['Hemolytic anemia, nonspherocytic, due to glucose phosphate isomerase deficiency']
NM_005859.4(PURA):c.289A>G (p.Lys97Glu)	587782994	PURA	[]	[]	['Neonatal hypotonia', 'Intellectual disability', 'Seizures', 'Delayed speech and language development', 'Global developmental delay', 'Mental retardation, autosomal dominant 31']
NM_005144.4(HR):c.-218A>G	267606869	HR	['CTCYAGG GCCGCAGG TTGGAGGG']	['CTCYAGGGCCGCAGGTTGGAGGG', 'GCTCYAGGGCCGCAGGTTGGAGG', 'GGCGCTCYAGGGCCGCAGGTTGG']	['Marie Unna hereditary hypotrichosis 1']
NM_000257.3(MYH7):c.789A>G (p.Ile263Met)	730880855	MYH7	[]	[]	['Cardiomyopathy']
NM_000060.3(BTD):c.683A>G (p.Asp228Gly)	587783004	BTD	[]	[]	['Biotinidase deficiency']
NM_000257.3(MYH7):c.1051A>G (p.Lys351Glu)	730880864	MYH7	[]	[]	['Cardiomyopathy']
NM_015713.4(RRM2B):c.190T>C (p.Trp64Arg)	515726182	RRM2B	[]	['TTCCTTCYGGACAGCAGAAGAGG']	['RRM2B-related mitochondrial disease']
NM_005957.4(MTHFR):c.971A>G (p.Asn324Ser)	267606887	MTHFR	['CGCGGYTGAGGGTGTAGAAGTGG']	['CGCGGYTGAGGGTGTAGAAGTGG']	['Homocystinuria due to MTHFR deficiency']
NM_015713.4(RRM2B):c.368T>C (p.Phe123Ser)	515726187	RRM2B	[]	[]	['RRM2B-related mitochondrial disease']
m.12770A>G	267606894	MT-ND5	[]	[]	['Juvenile myopathy, encephalopathy, lactic acidosis AND stroke']
NM_000257.3(MYH7):c.1805A>G (p.Asn602Ser)	730880880	MYH7	[]	[]	['Cardiomyopathy']
NM_018109.3(MTPAP):c.1432A>G (p.Asn478Asp)	267606900	MTPAP	['AATGGATYCTGAATGTACAGAGG']	['AATGGATYCTGAATGTACAGAGG']	['Ataxia, spastic, 4, autosomal recessive']
NM_000257.3(MYH7):c.2717A>G (p.Asp906Gly)	267606908	MYH7	[]	[]	['Primary familial hypertrophic cardiomyopathy']

					'Familial hypertrophic cardiomyopathy 1', 'Cardiomyopathy']
NM_003122.4(SPINK1):c.160T>C (p.Tyr54His)	515726207	SPINK1	[]	[]	['Hereditary pancreatitis']
NM_003159.2(CDKL5):c.404-2A>G	587783080	CDKL5	[]	[]	['not provided']
NM_003159.2(CDKL5):c.449A>G (p.Lys150Arg)	587783083	CDKL5	['ACAGTYT TAGGACAT CATTGTGG']	['ACAGTYTTAGGA CATCATTGTGG']	['not provided']
NM_016203.3(PRKAG2):c.1589A>G (p.His530Arg)	267606977	PRKAG2	[]	[]	['Familial hypertrophic cardiomyopathy 6', 'not provided']
NM_198965.1(PTHLH):c.534A>G (p.Ter178Trp)	267606987	PTHLH	[]	[]	['Brachydactyly type E2']
NM_000531.5(OTC):c.122A>G (p.Asp41Gly)	74518351	OTC	[]	[]	['not provided']
NM_001134363.2(RBM20):c.1909A>G (p.Ser637Gly)	267607005	RBM20	[]	[]	['Dilated cardiomyopathy 1DD']
NM_000553.4(WRN):c.403A>G (p.Lys135Glu)	267607008	WRN	[]	[]	['Werner syndrome']
NM_002880.3(RAF1):c.1279A>G (p.Ser427Gly)	730881002	RAF1	[]	['GCTGCGGCCCTC GCACCACTGGG', 'GGCTGCGGCCCT CGCACCACTGG']	['Rasopathy']
NM_002977.3(SCN9A):c.29A>G (p.Gln10Arg)	267607030	SCN9A	[]	['AAGCTCYGAGG TCCTGGGGGAGG']	['Primary erythromelalgia']
NM_016955.3(SEPSECS):c.1001A>G (p.Tyr334Cys)	267607036	SEPSECS	[]	[]	['Pontocerebellar hypoplasia type 2D']
NM_007373.3(SHOC2):c.4A>G (p.Ser2Gly)	267607048	SHOC2	[]	['TACYCATGGTGA CTCAAGCCTGG']	['Noonan-like syndrome with loose anagen hair', 'Rasopathy']
NM_005633.3(SOS1):c.1430A>G (p.Gln477Arg)	730881044	SOS1	[]	[]	['Rasopathy']
NM_007375.3(TARDBP):c.787A>G (p.Lys263Glu)	267607102	TARDBP	[]	[]	['FRONTOTEMPORAL DEMENTIA WITH TDP43 INCLUSIONS, TARDBP-RELATED']
NM_003286.2(TOP1):c.1598A>G (p.Asp533Gly)	267607131	-	[]	[]	[]
NM_021625.4(TRPV4):c.1805A>G (p.Tyr602Cys)	267607150	TRPV4	[]	[]	['Spondyloepiphyseal dysplasia Maroteaux type']
NM_000551.3(VHL):c	267607170	VHL	[]	[]	['Von Hippel-Lindau

.491A>G (p.Gln164Arg)					syndrome']
NM_001006657.1(WDR35):c.1877A>G (p.Glu626Gly)	267607174	WDR35	[]	[]	['Craniocutaneous dysplasia 2']
NM_024884.2(L2HGDH):c.293A>G (p.His98Arg)	267607206	L2HGDH	[]	[]	['L-2-hydroxyglutaric aciduria']
NM_002437.4(MPV17):c.262A>G (p.Lys88Glu)	267607256	MPV17	[]	[]	['Navajo neurohepatopathy']
NM_006888.4(CALM1):c.293A>G (p.Asn98Ser)	267607277	CALM1	[]	[]	['Catecholaminergic polymorphic ventricular tachycardia', 'Ventricular tachycardia, catecholaminergic polymorphic, 4']
NM_000487.5(ARSA):c.*96A>G	6151429	ARSA	[]	[]	['Metachromatic leukodystrophy', 'Arylsulfatase A pseudodeficiency', 'not provided']
NM_003122.4(SPINK1):c.194+2T>C	148954387	SPINK1	[]	[]	['Hereditary pancreatitis']
NM_000552.3(VWF):c.3437A>G (p.Tyr1146Cys)	267607326	VWF	[]	[]	['von Willebrand disease type 2', 'not provided']
NM_000489.4(ATRX):c.4826A>G (p.His1609Arg)	122445093	ATRX	[]	[]	['ATR-X syndrome']
NM_000489.4(ATRX):c.6488A>G (p.Tyr2163Cys)	122445098	ATRX	[]	[]	['ATR-X syndrome']
NM_000489.4(ATRX):c.6811A>G (p.Arg2271Gly)	122445112	ATRX	[]	[]	[]
NM_004380.2(CREBBP):c.3983-2A>G	587783486	CREBBP	[]	['GCAGCCCYAGG AAGTCCAGAAGG']	['Rubinstein-Taybi syndrome']
NM_004380.2(CREBBP):c.4508A>G (p.Tyr1503Cys)	587783497	CREBBP	[]	[]	['Rubinstein-Taybi syndrome']
NM_000051.3(ATM):c.3154-2A>G	730881357	ATM	[]	['AGCCYACGGGA AAAGA ACTGTGG']	['Hereditary cancer-predisposing syndrome']
NM_178151.2(DCX):c.1027-2A>G	587783518	DCX	[]	[]	['Heterotopia']
NM_178151.2(DCX):c.520A>G (p.Lys174Glu)	587783557	DCX	[]	[]	['Heterotopia']
NM_178151.2(DCX):c.538A>G (p.Lys180Glu)	587783560	DCX	[]	[]	['Heterotopia']
NM_178151.2(DCX):c.607A>G (p.Thr203Ala)	587783570	DCX	[]	[]	['Heterotopia']
NM_001257235.1(ALG13)	398122394	ALG13	[]	[]	['Congenital disorder

G13):c.8A>G (p.Asn3Ser)					of glycosylation type 1s']
NM_001256864.1(DNAJC6):c.801-2A>G	398122404	DNAJC6	[]	['AGGTATCYGAAACAGAAGGTTGG']	['Parkinson disease 19, juvenile-onset']
NM_001927.3(DES):c.1024A>G (p.Asn342Asp)	267607482	DES	[]	['GAATCGTYCTGCAGGAGAGGGGG']	['Myofibrillar myopathy 1', 'not provided']
NM_001927.3(DES):c.735+3A>G	267607483	DES	[]	[]	['Myofibrillar myopathy 1', 'not provided']
NM_001927.3(DES):c.1333A>G (p.Thr445Ala)	267607498	DES	[]	[]	['not provided']
NM_005249.4(FOXG1):c.757A>G (p.Asn253Asp)	587783641	FOXG1	[]	[]	['Rett syndrome, congenital variant']
NM_000252.2(MTM1):c.1406A>G (p.His469Arg)	587783789	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000391.3(TPP1):c.833A>G (p.Gln278Arg)	796053439	TPP1	[]	['CAGGTACYGCA CATCTAGACTGG']	['not provided']
NM_000252.2(MTM1):c.301A>G (p.Ser101Gly)	587783818	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1):c.343-2A>G	587783821	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1):c.529-2A>G	587783831	MTM1	[]	[]	['Severe X-linked myotubular myopathy']
NM_000252.2(MTM1):c.550A>G (p.Arg184Gly)	587783835	MTM1	[]	['GTTATTCYCCAA TGGTGATTGGG']	['Severe X-linked myotubular myopathy']
NM_000158.3(GBE1):c.691+2T>C	192044702	GBE1	[]	[]	['Glycogen storage disease, type IV']
NM_000252.2(MTM1):c.629A>G (p.Asp210Gly)	587783842	MTM1	[]	['TCATCAYCTGAG GCACGATACGG']	['Severe X-linked myotubular myopathy']
NM_000249.3(MLH1):c.545+3A>G	267607760	MLH1	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_000249.3(MLH1):c.589-2A>G	267607767	MLH1	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms', 'Hereditary cancer-predisposing syndrome']
NM_000249.3(MLH1):c.884+4A>G	267607777	MLH1	[]	['TGCTACAYTACC TGAGGTACAGG']	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_000081.3(LYST):c.10127A>G (p.Asn3376Ser)	80338669	LYST	[]	[]	['Ch\ xc3\ xa9diak-Higashi syndrome']
NM_000528.3(MAN2)	80338678	MAN2B1	[]	[]	['Deficiency of

B1):c.1831-2A>G					alpha-mannosidase']
NM_022132.4(MCCC2):c.569A>G (p.His190Arg)	119103225	MCCC2	[]	[]	['3-methylcrotonyl CoA carboxylase 2 deficiency']
m.3260A>G	199474663	MT-TL1	['TTAAGTT YTATGCGA TTACCGGG']	['TTAAGTTYTATG CGATTACCGGG']	['Cardiomyopathy with or without skeletal myopathy']
NM_014874.3(MFN2):c.827A>G (p.Gln276Arg)	119103264	MFN2	[]	[]	['Hereditary motor and sensory neuropathy with optic atrophy']
NM_004525.2(LRP2):c.770-2A>G	80338743	LRP2	[]	[]	['Donnai Barrow syndrome']
NM_005120.2(MED12):c.3020A>G (p.Asn1007Ser)	80338759	MED12	[]	[]	['X-linked mental retardation with marfanoid habitus syndrome']
NM_000834.3(GRIN2B):c.2172-2A>G	398122824	GRIN2B	[]	[]	['Mental retardation, autosomal dominant 6']
NM_000249.3(MLH1):c.1990-2A>G	267607883	MLH1	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms', 'not provided']
NM_000518.4(HBB):c.59A>G (p.Asn20Ser)	33972047	HBB	[]	['CACGYTCACCTT GCCCCACAGGG', 'CCACGYTCACCT TGCCCCACAGG']	['alpha Thalassemia']
NM_003688.3(CASK):c.2168A>G (p.Tyr723Cys)	398122844	CASK	[]	[]	['FG syndrome 4', 'Mental retardation and microcephaly with pontine and cerebellar hypoplasia']
NM_024675.3(PALB2):c.109-2A>G	730881897	PALB2	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000251.2(MSH2):c.1511-2A>G	267607962	MSH2	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms']
NM_003124.4(SCRN1):c.596-2A>G	398122922	SCRN1	[]	[]	['Sepiapterin reductase deficiency']
NM_022455.4(NSD1):c.4498-3A>G	587784120	NSD1	[]	[]	['Sotos syndrome 1']
NM_000455.4(STK11):c.889A>G (p.Arg297Gly)	730881978	STK11	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_001927.3(DES):c.1289-2A>G	398122940	DES	[]	[]	['Muscular dystrophy, limb-girdle, type 2r']
NM_000546.5(TP53):c.709A>G (p.Met237Val)	730882004	TP53	[]	['ACACAYGTAGTT GTAGTGGATGG']	['Li-Fraumeni syndrome', 'Hereditary cancer-predisposing syndrome']

NM_024876.3(ADCK4):c.857A>G (p.Asp286Gly)	398122979	ADCK4	[]	[]	['Nephrotic syndrome, type 9']
NM_022455.4(NSD1):c.6059A>G (p.Asn2020Ser)	587784178	NSD1	[]	[]	['Sotos syndrome 1']
NM_022455.4(NSD1):c.6356A>G (p.Asp2119Gly)	587784191	NSD1	[]	[]	['Sotos syndrome 1']
NM_007332.2(TRPA1):c.2564A>G (p.Asn855Ser)	398123010	-	[]	[]	['Familial episodic pain syndrome 1']
NM_001231.4(CASQ1):c.731A>G (p.Asp244Gly)	730882052	CASQ1	[]	['GGCTTGYCTGGG ATGGTCACAGG']	['Myopathy, vacuolar, with casq1 aggregates']
NM_004004.5(GJB2):c.487A>G (p.Met163Val)	80338949	GJB2	[]	[]	['Deafness, autosomal recessive 1A', 'not specified']
NM_130466.3(UBE3B):c.545-2A>G	398123022	UBE3B	[]	[]	['Kaufman oculocerebrofacial syndrome']
NM_000334.4(SCN4A):c.4078A>G (p.Met1360Val)	80338959	SCN4A	[]	['GATCAYGATGGT GATGTCGAAGG']	['Hyperkalemic Periodic Paralysis Type 1']
NM_000334.4(SCN4A):c.4108A>G (p.Met1370Val)	80338960	SCN4A	[]	['CCATCAYGGTGA CCATGTTGAGG']	['Hyperkalemic Periodic Paralysis Type 1']
NM_000334.4(SCN4A):c.4774A>G (p.Met1592Val)	80338962	SCN4A	[]	['TGTACAYGTTGA CCACGATGAGG']	['Hyperkalemic Periodic Paralysis Type 1', 'Familial hyperkalemic periodic paralysis']
NM_015250.3(BICD2):c.2321A>G (p.Glu774Gly)	398123030	BICD2	[]	[]	['Spinal muscular atrophy, lower extremity predominant 2, autosomal dominant']
NM_006012.2(CLPP):c.270+4A>G	398123035	CLPP	[]	[]	['Autosomal recessive hearing impairment with normal menstrual cycles']
NM_000179.2(MSH6):c.3439-2A>G	267608098	MSH6	[]	[]	['Hereditary Nonpolyposis Colorectal Neoplasms', 'Hereditary cancer-predisposing syndrome', 'not provided']
NM_002246.2(KCNK3):c.575A>G (p.Tyr192Cys)	398123043	KCNK3	[]	[]	['Primary pulmonary hypertension 4']
NM_001070.4(TUBG1):c.275A>G (p.Tyr92Cys)	398123046	TUBG1	[]	[]	['Cortical dysplasia, complex, with other brain malformations 4']
NM_000383.3(AIRE):	179363882	AIRE	[]	[]	['Polyglandular

c.254A>G (p.Tyr85Cys)					autoimmune syndrome, type 1', 'not provided']
NM_001651.3(AQP5): c.367A>G (p.Asn123Asp)	398123057	AQP5	[]	[]	['Diffuse palmoplantar keratoderma, Bothnian type']
NM_012160.4(FBXL4): c.1694A>G (p.Asp565Gly)	398123062	FBXL4	[]	['TATGYCCAGCTGCTGTAACCTGG']	['Mitochondrial DNA depletion syndrome 13 (encephalomyopathic type)']
NM_024531.4(SLC52A2): c.914A>G (p.Tyr305Cys)	398123068	SLC52A2	[]	[]	['Brown-Vialetto-Van Laere syndrome 2']
NM_001039550.1(DNAJB2): c.14A>G (p.Tyr5Cys)	730882140	DNAJB2	[]	['GATCTCGYAGTAGGATGCCATGG']	['Charcot-Marie-Tooth disease', 'Charcot-Marie-Tooth disease, axonal, type 2T']
NM_052859.3(RFT1): c.1222A>G (p.Met408Val)	796053522	RFT1	[]	['GCAYCACAAAA TTGTACCTGGG', 'AGCAYCACAAAA TTGTACCTGGG', 'CAGCAYCACAAAA ATTGTACCTGG']	['Congenital disorder of glycosylation type 1N']
NM_007294.3(BRCA1): c.5057A>G (p.His1686Arg)	730882166	BRCA1	[]	[]	['Breast-ovarian cancer, familial 1']
NM_000050.4(ASS1): c.496-2A>G	398123130	ASS1	[]	[]	['Citrullinemia type I']
NM_012233.2(RAB3GAP1): c.649-2A>G	730882183	RAB3GAP1	[]	[]	['Warburg micro syndrome 1']
NM_000159.3(GCDH): c.542A>G (p.Glu181Gly)	398123194	GCDH	[]	[]	['Glutaric aciduria, type 1', 'not provided']
NM_000288.3(PEX7): c.340-10A>G	267608255	PEX7	[]	[]	['Phytanic acid storage disease', 'Peroxisome biogenesis disorder 9B']
NM_000038.5(APC): c.221-2A>G	786201291	APC	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_000169.2(GLA): c.509A>G (p.Asp170Gly)	398123210	-	[]	[]	['Fabry disease']
NM_000169.2(GLA): c.548-2A>G	398123211	-	[]	['AACCYGTATGAGAAAACAATGGG', 'TAACCYGTATGAGAAAACAATGG']	['Fabry disease']
NM_006888.4(CALM1): c.389A>G (p.Asp130Gly)	730882252	CALM1	[]	[]	['Long QT syndrome 14']
NM_000169.2(GLA): c.647A>G (p.Tyr216Cys)	398123217	-	[]	[]	['Fabry disease']
NM_006306.3(SMC1)	587784423	SMC1A	[]	['AGCCYGTGCAA	['Congenital

A):c.616-2A>G				ACAGGGGAATGG'	muscular hypertrophy-cerebral syndrome']
NM_000255.3(MUT):c.1445-2A>G	398123276	MUT	[]	[]	['Methylmalonic aciduria due to methylmalonyl-CoA mutase deficiency', 'not provided']
NM_001083962.1(TCF4):c.991-2A>G	587784470	TCF4	[]	[]	['Pitt-Hopkins syndrome']
NM_000060.3(BTD):c.1205A>G (p.Asn402Ser)	201023772	BTD	[]	[]	['Biotinidase deficiency']
NC_000007.14:g.62535490A>G	483352872	-	[]	[]	['Isolated growth hormone deficiency type 1B']
NM_000271.4(NPC1):c.1832A>G (p.Asp611Gly)	483352887	NPC1	[]	[]	['Niemann-Pick disease type C1']
NM_152419.2(HGSNAT):c.372-2A>G	483352896	HGSNAT	[]	[]	['Mucopolysaccharidosis, MPS-III-C']
NM_001199397.1(NEK1):c.869-2A>G	483352906	NEK1	[]	[]	['Short rib-polydactyly syndrome, Majewski type']
NM_000350.2(ABCA4):c.67-2A>G	398123339	ABCA4	[]	[]	['Stargardt disease 1']
NM_004992.3(MECP2):c.27-2A>G	267608412	MECP2	[]	[]	['Rett disorder', 'not provided']
NM_003159.2(CDKL5):c.100-2A>G	267608423	CDKL5	[]	[]	['Early infantile epileptic encephalopathy 2', 'not provided']
NM_003159.2(CDKL5):c.125A>G (p.Lys42Arg)	267608429	CDKL5	[]	[]	['Early infantile epileptic encephalopathy 2', 'not provided']
NM_000487.5(ARSA):c.1108-2A>G	398123411	ARSA	[]	['GGCTCYGGGGG CAGAGTCAGGGG', 'GGGCTCYGGGGG CAGAGTCAGGG', 'AGGGCTCYGGGG GCAGAGTCAGG']	['Metachromatic leukodystrophy']
NM_003159.2(CDKL5):c.380A>G (p.His127Arg)	267608468	CDKL5	[]	[]	['Atypical Rett syndrome', 'not provided']
NM_000489.4(ATRX):c.134-2A>G	398123420	ATRX	[]	[]	['not provided']
NM_003159.2(CDKL5):c.464-2A>G	267608480	CDKL5	[]	[]	['Early infantile epileptic encephalopathy 2', 'not provided']
NM_000489.4(ATRX):c.536A>G (p.Asn179Ser)	398123425	ATRX	[]	[]	['not provided']
NM_000512.4(GALNS):c.1171A>G (p.Met391Val)	398123429	GALNS	[]	['CCGCCAYCAGC GTGTCGCCACGG']	['Mucopolysaccharidosis, MPS-IV-A', 'not provided']

NM_003159.2(CDKL5):c.578A>G (p.Asp193Gly)	267608500	CDKL5	[]	['ATGYCCACGGA CTTTCCATAGGG', 'CATGYCCACGGA CTTTCCATAGG']	['Early infantile epileptic encephalopathy 2']
NM_000521.3(HEXB):c.1243-2A>G	398123446	HEXB	[]	[]	['Sandhoff disease', 'not provided']
NM_003159.2(CDKL5):c.978-2A>G	267608553	CDKL5	[]	[]	['Early infantile epileptic encephalopathy 2', 'not provided']
NM_001164342.2(ZBTB20):c.1787A>G (p.His596Arg)	483353066	ZBTB20	[]	[]	['Primrose syndrome']
NM_000402.4(G6PD):c.188T>C (p.Ile63Thr)	398123552	-	[]	['ACACACAYATTC ATCATCATGGG']	['Anemia, nonspherocytic hemolytic, due to G6PD deficiency', 'not provided']
NM_001893.4(CSNK1D):c.130A>G (p.Thr44Ala)	104894561	CSNK1D	[]	[]	['Advanced sleep phase syndrome, familial, 2']
NM_000155.3(GALT):c.563A>G (p.Gln188Arg)	75391579	GALT	[]	['TTACCYGGCAGT GGGGGTGGGG', 'CTTACCYGGCAG TGGGGGTGGGG', 'CCTTACCYGGCA GTGGGGGTGGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase', 'not provided']
NM_007055.3(POLR3A):c.2554A>G (p.Met852Val)	267608671	POLR3A	[]	[]	['Hypomyelinating leukodystrophy 7']
NM_001848.2(COL6A1):c.805-2A>G	398123639	COL6A1	[]	['TTCTCCCYGGAA CACAAAACAGG']	['Ullrich congenital muscular dystrophy', 'Bethlem myopathy', 'not provided']
NM_001918.3(DBT):c.773-2A>G	398123674	DBT	[]	[]	['Maple syrup urine disease', 'not provided']
NM_001999.3(FBN2):c.3740T>C (p.Met1247Thr)	149054177	FBN2	['GAATGTA YGATAATG AACGGAG G']	['GAATGTAYGAT AATGAACGGAGG']	['not specified', 'Macular degeneration, early-onset']
NM_003482.3(KMT2D):c.5645-2A>G	398123750	KMT2D	[]	['GCAGTTCYGTGG GGAATGAAGG']	['Kabuki make-up syndrome', 'not provided']
NM_003494.3(DYSF):c.1398-2A>G	398123769	DYSF	[]	[]	['Limb-girdle muscular dystrophy, type 2B', 'not provided']
NM_015560.2(OPA1):c.1146A>G (p.Ile382Met)	143319805	OPA1	[]	[]	['Dominant hereditary optic atrophy', 'Optic Atrophy Type 1', 'not specified', 'not provided']
NM_203447.3(DOCK8):c.1418A>G (p.Lys473Arg)	112321280	DOCK8	[]	[]	['Hyperimmunoglobulin E recurrent infection syndrome, autosomal recessive']

NM_001145.4(ANG):c.121A>G (p.Lys41Glu)	121909537	-	['TGGTTYG GCATCATA GTGCTGGG', 'GTGGTTYG GCATCATA GTGCTGG']	['TGGTTYGGCATC ATAGTGCTGGG', 'GTGGTTYGGCAT CATAGTGCTGG']	['Amyotrophic lateral sclerosis type 9']
NM_004006.2(DMD):c.3432+3A>G	398123938	DMD	[]	[]	['Dilated cardiomyopathy 3B']
NM_006514.3(SCN10A):c.1661T>C (p.Leu554Pro)	138404783	SCN10A	[]	[]	['Episodic pain syndrome, familial, 2']
NM_004006.2(DMD):c.6763-2A>G	398124033	DMD	[]	[]	['Dilated cardiomyopathy 3B']
NM_001079802.1(FKTN):c.1112A>G (p.Tyr371Cys)	119464998	FKTN	[]	[]	[]
NM_004006.2(DMD):c.9224+61934A>G	398124084	DMD	[]	[]	['Dilated cardiomyopathy 3B']
NM_004006.2(DMD):c.9225-647A>G	398124091	DMD	[]	[]	['Duchenne muscular dystrophy', 'Becker muscular dystrophy', 'Dilated cardiomyopathy 3B']
NM_004006.2(DMD):c.9650-2A>G	398124100	DMD	[]	[]	['Duchenne muscular dystrophy', 'Becker muscular dystrophy', 'Dilated cardiomyopathy 3B']
NM_198578.3(LRRK2):c.3342A>G (p.Leu1114=)	35808389	LRRK2	[]	[]	['Parkinson disease 8, autosomal dominant']
NM_031229.2(RBCK1):c.1160A>G (p.Asn387Ser)	566912235	RBCK1	[]	[]	['Polyglucosan body myopathy 1 with or without immunodeficiency']
NM_000178.2(GSS):c.656A>G (p.Asp219Gly)	28938472	GSS	[]	[]	['Glutathione synthetase deficiency of erythrocytes, hemolytic anemia due to']
NM_025132.3(WDR19):c.407-2A>G	374400438	WDR19	[]	[]	['SENIOR-LOKEN SYNDROME 8']
NM_144997.5(FLCN):c.1433-2A>G	398124528	FLCN	[]	['CCCACYGGGGA GAAGGGCAGGGG', 'GCCCACYGGGGA GAAGGGCAGGGG', 'GGCCCACYGGGG AGAAGGGCAGG']	['Hereditary cancer-predisposing syndrome', 'not provided']
NM_144997.5(FLCN):c.250-2A>G	398124533	FLCN	[]	[]	['not provided']
NM_000146.3(FTL):c.-160A>G	398124633	FTL	[]	[]	['Hyperferritinemia cataract syndrome']
NM_003184.3(TAF2):c.1945T>C (p.Trp649Arg)	398124645	TAF2	[]	[]	['Mental retardation, autosomal recessive 40']
NM_013281.3(FLRT3)	398124654	-	[]	[]	['Hypogonadotropic

)c.1016A>G (p.Lys339Arg)					hypogonadism 21 with or without anosmia']
NM_002834.3(PTPN11):c.767A>G (p.Gln256Arg)	397507523	PTPN11	[]	[]	['Noonan syndrome 1', 'Rasopathy', 'not provided']
NM_000054.4(AVPR2):c.614A>G (p.Tyr205Cys)	104894749	AVPR2	['ACAYAGG TGCGACGG CCCCAGGG', 'GACAYAG GTGCGACG GCCCCAGG']	['ACAYAGGTGCG ACGGCCCCAGGG', 'GACAYAGGTGCG ACGGCCCCAGG']	['Nephrogenic diabetes insipidus', 'Nephrogenic diabetes insipidus, X-linked']
NM_031157.2(HNRNPA1):c.956A>G (p.Asn319Ser)	397518454	HNRNPA1	[]	[]	['Amyotrophic lateral sclerosis 20']
NM_000277.1(PAH):c.662A>G (p.Glu221Gly)	62514934	PAH	[]	[]	['Phenylketonuria', 'not provided']
NM_000219.5(KCNE1):c.176T>C (p.Leu59Pro)	141813529	KCNE1	[]	[]	['Congenital long QT syndrome']
NM_001165963.1(SCN1A):c.1076A>G (p.Asn359Ser)	794726713	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_013339.3(ALG6):c.391T>C (p.Tyr131His)	35383149	ALG6	[]	[]	['Congenital disorder of glycosylation type 1C', 'not specified']
NM_153767.3(KCNJ1):c.1013T>C (p.Met338Thr)	59172778	KCNJ1	[]	[]	['Bartter syndrome antenatal type 2']
NM_176824.2(BBS7):c.968A>G (p.His323Arg)	119466001	BBS7	[]	[]	['Bardet-Biedl syndrome 7']
NM_000199.3(SGSH):c.892T>C (p.Ser298Pro)	138504221	SGSH	[]	[]	['Mucopolysaccharidosis, MPS-III-A', 'not provided']
NM_000891.2(KCNJ2):c.220A>G (p.Thr74Ala)	199473652	KCNJ2			['Congenital long QT syndrome']
NM_001165963.1(SCN1A):c.1048A>G (p.Met350Val)	794726768	SCN1A	['ACAYATA TCCCTCTG GACATTGG']	['ACAYATATCCCT CTGGACATTGG']	['Severe myoclonic epilepsy in infancy']
NM_002863.4(PYGL):c.1016A>G (p.Asn339Ser)	113993976	PYGL	[]	[]	['Glycogen storage disease, type VI']
NM_001165963.1(SCN1A):c.2537A>G (p.Glu846Gly)	794726794	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_002693.2(POLG):c.2864A>G (p.Tyr955Cys)	113994099	POLG	[]	[]	['Autosomal dominant progressive external ophthalmoplegia with mitochondrial DNA deletions 1']
NM_000920.3(PC):c.1705A>G	113994144	PC	[]	[]	['Pyruvate carboxylase

(p.Thr569Ala)					deficiency']
NM_025265.3(TSEN2):c.926A>G (p.Tyr309Cys)	113994149	TSEN2	[]	['CAGAGCAYAGA CCAAGAAAAGG']	['Pontocerebellar hypoplasia type 2B']
NM_001039958.1(MESP2):c.271A>G (p.Lys91Glu)	113994156	MESP2	[]	[]	['Spondylocostal dysostosis 2']
NM_024649.4(BBS1):c.1340-2A>G	113994180	-	[]	[]	['Bardet-Biedl syndrome']
NM_033028.4(BBS4):c.157-2A>G	113994192	BBS4	[]	[]	['Bardet-Biedl syndrome', 'Bardet-Biedl syndrome 4']
NM_212472.2(PRKAR1A):c.1A>G (p.Met1Val)	281864779	PRKAR1A	[]	[]	['Carney complex, type 1']
NM_212472.2(PRKAR1A):c.178-2A>G	281864796	PRKAR1A	[]	[]	['Carney complex, type 1']
NM_212472.2(PRKAR1A):c.891+3A>G	281864799	PRKAR1A	[]	[]	['Carney complex, type 1']
NM_001165963.1(SCN1A):c.433A>G (p.Met145Val)	794726849	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy']
NM_005710.2(PQBP1):c.194A>G (p.Tyr65Cys)	121917899	PQBP1	[]	[]	['Renpenning syndrome 1']
NM_000921.4(PDE3A):c.1333A>G (p.Thr445Ala)	794726865	PDE3A	['CGAGGYG GTGGTGGT CCAAGTGG']	['CGAGGYGGTGG TGGTCCAAGTGG']	['Brachydactyly with hypertension']
NM_024312.4(GNPTAB):c.1285-2A>G	281864974	GNPTAB	[]	[]	['Pseudo-Hurler polydystrophy']
NM_024312.4(GNPTAB):c.2783A>G (p.Lys928Arg)	281865003	GNPTAB	[]	[]	['I cell disease']
NM_024312.4(GNPTAB):c.2867A>G (p.His956Arg)	281865005	GNPTAB	[]	[]	['Pseudo-Hurler polydystrophy']
NM_024312.4(GNPTAB):c.3053A>G (p.Asp1018Gly)	281865007	GNPTAB	[]	[]	['I cell disease']
NM_024312.4(GNPTAB):c.3458A>G (p.Asn1153Ser)	281865019	GNPTAB	[]	[]	['Pseudo-Hurler polydystrophy']
NM_024312.4(GNPTAB):c.118-2A>G	281865023	GNPTAB	[]	[]	['I cell disease']
NM_198578.3(LRRK2):c.5605A>G (p.Met1869Val)	281865052	LRRK2	[]	['TCAACAYAATAT TTCTAGGCAGG']	['Parkinson disease 8, autosomal dominant']
NM_139241.3(FGD4):c.1762-2A>G	281865065	FGD4	[]	[]	['Charcot-Marie-Tooth disease, type 4H']
NM_006121.3(KRT1):c.1445A>G (p.Tyr482Cys)	58420087	KRT1	[]	[]	['Bullous ichthyosiform erythroderma', 'not provided']
NM_000195.4(HPS1):c.716T>C (p.Leu239Pro)	281865080	HPS1	[]	[]	['Hermansky-Pudlak syndrome 1']
NM_000195.4(HPS1):	281865090	HPS1	[]	[]	['Hermansky-Pudlak

c.2003T>C (p.Leu668Pro)					syndrome 1']
NM_022081.5(HPS4): c.461A>G (p.His154Arg)	281865098	HPS4	[]	[]	['Hermansky-Pudlak syndrome 4']
NM_000277.1(PAH):c .1157A>G (p.Tyr386Cys)	62516141	PAH	[]	[]	['Phenylketonuria', 'not provided']
NM_025114.3(CEP29 0):c.2991+1655A>G	281865192	CEP290	['GAGATAY TCACAATT ACAACCTGG ']	['GATAYTCACAAT TACAACCTGGG', 'AGATAYTCACAA TTACAACCTGGG', 'GAGATAYTCACA ATTACAACCTGG']	['Leber congenital amaurosis 10', 'not provided']
NM_018319.3(TDP1): c.1478A>G (p.His493Arg)	119467003	TDP1	[]	[]	['Spinocerebellar ataxia autosomal recessive with axonal neuropathy']
NM_000051.3(ATM): c.5762_5763insNG_00 9830.1:g.91138_91274	774925473	ATM	[]	[]	['Ataxia- telangiectasia variant']
NM_004614.4(TK2):c. 562A>G (p.Thr188Ala)	281865495	TK2	[]	['AAGYCTCAGGA TTGGTCCGAAGG']	['Mitochondrial DNA depletion syndrome 2']
NM_003494.3(DYSF): c.3041A>G (p.Tyr1014Cys)	756328339	DYSF	[]	['CTAYACTCCCAG CCTGGGGGAGG', 'ATGCTAYACTCC CAGCCTGGGGG', 'GATGCTAYACTC CCAGCCTGGGG']	['Limb-girdle muscular dystrophy, type 2B']
NM_000531.5(OTC):c .1034A>G (p.Tyr345Cys)	72558492	OTC	['AGGTGAG YAATCTGT CAGCAGGG ']	['AGGTGAGYAAT CTGTCAGCAGGG']	['not provided']
NM_000518.4(HBB):c .199A>G (p.Lys67Glu)	34165323	HBB	[]	[]	['Hemoglobinopathy' ']
NM_153427.2(PITX2) :c.262A>G (p.Lys88Glu)	387906810	PITX2	[]	['TCTYGAACCAAA CCTGGGGGCGG', 'GATTCTYGAACC AAACCTGGGGG', 'CGATTCTYGAAC CAAACCTGGGG']	['Axenfeld-Rieger syndrome type 1']
NM_030964.3(SPRY4 (p.Lys177Arg)	78310959	SPRY4	[]	['AGTGTCYTGTCCA GCTCGGGTGGG', 'AAGTGTCYTGTCC AGCTCGGGTGG']	['Hypogonadotropic hypogonadism 17 with or without anosmia']
NM_002834.3(PTPN1 1):c.922A>G (p.Asn308Asp)	28933386	PTPN11	[]	[]	['Noonan syndrome', 'Noonan syndrome 1', 'Rasopathy', 'not provided']
NM_000518.4(HBB):c .-50-29A>G	34598529	HBB	[]	[]	['alpha Thalassemia', 'Beta thalassemia intermedia']
NM_207352.3(CYP4V 2):c.1393A>G (p.Arg465Gly)	144109267	CYP4V2	[]	['TTCCYGGGGCCA GCAGAGAAGGG', 'GTTCYGGGGCC AGCAGAGAAGG']	['Bietti crystalline corneoretinal dystrophy']
NM_001360.2(DHCR 7):c.1A>G	104886033	DHCR7	[]	[]	['Smith-Lemli-Opitz syndrome']

(p.Met1Val)					
NM_000495.4(COL4A5):c.1A>G (p.Met1Val)	104886050	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.2692A>G (p.Met898Val)	104886192	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.2746A>G (p.Ser916Gly)	104886193	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_004572.3(PKP2):c.2062T>C (p.Ser688Pro)	144601090	PKP2	[]	[]	['Arrhythmogenic right ventricular cardiomyopathy', 'not specified', 'not provided']
NM_000495.4(COL4A5):c.4790A>G (p.Tyr1597Cys)	104886298	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.1340-2A>G	104886319	COL4A5	[]	['CACCYGAGTAA GATAAAGAAAGG']	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.2042-18A>G	104886341	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.2147-2A>G	104886344	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.2510-33A>G	104886358	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000495.4(COL4A5):c.3107-2A>G	104886379	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000051.3(ATM):c.7268A>G (p.Glu2423Gly)	121434221	ATM	[]	[]	['Mantle cell lymphoma']
NM_000495.4(COL4A5):c.3605-2A>G	104886385	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_000096.3(CP):c.1209-2A>G	386134137	CP	[]	[]	['Deficiency of ferroxidase']
NM_000348.3(SRD5A2):c.692A>G (p.His231Arg)	121434251	SRD5A2	[]	[]	['3-Oxo-5 alpha-steroid delta 4-dehydrogenase deficiency']
NM_002739.3(PRKCG):c.76A>G (p.Arg26Gly)	386134157	PRKCG	[]	[]	['Spinocerebellar ataxia 14']
NM_000383.3(AIRE):c.247A>G (p.Lys83Glu)	121434255	AIRE	[]	[]	['AUTOIMMUNE POLYENDOCRINOPATHY SYNDROME, TYPE I']
NM_000495.4(COL4A5):c.466-2A>G	104886416	COL4A5	[]	['ACCCYAAAAGA AGCCATCAATGG']	['Alport syndrome, X-linked recessive']
NM_001127328.2(ACADM):c.589A>G (p.Thr197Ala)	121434279	ACADM	[]	[]	['Medium-chain acyl-coenzyme A dehydrogenase deficiency']
NM_000495.4(COL4A5):c.892-2A>G	104886453	COL4A5	[]	[]	['Alport syndrome, X-linked recessive']
NM_020533.2(MCOLN1):c.406-2A>G	104886461	MCOLN1	['TACYGTGGCAGAG AAGGGGA	['TACYGTGGGCA GAGAAGGGGAGG	['Ganglioside sialidase deficiency', 'not provided']

			GG']	'AGGTACYGTGGG CAGAGAAGGGG', 'CAGGTACYGTGG GCAGAGAAGGG']	
NM_018136.4(ASPM) :c.2761-25A>G	199422149	ASPM	[]	[]	['Primary autosomal recessive microcephaly 5']
NM_017780.3(CHD7): c.3082A>G (p.Ile1028Val)	121434338	CHD7	[]	[]	['CHARGE association', 'not provided']
NM_017780.3(CHD7): c.164A>G (p.His55Arg)	121434345	CHD7	[]	[]	['Kallmann syndrome 5']
NM_152783.4(D2HGDH): c.1315A>G (p.Asn439Asp)	121434362	D2HGDH	['GCAGGTY ACCATCTC CTGGAGGG , 'TGCAGGT YACCATCT CCTGGAGG '	['GCAGGTYACCAT CTCCTGGAGGG', 'TGCAGGTYACCA TCTCCTGGAGG']	['D-2- hydroxyglutaric aciduria 1']
NM_005006.6(NDUFS1): c.755A>G (p.Asp252Gly)	199422224	NDUFS1	[]	[]	['Mitochondrial complex I deficiency', 'not provided']
NM_002894.2(RBBP8) :c.1009A>G (p.Lys337Glu)	121434388	RBBP8	[]	[]	['Carcinoma of pancreas']
NM_004621.5(TRPC6) :c.428A>G (p.Asn143Ser)	121434391	TRPC6	[]	[]	['Focal segmental glomerulosclerosis 2']
NM_003705.4(SLC25A12): c.1769A>G (p.Gln590Arg)	121434396	SLC25A12	[]	[]	['Hypomyelination, global cerebral']
NM_001363.4(DKC1): c.127A>G (p.Lys43Glu)	199422243	DKC1	[]	[]	['Dyskeratosis congenita X-linked']
NM_001084.4(PLOD3) :c.668A>G (p.Asn223Ser)	121434414	PLOD3	[]	[]	['Bone fragility with contractures, arterial rupture, and deafness']
NM_006702.4(PNPLA6): c.3034A>G (p.Met1012Val)	121434415	PNPLA6	[]	[]	['Spastic paraplegia 39']
NR_001566.1(TERC): n.48A>G	199422262	TERC	[]	[]	['Dyskeratosis congenita autosomal dominant']
NM_004984.2(KIF5A) :c.767A>G (p.Asn256Ser)	121434441	KIF5A	[]	[]	['Spastic paraplegia 10']
NM_004984.2(KIF5A) :c.827A>G (p.Tyr276Cys)	121434443	KIF5A	[]	['GAACAYAGCTTT TCTGGGGGAGG']	['Spastic paraplegia 10']
m.10438A>G	121434456	MT-TR	[]	[]	['Mitochondrial encephalomyopathy' '
NM_198253.2(TERT): c.2537A>G (p.Tyr846Cys)	199422302	TERT	[]	[]	['Aplastic anemia']

m.12320A>G	121434463	MT-TL2	['TGGAGTY GCACCAAA ATTTTTGG']	['GAGTYGCACCA AAATTTTTGGGG', 'GGAGTYGCACCA AAATTTTTGGG', 'TGGAGTYGCACC AAAATTTTTGG']	['Mitochondrial myopathy']
m.4317A>G	121434465	MT-TI	[]	[]	[]
m.4269A>G	121434466	MT-TI	['ACAYATT TCTTAGGT TTGAGGGG' , 'GACAYAT TTCTTAGG TTTGAGGG']	['ACAYATTTCTTA GGTTTGAGGGG', 'GACAYATTTCTT AGGTTTGAGGG', 'AGACAYATTTCT TAGGTTTGAGG']	[]
m.4295A>G	121434467	MT-TI	[]	[]	['Primary familial hypertrophic cardiomyopathy', 'Deafness, nonsyndromic sensorineural, mitochondrial']
m.4300A>G	121434470	MT-TI	[]	[]	['Primary familial hypertrophic cardiomyopathy']
NM_001099274.1(TINF2):c.850A>G (p.Thr284Ala)	199422314	TINF2	[]	['TGACTGYGGGG CGCTCCTTATGG']	['Dyskeratosis congenita autosomal dominant']
NM_004044.6(ATIC): c.1277A>G (p.Lys426Arg)	121434478	ATIC	[]	['AGTGTACYTGAC AGCAATGGTGG']	['AICAR transformylase/IMP cyclohydrolase deficiency']
NM_001099274.1(TINF2):c.871A>G (p.Arg291Gly)	199422319	TINF2	[]	[]	['Dyskeratosis congenita autosomal dominant']
NM_015474.3(SAMHD1):c.760A>G (p.Met254Val)	121434521	SAMHD1	[]	[]	['Aicardi Goutieres syndrome 5']
NM_001103.3(ACTN2):c.26A>G (p.Gln9Arg)	121434525	ACTN2	[]	[]	['Dilated cardiomyopathy 1AA', 'Cardiomyopathy', 'Dilated cardiomyopathy', 'not specified']
NM_000155.3(GALT):c.812A>G (p.Glu271Gly)	111033765	GALT	[]	['CGCYCAGCAGG GGTCAGCTCAGG']	['Deficiency of UDPglucose- hexose-1-phosphate uridylyltransferase', 'not provided']
NM_000316.2(PTH1R):c.668A>G (p.His223Arg)	121434597	PTH1R	[]	[]	['Metaphyseal chondrodysplasia, Jansen type']
NM_006006.4(ZBTB16):c.1849A>G (p.Met617Val)	121434606	ZBTB16	[]	['GATCAYGGCCG AGTAGTCCCGG', 'TGATCAYGGCCG AGTAGTCCCGG']	['Skeletal defects, genital hypoplasia, and mental retardation']
NM_014305.3(TGDS): c.700T>C	544436734	TGDS	[]	[]	['Catel Manzke syndrome']

(p.Tyr234His)					
NM_002835.3(PTPN12):c.182A>G (p.Lys61Arg)	121434623	PTPN12	[]	[]	['Carcinoma of colon']
NM_000035.3(ALDOB):c.1027T>C (p.Tyr343His)	369586696	ALDOB	[]	[]	['Hereditary fructosuria']
NM_006180.4(NTRK2):c.2165A>G (p.Tyr722Cys)	121434633	NTRK2	[]	[]	['Obesity, hyperphagia, and developmental delay']
NM_000107.2(DDB2):c.730A>G (p.Lys244Glu)	121434639	DDB2	[]	[]	['Xeroderma pigmentosum, group E']
NM_000017.3(ACADS):c.1108A>G (p.Met370Val)	566325901	ACADS	[]	['AGCCCAYGCCG CCCAGGATCTGG']	['not provided']
NM_001151.3(SLC25A4):c.311A>G (p.Asp104Gly)	28999114	SLC25A4	[]	[]	['Autosomal dominant progressive external ophthalmoplegia with mitochondrial DNA deletions 2']
NM_012079.5(DGAT1):c.751+2T>C	148665132	DGAT1	[]	['ACCGCGGYGAG GACCTCTGTGGG']	['Diarrhea 7']
NM_002036.3(ACKR1):c.-67T>C	2814778	ACKR1	[]	[]	['White blood cell count quantitative trait locus 1']
NM_000492.3(CFTR):c.1666A>G (p.Ile556Val)	75789129	CFTR	[]	[]	['Cystic fibrosis', 'not specified']
NM_000155.3(GALT):c.574A>G (p.Ser192Gly)	111033830	GALT	[]	['TGCYGGCCCAT CCTGTCAAGG', 'CTGCYGGCCAT ACCTGTCAAGG']	['Deficiency of UDPglucose-hexose-1-phosphate uridylyltransferase']
NM_174916.2(UBR1):c.407A>G (p.His136Arg)	119477054	UBR1	[]	[]	['Johanson-Blizzard syndrome']
m.3274A>G	199474666	MT-TL1	[]	[]	[]
NM_000060.3(BTD):c.128A>G (p.His43Arg)	146011150	BTD	[]	[]	['Biotinidase deficiency']
NM_172107.2(KCNQ2):c.635A>G (p.Asp212Gly)	118192202	KCNQ2	[]	[]	['Benign familial neonatal seizures 1']
NM_006493.2(CLN5):c.1121A>G (p.Tyr374Cys)	148862100	CLN5	[]	[]	['Ceroid lipofuscinosis neuronal 5']
NM_000060.3(BTD):c.880A>G (p.Ile294Val)	35976361	BTD	[]	[]	['Biotinidase deficiency']
NM_000132.3(F8):c.1226A>G (p.Glu409Gly)	28933671	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.5600A>G (p.His1867Arg)	28933679	F8	[]	['GAGYGCACATCT TTTTCCTAGGG', 'TGAGYGCACATC TTTTCCTAGG']	['Hereditary factor VIII deficiency disease']
NM_000266.3(NDP):c.1A>G (p.Met1Val)	28933685	NDP	[]	[]	['Atrophia bulborum hereditaria']

NM_000133.3(F9):c.2 78A>G (p.Asp93Gly)	137852230	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_000133.3(F9):c.3 29A>G (p.Asp110Gly)	137852234	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_000133.3(F9):c.9 17A>G (p.Asn306Ser)	137852251	F9	[]	['GCTGCAYTGTAG TTGTGGTGAGG']	['Hereditary factor IX deficiency disease']
NM_000133.3(F9):c.1 180A>G (p.Met394Val)	137852262	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_000133.3(F9):c.1 231A>G (p.Ser411Gly)	137852277	F9	[]	[]	['Hereditary factor IX deficiency disease']
NM_000292.2(PHKA2) :c.896A>G (p.Asp299Gly)	137852289	PHKA2	[]	[]	['Glycogen storage disease type IXa1']
NM_000292.2(PHKA2) :c.565A>G (p.Lys189Glu)	137852295	PHKA2	[]	[]	['Glycogen storage disease IXa2']
NM_000032.4(ALAS2) :c.1702A>G (p.Ser568Gly)	137852306	ALAS2	[]	[]	['Hereditary sideroblastic anemia']
NM_001287223.1(SCN11A):c.3473T>C (p.Leu1158Pro)	141686175	SCN11A	[]	['CGTGCGCYGTCC CAGTTTGAAGG']	['Episodic pain syndrome, familial, 3']
NM_000402.4(G6PD): c.583A>G (p.Asn195Asp)	137852331	G6PD	[]	['ATGCGGTYCCAG CCTCTGCTGGG']	['Favism, susceptibility to', 'Anemia, nonspherocytic hemolytic, due to G6PD deficiency']
NM_000132.3(F8):c.8 72A>G (p.Glu291Gly)	137852359	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_018718.2(CEP41) :c.107T>C (p.Met36Thr)	368178632	CEP41	[]	[]	['Joubert syndrome 9/15, digenic']
NM_000132.3(F8):c.5 183A>G (p.Tyr1728Cys)	137852362	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.5 821A>G (p.Asn1941Asp)	137852369	F8	[]	['TAGCCATYGATT GCTGGAGAAGG']	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.3 28A>G (p.Met110Val)	137852385	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.3 98A>G (p.Tyr133Cys)	137852389	F8	[]	['TCAYATTCAGCT CCTATAGCAGG']	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.4 04A>G (p.Asp135Gly)	137852390	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.9 40A>G (p.Thr314Ala)	137852406	F8	[]	['TGAGCAGYAAG GAAAGTTATTGG']	['Hereditary factor VIII deficiency disease']
NM_000041.3(APOE): c.178A>G	28931576	APOE	[]	['ACAGTGYCTGCA CCCAGCGCAGG']	[]

(p.Thr60Ala)					
NM_000132.3(F8):c.1 682A>G (p.Asp561Gly)	137852420	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.1 892A>G (p.Asn631Ser)	137852429	F8	['ATGYTGG AGGCTTGG AACTCTGG']	['ATGYTGGAGGCT TGGAACTCTGG']	['Hereditary factor VIII deficiency disease']
NM_012082.3(ZFPM2)):c.1969A>G (p.Ser657Gly)	28374544	-	[]	[]	['Tetralogy of Fallot']
NM_000098.2(CPT2): c.638A>G (p.Asp213Gly)	74315300	CPT2	[]	[]	['CARNITINE PALMITOYLTRANSFERASE II DEFICIENCY, LATE-ONSET']
NM_000396.3(CTSK): c.990A>G (p.Ter330Trp)	74315301	CTSK	[]	['GAGYCACATCTT GGGAAGCTGG']	['Pyknodysostosis']
NM_000132.3(F8):c.6 113A>G (p.Asn2038Ser)	137852454	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000132.3(F8):c.6 278A>G (p.Asp2093Gly)	137852457	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_024009.2(GJB3): c.421A>G (p.Ile141Val)	74315320	GJB3	['CAAYGAT GAGCTTGA AGATGAGG']	['CAAYGATGAGC TTGAAGATGAGG']	['Deafness, autosomal recessive']
NM_000132.3(F8):c.1 04A>G (p.Tyr35Cys)	137852476	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_000194.2(HPRT1)):c.602A>G (p.Asp201Gly)	137852479	HPRT1	[]	[]	['Partial hypoxanthine-guanine phosphoribosyltransferase deficiency']
NM_000261.1(MYOC)):c.1267A>G (p.Lys423Glu)	74315336	MYOC	[]	[]	['Primary open angle glaucoma juvenile onset 1']
NM_014625.3(NPHS2)):c.479A>G (p.Asp160Gly)	74315346	NPHS2	[]	[]	['Nephrotic syndrome, idiopathic, steroid-resistant']
NM_000194.2(HPRT1)):c.155A>G (p.Asp52Gly)	137852502	HPRT1	[]	[]	['Partial hypoxanthine-guanine phosphoribosyltransferase deficiency']
NM_002764.3(PRPS1) :c.341A>G (p.Asn114Ser)	137852540	PRPS1	[]	['TAGCATAYTTGC AACAAGCTTGG']	['Phosphoribosylpyrophosphate synthetase superactivity']
NM_000055.2(BCHE) :c.293A>G (p.Asp98Gly)	1799807	BCHE	[]	[]	['Postanesthetic apnea']
NM_170784.2(MKKS) :c.169A>G (p.Thr57Ala)	74315399	MKKS	[]	[]	['Bardet-Biedl syndrome 6']

NM_000311.3(PRNP): c.650A>G (p.Gln217Arg)	74315406	PRNP	[]	[]	['Gerstmann- Straussler-Scheinker syndrome', 'Genetic prion diseases']
NM_000311.3(PRNP): c.560A>G (p.His187Arg)	74315413	PRNP	[]	[]	['Gerstmann- Straussler-Scheinker syndrome', 'Genetic prion diseases', 'Spongiform encephalopathy with neuropsychiatric features']
NM_000044.3(AR):c.2 291A>G (p.Tyr764Cys)	137852567	AR	[]	[]	['Reifenstein syndrome']
NM_000044.3(AR):c.2 362A>G (p.Met788Val)	137852570	AR	[]	[]	[]
NM_000044.3(AR):c.2 632A>G (p.Thr878Ala)	137852578	AR	[]	[]	['Malignant tumor of prostate']
NM_020436.3(SALL4 (p.His888Arg)	74315429	SALL4	[]	[]	['Duane-radial ray syndrome']
NM_000044.3(AR):c.2 708A>G (p.Gln903Arg)	137852582	AR	[]	[]	['Malignant tumor of prostate']
NM_000211.4(ITGB2) :c.1052A>G (p.Asn351Ser)	137852613	ITGB2	[]	[]	['Leukocyte adhesion deficiency']
NM_000215.3(JAK3): c.299A>G (p.Tyr100Cys)	137852624	JAK3	[]	['AATCCTGYACAG CAGGACTTGGG']	['Severe combined immunodeficiency, autosomal recessive, T cell-negative, B cell-positive, NK cell-negative']
NM_001166107.1(HM GCS2):c.500A>G (p.Tyr167Cys)	137852640	HMGCS2	[]	['ACCACCGYAGC AGGCATTGGTGG']	['mitochondrial 3- hydroxy-3- methylglutaryl-CoA synthase deficiency']
NM_002047.2(GARS) :c.374A>G (p.Glu125Gly)	137852645	GARS	[]	[]	['Charcot-Marie- Tooth disease type 2D', 'Distal hereditary motor neuropathy type 5']
NM_033163.3(FGF8): c.298A>G (p.Lys100Glu)	137852662	FGF8	[]	[]	['Kallmann syndrome 6']
NM_002180.2(IGHM BP2):c.638A>G (p.His213Arg)	137852666	IGHMBP 2	[]	[]	['Werdnig-Hoffmann disease']
NM_004387.3(NKX2- 5):c.896A>G (p.Asp299Gly)	137852683	NKX2-5	[]	[]	['Atrial septal defect 7 with or without atrioventricular conduction defects']
NM_004387.3(NKX2- 5):c.547A>G (p.Lys183Glu)	137852686	NKX2-5	[]	[]	[]

NM_000310.3(PPT1): c.236A>G (p.Asp79Gly)	137852697	PPT1	[]	[]	['Ceroid lipofuscinosis neuronal 1']
NM_000336.2(SCNN1B):c.863A>G (p.Asn288Ser)	137852712	SCNN1B	[]	[]	['Bronchiectasis']
NM_000579.3(CCR5): c.-301+246A>G	1799987	-	[]	[]	['Human immunodeficiency virus type 1, susceptibility to']
NM_001204.6(BMPR2):c.1454A>G (p.Asp485Gly)	137852745	BMPR2	[]	[]	['Primary pulmonary hypertension']
NM_005591.3(MRE11A):c.350A>G (p.Asn117Ser)	137852760	MRE11A	[]	[]	['Hereditary cancer-predisposing syndrome', 'Ataxia-telangiectasia-like disorder']
NM_003476.4(CSRP3):c.206A>G (p.Lys69Arg)	137852764	CSRP3	[]	[]	['Dilated cardiomyopathy 1M', 'Cardiomyopathy', 'Familial hypertrophic cardiomyopathy 12']
NM_000519.3(HBD):c.-81A>G	35518301	HBD	[]	[]	[]
NM_005633.3(SOS1): c.1654A>G (p.Arg552Gly)	137852814	SOS1	[]	['GCATCCYTTCCA GTGTACTCCG']	['Noonan syndrome', 'Noonan syndrome 4', 'Rasopathy', 'not provided']
NM_003688.3(CASK):c.2129A>G (p.Asp710Gly)	137852818	CASK	[]	[]	['FG syndrome 4']
NM_031443.3(CCM2):c.1A>G (p.Met1Val)	137852842	CCM2	[]	[]	['Cerebral cavernous malformations 2']
NM_182760.3(SUMF1):c.1A>G (p.Met1Val)	137852855	SUMF1	[]	[]	['Multiple sulfatase deficiency']
NM_001171993.1(HPD):c.362A>G (p.Tyr121Cys)	137852865	HPD	[]	['CCTCAYATCCAG GCAAGAATTGG']	['4-Hydroxyphenylpyruvate dioxygenase deficiency']
NM_024996.5(GFM1):c.521A>G (p.Asn174Ser)	119470018	GFM1	['TTGYTAA TAAAAGTT AGAAACG G']	['TTGYTAATAAAA GTTAGAAACGG']	['Combined oxidative phosphorylation deficiency 1']
NM_000158.3(GBE1): c.1634A>G (p.His545Arg)	137852889	GBE1	[]	[]	['Glycogen storage disease, type IV', 'GLYCOGEN STORAGE DISEASE IV, FATAL PERINATAL NEUROMUSCULAR']
NM_000158.3(GBE1): c.1883A>G (p.His628Arg)	137852891	GBE1	[]	[]	['Glycogen storage disease, type IV', 'GLYCOGEN STORAGE DISEASE IV,

					CHILDHOOD NEUROMUSCULAR']
m.8344A>G	118192098	MT-TK	[]	[]	['Parkinson disease, mitochondrial', 'Leigh disease', 'Myoclonus with epilepsy with ragged red fibers']
NM_000540.2(RYR1): c.10100A>G (p.Lys3367Arg)	118192126	RYR1	[]	[]	['Central core disease', 'not provided']
NM_000540.2(RYR1): c.14572A>G (p.Asn4858Asp)	118192144	RYR1	[]	[]	['Central core disease', 'not provided']
NM_012464.4(TLL1): c.1885A>G (p.Ile629Val)	137852953	TLL1	['GGTTAYG GTGCCGTT AAGTTTGG']	['GGTTAYGGTGCC GTTAAGTTTGG']	['Atrial septal defect 6']
NM_025243.3(SLC19A3):c.130A>G (p.Lys44Glu)	137852957	SLC19A3	[]	[]	['Basal ganglia disease, biotin-responsive']
NM_138691.2(TMC1):c.1763+3A>G	370898981	TMC1	[]	['TGGCCYACCAG ATCATGCCTTGG']	['Deafness, autosomal recessive 7']
NM_000540.2(RYR1): c.13909A>G (p.Thr4637Ala)	118192166	RYR1	[]	[]	['Central core disease', 'not provided']
NM_000540.2(RYR1): c.14387A>G (p.Tyr4796Cys)	118192167	RYR1	[]	['CCATAYACCAGC CCAGGTACAGG']	['Malignant hyperthermia susceptibility type 1', 'Central core disease', 'not provided']
NM_032667.6(BSCL2):c.263A>G (p.Asn88Ser)	137852972	-	[]	['CGAGACAYTGG CAACAGGGAAGG']	['Distal hereditary motor neuronopathy type 5', 'Silver spastic paraplegia syndrome', 'Charcot-Marie-Tooth disease, type 2']
NM_014795.3(ZEB2): c.3356A>G (p.Gln1119Arg)	137852983	ZEB2	[]	[]	['Mowat-Wilson syndrome']
NM_000540.2(RYR1): c.14740A>G (p.Arg4914Gly)	118192184	RYR1	[]	[]	['Central core disease', 'not provided']
NM_172107.2(KCNQ2):c.356A>G (p.Glu119Gly)	118192193	KCNQ2	[]	['CTTCYCATACTC CTTGATGGTGG', 'GCTCTTCYCATA CTCCTTGATGG']	['Benign familial neonatal seizures 1']
NM_172107.2(KCNQ2):c.622A>G (p.Met208Val)	118192201	KCNQ2	[]	['GGATCAYCCGC AGAATCTGCAGG']	['Benign familial neonatal seizures 1']
NM_172107.2(KCNQ2):c.773A>G (p.Asn258Ser)	118192207	KCNQ2	[]	[]	['Benign familial neonatal seizures 1']
NM_004006.2(DMD): c.8734A>G	1800278	DMD	[]	[]	['Duchenne muscular dystrophy',

(p.Asn2912Asp)						'not specified']
NM_004006.2(DMD): c.8762A>G (p.His2921Arg)	1800279	DMD	[]	[]		['Becker muscular dystrophy', 'not specified']
NM_001080463.1(DY NC2H1):c.11284A>G (p.Met3762Val)	137853026	DYNC2H 1	[]	[]		['Short-rib thoracic dysplasia 3 with or without polydactyly']
NM_001080463.1(DY NC2H1):c.9044A>G (p.Asp3015Gly)	137853027	DYNC2H 1	[]	['ATAYCTCTAATT ACATCAGGTGG', 'AGAATAYCTCTA ATTACATCAGG']		['Short-rib thoracic dysplasia 3 with or without polydactyly']
NM_001080463.1(DY NC2H1):c.4610A>G (p.Gln1537Arg)	137853033	DYNC2H 1	['ACCYGTG AAGGGAA CAGAGATG G']	['ACCYGTGAAGG GAACAGAGATGG']		['Short-rib thoracic dysplasia 3 with or without polydactyly']
NM_001080463.1(DY NC2H1):c.5959A>G (p.Thr1987Ala)	137853035	DYNC2H 1	[]	[]		['Short-rib thoracic dysplasia 3 with or without polydactyly']
NM_001430.4(EPAS1 (p.Met535Val)	137853037	EPAS1	[]	[]		['Erythrocytosis, familial, 4']
NM_172107.2(KCNQ 2):c.1764-2A>G	118192238	KCNQ2	[]	[]		['Benign familial neonatal seizures 1']
NM_004519.3(KCNQ 3):c.914A>G (p.Asp305Gly)	118192248	KCNQ3	[]	[]		['Benign familial neonatal seizures 2']
NM_004519.3(KCNQ 3):c.1403A>G (p.Asn468Ser)	118192252	KCNQ3	['TCTTTAY TGTTTAAAG CCAACAGG']	['TCTTTAYTGTTT AAGCCAACAGG']		['Benign familial neonatal seizures 2', 'not specified', 'not provided']
NM_004519.3(KCNQ 3):c.2462A>G (p.Asn821Ser)	118192254	KCNQ3	[]	[]		['Benign familial neonatal seizures 2', 'not specified', 'not provided']
NM_138701.3(MPLKI P):c.430A>G (p.Met144Val)	137853117	MPLKIP	[]	[]		['Trichothiodystrophy, nonphotosensitive 1']
NM_005094.3(SLC27 A4):c.899A>G (p.Gln300Arg)	137853134	SLC27A4	[]	[]		['Ichthyosis prematurity syndrome']
NM_194456.1(KRIT1 (p.Asp137Gly)	137853139	KRIT1	[]	[]		['Cerebral cavernous malformations 1']
NM_000351.4(STS):c. 1331A>G (p.His444Arg)	137853169	STS	[]	[]		['X-linked ichthyosis with steryl-sulfatase deficiency']
NM_152416.3(NDUF AF6):c.296A>G (p.Gln99Arg)	137853184	NDUFAF 6	[]	[]		['Leigh syndrome due to mitochondrial complex I deficiency']
NM_144573.3(NEXN (p.Tyr652Cys)	137853197	NEXN	[]	['ATAYACTCTCCT CCATCTTCTGG']		['Dilated cardiomyopathy 1CC', 'Cardiomyopathy', 'not specified']
NM_000476.2(AK1):c. 491A>G	137853203	AK1	[]	['TTCTCAYAGAAG GCGATGACGGG']		['Adenylate kinase deficiency,

(p.Tyr164Cys)				'TTTCTCAYAGAA GGCGATGACGG']	hemolytic anemia due to']
NM_013411.4(AK2):c .1A>G (p.Met1Val)	137853206	AK2	[]	[]	['Reticular dysgenesis']
NM_000308.2(CTSA): c.746+3A>G	786200859	CTSA	[]	['TCCCAAYACCTGT TCCCAGAAGG']	['Galactosialidosis, adult']
NM_002890.2(RASA1 (p.Lys400Glu)	137853215	RASA1	[]	[]	[]
NM_002890.2(RASA1 (p.Ile401Val)	137853216	RASA1	[]	[]	[]
NM_000515.4(GH1):c .413A>G (p.Asp138Gly)	137853221	GH1	[]	[]	['Kowarski syndrome']
NM_017636.3(TRPM4 (p.Lys914Arg)	172151858	TRPM4	[]	[]	['Progressive familial heart block type 1B']
NM_000545.6(HNF1A (p.Tyr122Cys)	137853237	HNF1A	[]	[]	['Maturity-onset diabetes of the young, type 3']
NM_003494.3(DYSF): c.1285-2A>G	786200897	DYSF	[]	['CAGCYAGAAGA CACAGGGAGGGG' , 'ACAGCYAGAAGA CACAGGGAGGG', 'CACAGCYAGAAG ACACAGGGAGG']	['Limb-girdle muscular dystrophy, type 2B']
NM_005055.4(RAPSN <td>786200905</td> <td>RAPSN</td> <td>[]</td> <td>[]</td> <td>['MYASTHENIC SYNDROME, CONGENITAL, 11, ASSOCIATED WITH ACETYLCHOLINE RECEPTOR DEFICIENCY']</br></td>	786200905	RAPSN	[]	[]	['MYASTHENIC SYNDROME, CONGENITAL, 11, ASSOCIATED WITH
NM_000276.3(OCRL) :c.1436A>G (p.Tyr479Cys)	137853262	OCRL	[]	[]	['Dent disease 2']
NM_004463.2(FGD1): c.1396A>G (p.Met466Val)	137853267	FGD1	[]	[]	['Aarskog syndrome']
NM_153252.4(BRWD 3):c.4786A>G (p.Lys1596Glu)	137853272	BRWD3	[]	[]	['Mental retardation, X-linked 93']
NM_206933.2(USH2A <td>786200928</td> <td>USH2A</td> <td>[]</td> <td>['CTCTTAYCTTGG GAAAGGAGAGG']</td> <td>['Usher syndrome, type 2A']</td>	786200928	USH2A	[]	['CTCTTAYCTTGG GAAAGGAGAGG']	['Usher syndrome, type 2A']
NM_000362.4(TIMP3) :c.572A>G (p.Tyr191Cys)	137853299	-	['TGCAGYA GCCGCCCT TCTGCCGG']	['TGCAGYAGCCG CCCTTCTGCCGG']	['Sorsby fundus dystrophy']
NM_006785.3(MALT 1):c.1019-2A>G	786200953	MALT1	['CGCYTTG AAAAAAA AAGAAAG GG']	['CGCYTTGAAAA AAAAAGAAAGG , 'TCGCYTTGAAAA AAAAAGAAAGG']	['Combined immunodeficiency']
NM_003639.4(IKBKG (p.Ter420Trp)	137853321	IKBKG	[]	[]	['Incontinentia pigmenti syndrome', 'Ectodermal

					dysplasia, anhidrotic, with immunodeficiency, osteopetrosis, and lymphedema']
NM_003639.4(IKBKG):c.1219A>G (p.Met407Val)	137853322	IKBKG	[]	['CCAYATCAGGG GCCTGATACTGG']	['Incontinentia pigmenti syndrome']
NM_001014797.2(KCNMA1):c.1301A>G (p.Asp434Gly)	137853333	KCNMA1	[]	[]	['Generalized epilepsy and paroxysmal dyskinesia']
NM_016218.2(POLK):c.1679A>T (p.Glu560Val)	757103131	POLK	[]	[]	['Malignant tumor of prostate']
NM_003000.2(SDHB):c.541-2A>G	786201161	SDHB	[]	[]	['Hereditary cancer-predisposing syndrome']
NM_032383.4(HPS3):c.2482-2A>G	397507168	-	[]	[]	['Hermansky-Pudlak syndrome 3']
NM_000060.3(BTD):c.968A>G (p.His323Arg)	397507176	BTD	[]	[]	['Biotinidase deficiency', 'not provided']
NM_004315.4(ASAHI):c.155A>G (p.Tyr52Cys)	137853595	ASAHI	[]	[]	['Farber lipogranulomatosis']
NM_004315.4(ASAHI):c.1006A>G (p.Asn336Asp)	137853596	ASAHI	[]	[]	['Farber lipogranulomatosis']
NM_000059.3(BRCA2):c.1799A>G (p.Tyr600Cys)	397507276	BRCA2	[]	[]	['Breast-ovarian cancer, familial 2']
NM_022912.2(REEP1):c.183-2A>G	387906264	REEP1	[]	[]	['Spastic paraplegia 31, autosomal dominant']
NM_000022.2(ADA):c.219-2A>G	387906267	ADA	[]	['CCCCYGGGAAG GGAAGAAAGGGG', 'GCCCCYGGGAAG GGAAGAAAGGG', 'AGCCCCYGGGAA GGGAAGAAAGG']	['Severe combined immunodeficiency due to ADA deficiency']
NM_018249.5(CDK5RAP2):c.4005-15A>G	387906274	CDK5RAP2	[]	[]	['Primary autosomal recessive microcephaly 3']
NM_032119.3(ADGRV1):c.14973-2A>G	371981035	ADGRV1	[]	[]	['Usher syndrome, type 2C']
NM_020366.3(RPGRI1):c.3341A>G (p.Asp1114Gly)	17103671	RPGRIP1	[]	[]	['Leber congenital amaurosis 6', 'not specified']
NM_000492.3(CFTR):c.3717+4A>G	387906362	CFTR	[]	['TCAAATCYCACC CTCTGGCCAGG']	['Cystic fibrosis']
NM_000492.3(CFTR):c.273+4A>G	387906374	CFTR	[]	[]	['Cystic fibrosis']
NM_002769.4(PRSS1):c.65A>G (p.Asp22Gly)	397507442	-	[]	['CTTGYCATCATC ATCAAAGGGGG', 'TCTTGYCATCATC ATCAAAGGGG', 'ATCTTGYCATCA	['Hereditary pancreatitis']

				TCATCAAAGGG', 'GATCTTGYCATC ATCATCAAAGG']	
NM_004006.2(DMD): c.4675-2A>G	794727575	DMD	[]	[]	['Duchenne muscular dystrophy', 'Becker muscular dystrophy']
NM_000132.3(F8):c.1 418A>G (p.Tyr473Cys)	387906444	F8	[]	[]	['Hereditary factor VIII deficiency disease']
NM_004333.4(BRAF): c.2126A>G (p.Gln709Arg)	397507486	BRAF	[]	[]	['Rasopathy']
NM_002834.3(PTPN1 1):c.124A>G (p.Thr42Ala)	397507501	PTPN11	[]	[]	['Noonan syndrome', 'Noonan syndrome 1', 'Rasopathy', 'not provided']
NM_000834.3(GRIN2 B):c.1238A>G (p.Glu413Gly)	527236034	GRIN2B	[]	[]	['Mental retardation, autosomal dominant 6']
NM_004830.3(MED23):c.3638A>G (p.His1213Arg)	527236035	MED23	[]	[]	['Mental retardation, autosomal recessive 18']
NM_002834.3(PTPN1 1):c.844A>G (p.Ile282Val)	397507529	PTPN11	[]	[]	['Noonan syndrome 1', 'Rasopathy', 'not provided']
NM_000406.2(GNRH R):c.851A>G (p.Tyr284Cys)	28933074	GNRHR	[]	[]	[]
NM_002834.3(PTPN1 1):c.1510A>G (p.Met504Val)	397507547	PTPN11	[]	[]	['Noonan syndrome', 'Noonan syndrome 1', 'Rasopathy', 'not provided']
NM_004629.1(FANC G):c.925-2A>G	397507561	FANCG	[]	[]	['Fanconi anemia, complementation group G']
NM_000492.3(CFTR): c.3140-26A>G	76151804	CFTR	[]	[]	['Cystic fibrosis']
NM_024598.3(USB1): c.502A>G (p.Arg168Gly)	137853971	USB1	[]	['CCACCYGGTTTT CTCTTGATTGG']	['Poikiloderma with neutropenia']
NM_000067.2(CA2):c. 754A>G (p.Asn252Asp)	2228063	CA2	[]	['TGTYCTTCAGTG GCTGAGCTGGG', 'CTGTYCTTCAGT GGCTGAGCTGG']	[]
NM_000138.4(FBN1): c.5096A>G (p.Tyr1699Cys)	387906622	FBN1	[]	[]	['Geleophysic dysplasia 2']
NM_001194958.2(KC NJ18):c.1097A>G (p.Lys366Arg)	527236159	KCNJ18	[]	[]	['Thyrotoxic periodic paralysis', 'Thyrotoxic periodic paralysis 2']
NM_000138.4(FBN1): c.5087A>G (p.Tyr1696Cys)	387906625	FBN1	[]	[]	['Geleophysic dysplasia 2']
NM_000138.4(FBN1): c.5099A>G (p.Tyr1700Cys)	387906626	FBN1	[]	[]	[]
NM_001244710.1(GF	387906638	GFPT1	[]	[]	['Congenital

PT1):c.43A>G (p.Thr15Ala)					myasthenic syndrome with tubular aggregates 1']
NM_002292.3(LAMB2):c.440A>G (p.His147Arg)	387906644	LAMB2	[]	[]	['Nephrotic syndrome, type 5, with or without ocular abnormalities']
NM_005188.3(CBL):c.1112A>G (p.Tyr371Cys)	387906666	CBL	[]	[]	[]
NM_000313.3(PROS1):c.701A>G (p.Tyr234Cys)	387906675	PROS1	['GATTAYA TCTGTAGC CTTCGGGG' , 'AGATTAY ATCTGTAG CCTTCGGG' , 'GAGATTA YATCTGTA GCCTTCGG']	['GATTAYATCTGT AGCCTTCGGGG', 'AGATTAYATCTG TAGCCTTCGGG', 'GAGATTAYATCT GTAGCCTTCGG']	['Thrombophilia due to protein S deficiency, autosomal recessive']
NM_032018.6(SPRTN):c.350A>G (p.Tyr117Cys)	527236213	SPRTN	[]	[]	['Ruijs-aalfs syndrome']
NM_022464.4(SIL1):c.645+2T>C	548535414	SIL1	[]	[]	['Marinesco-Sjogren syndrome']
NM_001040142.1(SCN2A):c.4419A>G (p.Ile1473Met)	387906685	SCN2A	[]	[]	['Early infantile epileptic encephalopathy 11']
NM_001040142.1(SCN2A):c.754A>G (p.Met252Val)	387906687	SCN2A	[]	[]	['Benign familial neonatal-infantile seizures']
m.10450A>G	387906731	MT-TR	[]	[]	['Mitochondrial encephalomyopathy']
m.5816A>G	387906732	MT-TC	[]	[]	[]
m.608A>G	387906735	MT-TF	['TTCAGYG TATTGCTT TGAGGAGG']	['TTCAGYGTATTG CTTTGAGGAGG']	[]
NM_001376.4(DYNC1H1):c.917A>G (p.His306Arg)	387906738	DYNC1H1	[]	[]	['Charcot-Marie-Tooth disease, axonal, type 2O', 'Charcot-Marie-Tooth disease', 'Spinal muscular atrophy, lower extremity predominant 1, autosomal dominant']
NM_001376.4(DYNC1H1):c.2011A>G (p.Lys671Glu)	387906742	DYNC1H1	[]	[]	['Spinal muscular atrophy, lower extremity predominant 1,

					autosomal dominant']
NM_001376.4(DYNC1H1):c.2909A>G (p.Tyr970Cys)	387906743	DYNC1H1	[]	['ATTCAAGYAGAT TACCTGATTGG']	['Spinal muscular atrophy, lower extremity predominant 1, autosomal dominant']
NM_001354.5(AKR1C2):c.235A>G (p.Ile79Val)	387906750	AKR1C2	[]	[]	['46,XY sex reversal 8']
NM_007315.3(STAT1):c.604A>G (p.Met202Val)	387906762	STAT1	[]	[]	['Immunodeficiency 31C']
NM_007315.3(STAT1):c.494A>G (p.Asp165Gly)	387906764	STAT1	[]	[]	['Immunodeficiency 31C']
NM_007315.3(STAT1):c.862A>G (p.Thr288Ala)	387906765	STAT1	[]	[]	['Immunodeficiency 31C']
NM_002052.4(GATA4):c.928A>G (p.Met310Val)	387906772	GATA4	[]	['TCCGCAYTGCAA GAGGCCTGGGG', 'TTCCGCAYTGCA AGAGGCCTGGG']	['Atrial septal defect 2']
NM_021615.4(CHST6):c.329A>G (p.Tyr110Cys)	72547544	CHST6	[]	[]	['Macular corneal dystrophy Type I']
NM_000209.3(PDX1):c.533A>G (p.Glu178Gly)	387906777	PDX1	[]	[]	['Pancreatic agenesis, congenital']
NM_000890.3(KCNJ5):c.472A>G (p.Thr158Ala)	387906778	KCNJ5	[]	[]	['Andersen Tawil syndrome', 'Familial hyperaldosteronism type 3']
NM_001184.3(ATR):c.6431A>G (p.Gln2144Arg)	387906797	ATR	[]	[]	['Cutaneous telangiectasia and cancer syndrome, familial']
NM_000382.2(ALDH3A2):c.1157A>G (p.Asn386Ser)	72547575	ALDH3A2	[]	[]	['Sjogren-Larsson syndrome']
NM_001005862.2(ERBB2):c.2480A>G (p.Asn827Ser)	28933370	ERBB2	[]	[]	['Neoplasm of ovary']
NM_006194.3(PAX9):c.271A>G (p.Lys91Glu)	28933373	PAX9	[]	[]	['Tooth agenesis, selective, 3']
NM_001083116.1(PRF1):c.755A>G (p.Asn252Ser)	28933375	PRF1	[]	[]	['Hemophagocytic lymphohistiocytosis, familial, 2', 'Malignant lymphoma, non-Hodgkin']
NM_005257.5(GATA6):c.1354A>G (p.Thr452Ala)	387906817	GATA6	[]	[]	['Pancreatic agenesis and congenital heart disease']
NM_000414.3(HSD17B4):c.650A>G (p.Tyr217Cys)	387906825	HSD17B4	[]	['TGCCACAYACTC TGGCTTCAGGG']	['Gonadal dysgenesis with auditory

					dysfunction, autosomal recessive inheritance']
NM_004153.3(ORC1):c.380A>G (p.Glu127Gly)	387906826	ORC1	[]	[]	['Meier-Gorlin syndrome 1']
NM_002552.4(ORC4):c.521A>G (p.Tyr174Cys)	387906847	ORC4	[]	[]	['Meier-Gorlin syndrome 2']
NM_004544.3(NDUFA10):c.1A>G (p.Met1Val)	387906872	NDUFA10	[]	[]	['Leigh syndrome due to mitochondrial complex I deficiency']
NM_004544.3(NDUFA10):c.425A>G (p.Gln142Arg)	387906873	NDUFA10	[]	[]	['Leigh syndrome due to mitochondrial complex I deficiency']
NM_006796.2(AFG3L2):c.1847A>G (p.Tyr616Cys)	387906889	AFG3L2	['GTAYAGA GGTATTGT TCTTTTGG']	['GTAYAGAGGTA TTGTTCTTTTGG']	['Spastic ataxia 5, autosomal recessive']
NM_006587.3(CORIN):c.949A>G (p.Lys317Glu)	387906894	CORIN	[]	[]	['Preeclampsia/eclampsia 5']
NM_006587.3(CORIN):c.1414A>G (p.Ser472Gly)	387906895	CORIN	[]	['GGATAACYTGTA CTGTTGTAGGG']	['Preeclampsia/eclampsia 5']
NM_015560.2(OPA1):c.1294A>G (p.Ile432Val)	387906899	OPA1	[]	[]	['Optic Atrophy Type 1']
NM_021625.4(TRPV4):c.590A>G (p.Lys197Arg)	387906903	TRPV4	[]	[]	['Metatrophic dysplasia']
NM_021625.4(TRPV4):c.826A>G (p.Lys276Glu)	387906907	TRPV4	[]	[]	['Metatrophic dysplasia']
NM_024022.2(TMPRS3):c.308A>G (p.Asp103Gly)	387906915	TMPRS3	[]	[]	['Deafness, autosomal recessive 8']
NM_019109.4(ALG1):c.1129A>G (p.Met377Val)	387906925	ALG1	[]	[]	['Congenital disorder of glycosylation type 1K']
NM_006886.3(ATP5E):c.35A>G (p.Tyr12Cys)	387906929	-	[]	[]	['Nuclearly-encoded mitochondrial complex V (ATP synthase) deficiency 3']
NM_032578.3(MYPN):c.59A>G (p.Tyr20Cys)	140148105	MYPN	[]	[]	['Primary dilated cardiomyopathy', 'Cardiomyopathy', 'Dilated cardiomyopathy 1KK', 'Familial hypertrophic cardiomyopathy 22', 'not provided']
NM_016013.3(NDUFAF1):c.758A>G (p.Lys253Arg)	387906957	NDUFAF1	[]	['ACCYTGACCTCC TGCCAGTAGGG', 'TACCYTGACCTC']	['Mitochondrial complex I deficiency']

				CTGCCAGTAGG']	
NM_032580.3(HES7): c.172A>G (p.Ile58Val)	387906979	HES7	[]	[]	['Spondylocostal dysostosis 5']
NM_024700.3(SNIP1) :c.1097A>G (p.Glu366Gly)	387906986	SNIP1	[]	[]	['Psychomotor retardation, epilepsy, and craniofacial dysmorphism']
NM_016952.4(CDON) :c.2368A>G (p.Thr790Ala)	387906997	CDON	[]	[]	['Holoprosencephaly 11']
NM_031427.3(DNAL 1):c.449A>G (p.Asn150Ser)	387907021	DNAL1	['AGGGAYT GCCTACAA ACACCAGG ']	['AGGGAYTGCCT ACAAACACCAGG' ']	['Kartagener syndrome', 'Ciliary dyskinesia, primary, 16']
NM_020320.3(RARS2)):c.1024A>G (p.Met342Val)	387907048	RARS2	[]	[]	['Pontocerebellar hypoplasia type 6']
NM_000138.4(FBN1): c.6431A>G (p.Asn2144Ser)	137854461	FBN1	[]	[]	['Marfan syndrome']
NM_000138.4(FBN1): c.2261A>G (p.Tyr754Cys)	137854479	FBN1	[]	[]	['Marfan syndrome']
NM_212482.1(FN1):c. 2918A>G (p.Tyr973Cys)	137854488	FN1	['GAAGTAA YAGGTGAC CCCAGGGG ']	['GAAGTAAAYAGG TGACCCCAGGGG' ']	['Glomerulopathy with fibronectin deposits 2']
NM_198994.2(TGM6) :c.980A>G (p.Asp327Gly)	387907098	TGM6	[]	[]	['Spinocerebellar ataxia 35']
NM_001363.4(DKC1): c.1069A>G (p.Thr357Ala)	137854492	DKC1	['GCAGGYA GAGATGAC CGCTGTGG' ']	['GCAGGYAGAGA TGACCGCTGTGG']	['Dyskeratosis congenita X-linked']
NM_052873.2(IFT43): c.1A>G (p.Met1Val)	387907107	IFT43	[]	[]	['Cranoectodermal dysplasia 3']
NM_201269.2(ZNF64 4):c.2014A>G (p.Ser672Gly)	387907109	ZNF644	[]	[]	['Myopia 21, autosomal dominant']
NM_018699.3(PRDM 5):c.320A>G (p.Tyr107Cys)	387907111	PRDM5	[]	[]	['Brittle cornea syndrome 2']
NM_000132.3(F8):c.5 822A>G (p.Asn1941Ser)	28933682	F8	[]	['TAGCCAYTGATT GCTGGAGAAGG']	['Hereditary factor VIII deficiency disease']
NM_016464.4(TMEM 138):c.287A>G (p.His96Arg)	387907132	TMEM13 8	['GACAYGA AGGGAGAT GCTGAGGG ']	['GACAYGAAGGG AGATGCTGAGGG', 'AGACAYGAAGGG AGATGCTGAGG']	['Joubert syndrome 16']
NM_016464.4(TMEM 138):c.389A>G (p.Tyr130Cys)	387907135	TMEM13 8	[]	['CAGYACAACAC TGCTGCTGTGGG', 'GCAGYACAACAC TGCTGCTGTGG']	['Joubert syndrome 16']
NM_177965.3(C8orf3 7):c.545A>G (p.Gln182Arg)	387907137	C8orf37	[]	[]	['Retinitis pigmentosa 64']
NM_001077488.3(GN AS):c.1A>G (p.Met1Val)	137854530	GNAS	[]	['GCCAYGGCGG CGGCGGCGGCGG' ']	['Pseudohypoparathy roidism type 1A']

NM_006920.4(SCN1A):c.2557-2A>G	727504140	SCN1A	[]	[]	['Severe myoclonic epilepsy in infancy', 'Generalized epilepsy with febrile seizures plus, type 2']
NM_000308.2(CTSA):c.200A>G (p.Gln67Arg)	137854541	CTSA	[]	[]	['Combined deficiency of sialidase AND beta galactosidase']
NM_000308.2(CTSA):c.1238A>G (p.Tyr413Cys)	137854543	CTSA	[]	[]	['Combined deficiency of sialidase AND beta galactosidase']
NM_000308.2(CTSA):c.1411A>G (p.Lys471Glu)	137854549	CTSA	[]	[]	['Galactosialidosis, late infantile']
NM_000267.3(NF1):c.4267A>G (p.Lys1423Glu)	137854550	NF1	[]	[]	['Neurofibromatosis, type 1']
NM_000267.3(NF1):c.1466A>G (p.Tyr489Cys)	137854557	NF1	['ACTTAYA GCTTCTTG TCTCCAGG']	['ACTTAYAGCTTC TTGTCTCCAGG']	['Neurofibromatosis, type 1']
NM_018105.2(THAP1):c.70A>G (p.Lys24Glu)	387907176	THAP1	[]	['CCTCACTYGTGG AAAGAAACGGG']	['Dystonia 6, torsion']
NM_000492.3(CFTR):c.1393-2A>G	397508201	CFTR	[]	[]	['Cystic fibrosis']
NM_001172646.1(PLCB4):c.986A>G (p.Asn329Ser)	387907179	PLCB4	[]	[]	['Auriculocondylar syndrome 2']
NM_005850.4(SF3B4):c.1A>G (p.Met1Val)	387907185	SF3B4	[]	[]	['Nager syndrome']
NM_014714.3(IFT140):c.932A>G (p.Tyr311Cys)	387907193	IFT140	[]	[]	['Renal dysplasia, retinal pigmentary dystrophy, cerebellar ataxia and skeletal dysplasia']
NM_005006.6(NDUF S1):c.1783A>G (p.Thr595Ala)	387907199	NDUFS1	[]	[]	['Mitochondrial complex I deficiency']
NM_000397.3(CYBB):c.1499A>G (p.Asp500Gly)	137854593	CYBB	[]	['TCACAYCTTTCT CCTCATCATGG']	['Chronic granulomatous disease, X-linked', 'not provided']
NM_033360.3(KRAS):c.439A>G (p.Lys147Glu)	387907206	KRAS	[]	[]	['Cardiofaciocutaneous syndrome 2']
NM_000335.4(SCN5A):c.5381A>G (p.Tyr1794Cys)	137854614	SCN5A	[]	[]	['Long QT syndrome 3', 'Congenital long QT syndrome']
NM_000076.2(CDKN1C):c.832A>G (p.Lys278Glu)	387907226	CDKN1C	[]	['CGCTYGGCGAA GAAATCTGCGGG', 'GCGCTYGGCGAA GAAATCTGCGG']	['Intrauterine growth retardation, metaphyseal dysplasia, adrenal hypoplasia congenita, and genital anomalies']

NM_022912.2(REEP1):c.304-2A>G	387907242	REEP1	[]	['TCCYGTCAAAGG AAAAACAGAGG']	['Distal hereditary motor neuropathy type 5B']
NM_198253.2(TERT):c.2705A>G (p.Lys902Arg)	387907250	TERT	[]	[]	['PULMONARY FIBROSIS AND/OR BONE MARROW FAILURE, TELOMERE-RELATED, 1']
NM_005349.3(RBPJ):c.188A>G (p.Glu63Gly)	387907270	RBPJ	[]	[]	['Adams-Oliver syndrome 3']
NM_005349.3(RBPJ):c.505A>G (p.Lys169Glu)	387907271	RBPJ	[]	[]	['Adams-Oliver syndrome 3']
NM_022787.3(NMNAT1):c.817A>G (p.Asn273Asp)	387907291	NMNAT1	[]	['TGTYTCTCTGCA AAGGGGCCAGG']	['Leber congenital amaurosis 9']
NM_000492.3(CFTR):c.1A>G (p.Met1Val)	397508328	CFTR	['TGCA YGG TCTCTCGG GCGCTGGG ']	['GCAYGGTCTCTC GGGCGCTGGGG', 'TGCA YGGTCTCT CGGGCGCTGGG', 'CTGCA YGGTCTC TCGGGCGCTGG']	['Cystic fibrosis']
NM_020921.3(NIN):c.5126A>G (p.Asn1709Ser)	387907308	NIN	[]	[]	['Seckel syndrome 7']
NM_021629.3(GNB4):c.265A>G (p.Lys89Glu)	387907341	GNB4	[]	[]	['Charcot-Marie-Tooth disease, dominant intermediate F']
NM_000355.3(TCN2):c.580+624A>T	372866837	TCN2	[]	[]	[]
NM_032415.5(CARD11):c.401A>G (p.Glu134Gly)	387907351	CARD11	[]	[]	['B-CELL EXPANSION WITH NFKB AND T-CELL ANERGY']
NM_005430.3(WNT1):c.624+4A>G	387907354	WNT1	[]	[]	['Osteogenesis imperfecta type 15']
NM_207352.3(CYP4V2):c.367A>G (p.Met123Val)	149684063	CYP4V2	[]	[]	['Bietti crystalline corneoretinal dystrophy', 'not provided']
NM_031885.3(BBS2):c.472-2A>G	137854887	BBS2	[]	[]	['Bardet-Biedl syndrome 2']
NM_015268.3(DNAJC13):c.2564A>G (p.Asn855Ser)	387907571	DNAJC13	[]	[]	['Parkinson disease, late-onset', 'Essential tremor', 'PARKINSON DISEASE 21']
NM_001287.5(CLCN7):c.296A>G (p.Tyr99Cys)	387907576	CLCN7	[]	['TGTCAYAGTCCA AGCTCTGCAGG']	['Osteopetrosis autosomal dominant type 2', 'Osteopetrosis autosomal recessive 4']

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EQUIVALENTS AND SCOPE

[00416] Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents of the embodiments described herein. The scope of the present disclosure is not intended to be limited to the above description, but rather is as set forth in the appended claims.

[00417] Articles such as “a,” “an,” and “the” may mean one or more than one unless indicated to the contrary or otherwise evident from the context. Claims or descriptions that include “or” between two or more members of a group are considered satisfied if one, more than one, or all of the group members are present, unless indicated to the contrary or otherwise evident from the context. The disclosure of a group that includes “or” between two or more group members provides embodiments in which exactly one member of the group is present, embodiments in which more than one members of the group are present, and embodiments in which all of the group members are present. For purposes of brevity those embodiments have not been individually spelled out herein, but it will be understood that each of these embodiments is provided herein and may be specifically claimed or disclaimed.

[00418] It is to be understood that the invention encompasses all variations, combinations, and permutations in which one or more limitation, element, clause, or descriptive term, from one or more of the claims or from one or more relevant portion of the description, is introduced into another claim. For example, a claim that is dependent on another claim can be modified to include one or more of the limitations found in any other claim that is

dependent on the same base claim. Furthermore, where the claims recite a composition, it is to be understood that methods of making or using the composition according to any of the methods of making or using disclosed herein or according to methods known in the art, if any, are included, unless otherwise indicated or unless it would be evident to one of ordinary skill in the art that a contradiction or inconsistency would arise.

[00419] Where elements are presented as lists, *e.g.*, in Markush group format, it is to be understood that every possible subgroup of the elements is also disclosed, and that any element or subgroup of elements can be removed from the group. It is also noted that the term “comprising” is intended to be open and permits the inclusion of additional elements or steps. It should be understood that, in general, where an embodiment, product, or method is referred to as comprising particular elements, features, or steps, embodiments, products, or methods that consist, or consist essentially of, such elements, features, or steps, are provided as well. For purposes of brevity those embodiments have not been individually spelled out herein, but it will be understood that each of these embodiments is provided herein and may be specifically claimed or disclaimed.

[00420] Where ranges are given, endpoints are included. Furthermore, it is to be understood that unless otherwise indicated or otherwise evident from the context and/or the understanding of one of ordinary skill in the art, values that are expressed as ranges can assume any specific value within the stated ranges in some embodiments, to the tenth of the unit of the lower limit of the range, unless the context clearly dictates otherwise. For purposes of brevity, the values in each range have not been individually spelled out herein, but it will be understood that each of these values is provided herein and may be specifically claimed or disclaimed. It is also to be understood that unless otherwise indicated or otherwise evident from the context and/or the understanding of one of ordinary skill in the art, values expressed as ranges can assume any subrange within the given range, wherein the endpoints of the subrange are expressed to the same degree of accuracy as the tenth of the unit of the lower limit of the range.

[00421] In addition, it is to be understood that any particular embodiment of the present invention may be explicitly excluded from any one or more of the claims. Where ranges are given, any value within the range may explicitly be excluded from any one or more of the claims. Any embodiment, element, feature, application, or aspect of the compositions and/or methods of the invention, can be excluded from any one or more claims. For purposes of brevity, all of the embodiments in which one or more elements, features, purposes, or aspects is excluded are not set forth explicitly herein.

CLAIMS

What is claimed is:

1. A fusion protein comprising: (i) a Cas9 domain; (ii) a cytidine deaminase domain; and (iii) a uracil glycosylase inhibitor (UGI) domain.
2. The fusion protein of claim 1, wherein the Cas9 domain comprises an amino acid sequence that is at least 85% identical to the amino acid sequence provided in SEQ ID NO: 674.
3. The fusion protein of claim 1, wherein the Cas9 domain is a Cas9 nickase domain that cuts a nucleotide target strand of a nucleotide duplex, wherein the nucleotide target strand is the strand that binds to a gRNA of the Cas9 nickase domain.
4. The fusion protein of claim 1, wherein the Cas9 domain is an nCas9 domain that comprises a D10A mutation in the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
5. The fusion protein of claim 1, wherein the Cas9 domain is an nCas9 domain that comprises one or more of N496A, R660A, Q694A, and Q926A of the amino acid sequence provided in SEQ ID NO 10, or one or more corresponding mutations in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
6. The fusion protein of claim 1, wherein the cytidine deaminase domain is a deaminase from the apolipoprotein B mRNA-editing complex (APOBEC) family deaminase.
7. The fusion protein of claim 6, wherein the APOBEC family deaminase is selected from the group consisting of APOBEC1 deaminase, APOBEC2 deaminase, APOBEC3A deaminase, APOBEC3B deaminase, APOBEC3C deaminase, APOBEC3D deaminase, APOBEC3F deaminase, APOBEC3G deaminase, and APOBEC3H deaminase.

8. The fusion protein of claim 1, wherein the cytidine deaminase domain comprises an amino acid sequence that is at least 85% identical to an amino acid sequence of SEQ ID NO: 266-284, 607-610, 5724-5736, or 5738-5741.
9. The fusion protein of claim 1, wherein the cytidine deaminase domain comprises an amino acid sequence of SEQ ID NO: 266-284, 607-610, 5724-5736, or 5738-5741.
10. The fusion protein of claim 1, wherein the cytidine deaminase domain is a rat APOBEC1 (rAPOBEC1) deaminase comprising one or more mutations selected from the group consisting of W90Y, R126E, and R132E of SEQ ID NO: 284, or one or more corresponding mutations in another APOBEC deaminase.
11. The fusion protein of claim 1, wherein the cytidine deaminase domain is a human APOBEC1 (hAPOBEC1) deaminase comprising one or more mutations selected from the group consisting of W90Y, Q126E, and R132E of SEQ ID NO: 5724, or one or more corresponding mutations in another APOBEC deaminase.
12. The fusion protein of claim 1, wherein the cytidine deaminase domain is a human APOBEC3G (hAPOBEC3G) deaminase comprising one or more mutations selected from the group consisting of W285Y, R320E, and R326E of SEQ ID NO: 275, or one or more corresponding mutations in another APOBEC deaminase.
13. The fusion protein of claim 1, wherein the cytidine deaminase domain is an activation-induced deaminase (AID).
14. The fusion protein of claim 1, wherein the cytidine deaminase domain is a cytidine deaminase 1 from *Petromyzon marinus* (pmCDA1).
15. The fusion protein of claim 1, wherein the UGI domain comprises a domain capable of inhibiting UDG activity.
16. The fusion protein of claim 1, wherein the UGI domain comprises an amino acid sequence that is at least 85% identical to SEQ ID NO: 600.

17. The fusion protein of claim 1, wherein the UGI domain comprises an amino acid sequence as set forth in SEQ ID NO: 600.
18. The fusion protein of claim 1, wherein the fusion protein comprises the structure: NH₂-[cytidine deaminase domain]-[Cas9 domain]-[UGI domain]-COOH, and wherein each instance of “-” comprises an optional linker.
19. The fusion protein of claim 1, wherein the cytidine deaminase domain of (ii) and the nCas9 domain of (i) are linked via a linker comprising the amino acid sequence (GGGS)_n (SEQ ID NO: 265), (GGGGGS)_n (SEQ ID NO: 5), (G)_n, (EAAAK)_n (SEQ ID NO: 6), (GGS)_n, (SGGS)_n (SEQ ID NO: 4288), SGSETPGTSESATPES (SEQ ID NO: 7), or (XP)_n motif, or a combination thereof, wherein _n is independently an integer between 1 and 30, inclusive, and wherein X is any amino acid.
20. The fusion protein of claim 1, wherein the cytidine deaminase domain of (ii) and the nCas9 domain of (i) are linked via a linker comprising the amino acid sequence: SGSETPGTSESATPES (SEQ ID NO: 7).
21. The fusion protein of claim 1 further comprising a nuclear localization sequence (NLS).
22. The fusion protein of claim 21, wherein the NLS comprises the amino acid sequence PKKKRKV (SEQ ID NO: 741) or MDSLLMNRRKFLYQFKNVRWAKGRRETYLC (SEQ ID NO: 742)
23. The fusion protein of claim 21, wherein the fusion protein comprises the structure: NH₂-[cytidine deaminase domain]-[nCas9 domain]-[UGI domain]-[NLS]-COOH, and wherein each instance of “-” comprises an optional linker.
24. The fusion protein of claim 21, wherein the UGI domain and the NLS are linked via a linker comprising the amino acid sequence: SGGS (SEQ ID NO: 4288), or wherein the nCas9 domain and the UGI domain are linked via a linker comprising the amino acid sequence: SGGS (SEQ ID NO: 4288).

25. The fusion protein of claim 1, wherein the fusion protein comprises the amino acid sequence set forth in SEQ ID NO: 594.
26. A complex comprising the fusion protein of claim 1 and a guide RNA bound to the nCas9 domain of the fusion protein.
27. A method comprising contacting a nucleic acid molecule with the fusion protein of claim 1 and a guide RNA, wherein the guide RNA comprises a sequence of at least 10 contiguous nucleotides that is complementary to a target sequence in the genome of an organism and comprises a target base pair.
28. The method of claim 27, wherein the target base pair comprises a T to C point mutation associated with a disease or disorder, and wherein the deamination of the mutant C base results in a sequence that is not associated with a disease or disorder.
29. The method of claim 27, wherein the contacting results in less than 20% indel formation upon base editing.
30. The method of claim 27, wherein the contacting results in at least 2:1 intended to unintended product upon base editing.
31. A fusion protein comprising: (i) a nuclease-inactive Cas9 (dCas9) domain and (ii) an apolipoprotein B mRNA-editing complex 1 (APOBEC1) deaminase domain, wherein the deaminase domain is fused to the N-terminus of the dCas9 domain via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).
32. The fusion protein of claim 31, wherein the nuclease-inactive Cas9 (dCas9) domain of (i) comprises the amino acid sequence that is at least 85% identical to the amino acid sequence set forth in SEQ ID NO: 263.
33. The fusion protein of claim 31, wherein the nuclease-inactive Cas9 (dCas9) domain of (i) comprises the amino acid sequence set forth in SEQ ID NO: 263.

34. The fusion protein of any one of claims 31-33, wherein the deaminase is a rat APOBEC1 deaminase that is at least 85% identical the amino acid sequence as set forth in (SEQ ID NO: 284).
35. The fusion protein of any one of claims 31-34, wherein the deaminase is rat APOBEC1 deaminase comprising the amino acid sequence as set forth in (SEQ ID NO: 284).
36. The fusion protein of any one of claims 31-33, wherein the deaminase is a human APOBEC1 deaminase that is at least 85% identical to the amino acid sequence as set forth in (SEQ ID NO: 282).
37. The fusion protein of any one of claims 31-33, wherein the deaminase is a human APOBEC1 deaminase comprising the amino acid sequence as set forth in (SEQ ID NO: 282).
38. The fusion protein of any one of claims 31-37, wherein the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to SEQ ID NO: 600.
39. The fusion protein of any one of claims 31-38, wherein the UGI domain comprises the amino acid sequence as set forth in SEQ ID NO: 600
40. The fusion protein of any one of claims 31-37, wherein the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of SEQ ID NOs: 322-324.
41. The fusion protein of any one of claims 31-37, wherein the UGI domain comprises the amino acid sequence as set forth in any one of SEQ ID NOs: 322-324.
42. The fusion protein of any one of claims 31-41, wherein the fusion protein comprises amino acid residues 11-1629 of the amino acid sequence set forth in SEQ ID NO: 591.

43. The fusion protein of any one of claims 31-41, wherein the fusion protein comprises the amino acid sequence set forth in any one of SEQ ID NOs: 591-593, 611, 612, 615, 657, 658, and 5737.
44. A fusion protein comprising: (i) a nuclease-inactive Cas9 (dCas9) domain; (ii) a nucleic acid editing domain; and (iii) a uracil glycosylase inhibitor (UGI) domain.
45. The fusion protein of claim 44, wherein the amino acid sequence of the dCas9 domain comprises a D10X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for D.
46. The fusion protein of claim 44 or 45, wherein the amino acid sequence of the dCas9 domain comprises a D10A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
47. The fusion protein of any one of claims 44-46, wherein the amino acid sequence of the dCas9 domain comprises an H840X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for H.
48. The fusion protein of any one of claims 44-47, wherein the amino acid sequence of the dCas9 domain comprises an H840A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
49. The fusion protein of any one of claims 44-48, wherein the dCas9 domain comprises an amino acid sequence that is at least 85% identical to the amino acid sequence as set forth in SEQ ID NO: 263.
50. The fusion protein of any one of claims 44-49, wherein the dCas9 domain comprises the amino acid sequence as set forth in SEQ ID NO: 263.

51. The fusion protein of any one of claims 44-50, wherein the dCas9 domain comprises one or more of a N497X, R661X, Q695X, and Q926X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid.
52. The fusion protein of any one of claims 44-51, wherein the dCas9 domain comprises one or more of a N497A, R661A, Q695A, and Q926A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
53. The fusion protein of any one of claims 44-52, wherein the dCas9 domain comprises a N497A, R661A, Q695A, and Q926A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
54. The fusion protein of any one of claims 44-53, wherein the dCas9 domain comprises a *Staphylococcus aureus* (SaCas9).
55. The fusion protein of claim 54, wherein the SaCas9 comprises the amino acid sequence SEQ ID NO: 4273.
56. The fusion protein of claim 54 or 55, wherein the SaCas9 domain comprises one or more of a E781K, N967K, or R1014H mutation of SEQ ID NO: 4273, or one or more corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
57. The fusion protein of any one of claims 44-53, wherein the dCas9 domain comprises one or more of a D1134E, R1334Q, and T1336R mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
58. The fusion protein of any one of claims 44-53, wherein the dCas9 domain comprises one or more of a D1134V, R1334Q, and T1336R mutation of SEQ ID NO: 4276, or a

corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.

59. The fusion protein of any one of claims 44-53, wherein the dCas9 domain comprises one or more of a D1134V, G1217R, R1334Q, and T1336R mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.

60. The fusion protein of any of claims 44-59, wherein the nucleic acid editing domain is fused to the N-terminus of the dCas9 domain.

61. The fusion protein of any one of claims 44-60, wherein the UGI domain is fused to the C-terminus of the dCas9 domain.

62. The fusion protein of any one of claims 44-61, wherein the dCas9 domain and the nucleic acid editing domain are fused via a linker.

63. The fusion protein of any one of claims 44-62, wherein the dCas9 domain and the UGI domain are fused via a linker.

64. The fusion protein of claim 62 or 63, wherein the linker comprises the amino acid sequence (GGGGS)_n (SEQ ID NO: 5), (G)_n, (EAAAK)_n (SEQ ID NO: 6), (GGS)_n, SGSETPGTSESATPES (SEQ ID NO: 7), SGGGS (SEQ ID NO: 4288), (XP)_n, or any combination thereof, wherein n is independently an integer between 1 and 30, and wherein X is any amino acid.

65. The fusion protein of claim 62 or 63, wherein the linker comprises a covalent bond.

66. The fusion protein of claim 64, wherein the linker comprises the amino acid sequence (GGS)_n, wherein n is 1, 3, or 7.

67. The fusion protein of claim 64, wherein the linker comprises the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).

68. The fusion protein of claim 62, wherein the dCas9 domain and the nucleic acid editing domain are fused via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).
69. The fusion protein of claim 62, wherein the dCas9 domain and the nucleic acid editing domain are fused via a linker comprising the amino acid sequence (GGS)_n, wherein n is 1, 3, or 7.
70. The fusion protein of claim 63, wherein the dCas9 domain and the UGI domain are fused via a linker comprising the amino acid sequence (GGGGGS)_n (SEQ ID NO: 5), (G)_n, (EAAAK)_n (SEQ ID NO: 6), (GGS)_n, SGSETPGTSESATPES (SEQ ID NO: 7), SGGS (SEQ ID NO: 4288), (XP)_n, or any combination thereof, wherein n is independently an integer between 1 and 30, and wherein X is any amino acid.
71. The fusion protein of claim 63, wherein the dCas9 domain and the UGI domain are fused via a linker comprising the amino acid sequence SGGS (SEQ ID NO: 4288).
72. The fusion protein of any one of claims 44-71, wherein the fusion protein comprises the structure [nucleic acid editing domain]-[optional linker]-[dCas9 domain]-[optional linker]-[UGI].
73. The fusion protein of any one of claims 44-67 wherein the fusion protein comprises the structure [nucleic acid editing domain]-[optional linker]-[UGI]-[optional linker]-[dCas9]; [UGI]-[optional linker]-[nucleic acid editing domain]-[optional linker]-[dCas9]; [UGI]-[optional linker]-[dCas9]-[optional linker]-[nucleic acid editing domain]; [dCas9]-[optional linker]-[UGI]-[optional linker]-[nucleic acid editing domain]; or [dCas9]-[optional linker]-[nucleic acid editing domain]-[optional linker]-[UGI].
74. The fusion protein of any one of claims 44-73, wherein the nucleic acid editing domain comprises a deaminase.
75. The fusion protein of claim 74 wherein the deaminase is a cytidine deaminase.

76. The fusion protein of claim 74 or 75, wherein the deaminase is an apolipoprotein B mRNA-editing complex (APOBEC) family deaminase.
77. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC1 deaminase.
78. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC2 deaminase.
79. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC3A deaminase.
80. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC3B deaminase.
81. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC3C deaminase.
82. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC3D deaminase.
83. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC3F deaminase.
84. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC3G deaminase.
85. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC3H deaminase.
86. The fusion protein of any one of claims 74-76, wherein the deaminase is an APOBEC4 deaminase.

87. The fusion protein of claim 74 or 75, wherein the deaminase is an activation-induced deaminase (AID).
88. The fusion protein of claim 74 or 75, wherein the deaminase is an APOBEC deaminase comprising one or more mutations selected from the group consisting of H121R, H122R, R126A, R126E, R118A, W90A, W90Y, and R132E of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase.
89. The fusion protein of claim 74 or 75, wherein the deaminase is an APOBEC deaminase comprising a W90Y, a R126E, and a R132E mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase.
90. The fusion protein of claim 74 or 75, wherein the deaminase comprises one or more mutations selected from the group consisting of D316R, D317R, R320A, R320E, R313A, W285A, W285Y, R326E of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase.
91. The fusion protein of claim 74 or 75, wherein the deaminase is an APOBEC deaminase comprising a W285Y, a R320E, and a R326E mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase.
92. The fusion protein of any one of claims 74-91, wherein the deaminase is from a human, chimpanzee, gorilla, monkey, cow, dog, rat, or mouse.
93. The fusion protein of any one of claims 74-92, wherein the deaminase is from a human.
94. The fusion protein of any one of claims 74-92, wherein the deaminase is from a rat.
95. The fusion protein of claim 74 or 75, wherein the deaminase is an cytidine deaminase 1 from *Petromyzon marinus* (pmCDA1).
96. The fusion protein of any one of claims 74-76, wherein the deaminase is a rat APOBEC1 deaminase comprising the amino acid sequence set forth in (SEQ ID NO: 284).

97. The fusion protein of any one of claims 74-76, wherein the deaminase is a human APOBEC1 deaminase comprising the amino acid sequence set forth in (SEQ ID NO: 282).
98. The fusion protein of claim 95, wherein the pmCDA1 comprises an amino acid sequence set forth in (SEQ ID NO: 5738).
99. The fusion protein of claim 84, wherein the APOBEC3G is a human APOBEC3G comprising the amino acid sequence set forth in (SEQ ID NO: 275).
100. The fusion protein of claim 84, wherein the APOBEC3G is a human APOBEC3G variant comprising the amino acid sequence set forth in any one of (SEQ ID NOs: 5739-5741).
101. The fusion protein of claim 74 or 75, wherein the deaminase is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of the amino acid sequences set forth in SEQ ID NOs: 266-284, 607-610, 5724-5736, and 5738-5741.
102. The fusion protein of claim 74 or 75, wherein the deaminase comprises the amino acid sequence set forth in any one of SEQ ID NOs: 266-284, 607-610, 5724-5736, and 5738-5741.
103. The fusion protein of any one of claims 44-102, wherein the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to SEQ ID NO: 600.
104. The fusion protein of any one of claims 44-103, wherein the UGI domain comprises the amino acid sequence as set forth in SEQ ID NO: 600
105. The fusion protein of any one of claims 44-102, wherein the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least

95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of SEQ ID NOs: 322-324.

106. The fusion protein of any one of claims 44- 102, wherein the UGI domain comprises the amino acid sequence as set forth in any one of SEQ ID NOs: 322-324.

107. A fusion protein comprising: (i) a Cas9 nickase domain and (ii) an apolipoprotein B mRNA-editing complex 1 (APOBEC1) deaminase domain, wherein the deaminase domain is fused to the N-terminus of the Cas9 nickase domain via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).

108. The fusion protein of claim 107, wherein the deaminase is rat APOBEC1 (SEQ ID NO: 284).

109. The fusion protein of claim 107 or 108, wherein the deaminase is human APOBEC1 (SEQ ID NO: 282).

110. A fusion protein comprising: (i) a Cas9 nickase domain and (ii) an apolipoprotein B mRNA-editing complex 3G (APOBEC3G) deaminase domain, wherein the deaminase domain is fused to the N-terminus of the Cas9 nickase domain via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).

111. The fusion protein of claim 110, wherein the deaminase is a human APOBEC3G deaminase comprising an amino acid sequence at least 85% identical to the amino acid sequence set forth in (SEQ ID NO: 275).

112. The fusion protein of claim 110 or 111, wherein the deaminase is a human APOBEC3G (SEQ ID NO: 275).

113. The fusion protein of claim 110, wherein the APOBEC3G is a human APOBEC3G variant comprising an amino acid sequence that is at least 85% identical to the amino acid sequence as set forth in any one of (SEQ ID NOs: 5739-5741).

114. The fusion protein of claim 110, wherein the APOBEC3G is a human APOBEC3G variant comprising the amino acid sequence set forth in any one of (SEQ ID NOs: 5739-5741).
115. A fusion protein comprising: (i) a Cas9 nickase domain and (ii) pmCDA1 domain, wherein the deaminase domain is fused to the N-terminus of the Cas9 nickase domain via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).
116. The fusion protein of claim 115, wherein the pmCDA1 comprises an amino acid sequence that is at least 85% identical to the amino acid sequence as set forth in (SEQ ID NO: 5738).
117. The fusion protein of claim 115 or 116, wherein the pmCDA1 comprises an amino acid sequence set forth in (SEQ ID NO: 5738).
118. The fusion protein of any one of claims 107-117, wherein the amino acid sequence of the Cas9 nickase domain comprises a D10X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for D.
119. The fusion protein of any one of claims 107-118, wherein the amino acid sequence of the Cas9 nickase domain comprises a D10A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
120. The fusion protein of any one of claims 107-119, wherein the amino acid sequence of the Cas9 nickase domain comprises a histidine at amino acid position 840 of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding amino acid position in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
121. The fusion protein of any one of claims 107-120, wherein the amino acid sequence of the Cas9 nickase domain comprises the amino acid sequence that is at least 85% identical to the amino acid sequence as set forth in SEQ ID NO: 674.

122. The fusion protein of any one of claims 107- 121, wherein the amino acid sequence of the Cas9 nickase domain comprises the amino acid sequence as set forth in SEQ ID NO: 674.

123. The fusion protein of any one of claims 107-122, wherein the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to SEQ ID NO: 600.

124. The fusion protein of any one of claims 107-123, wherein the UGI domain comprises the amino acid sequence as set forth in SEQ ID NO: 600

125. The fusion protein of any one of claims 107-122, wherein the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of SEQ ID NOs: 322-324.

126. The fusion protein of any one of claims 107-122, wherein the UGI domain comprises the amino acid sequence as set forth in any one of SEQ ID NOs: 322-324.

127. The fusion protein of any one of claims 107-126, wherein the fusion protein comprises the amino acid sequence set forth in any one of SEQ ID NOs: 594, 5743, 5745, and 5746.

128. A fusion protein comprising: (i) a Cas9 nickase (nCas9) domain; (ii) a nucleic acid editing domain; and (iii) a uracil glycosylase inhibitor (UGI) domain.

129. The fusion protein of claim 128, wherein the amino acid sequence of the Cas9 nickase domain comprises a D10X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid except for D.

130. The fusion protein of claim 128 or 129, wherein the amino acid sequence of the Cas9 nickase domain comprises a D10A mutation of the amino acid sequence provided in SEQ ID

NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.

131. The fusion protein of any one of claims 128-130, wherein the amino acid sequence of the Cas9 nickase domain comprises a histidine at amino acid position 840 of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding amino acid position in any of the amino acid sequences provided in SEQ ID NOs: 11-260.

132. The fusion protein of any one of claims 128-131, wherein the amino acid sequence of the Cas9 nickase domain comprises an amino acid sequence that is at least 85% identical to the amino acid sequence as set forth in SEQ ID NO: 674.

133. The fusion protein of any one of claims 128-131, wherein the amino acid sequence of the Cas9 nickase domain comprises the amino acid sequence as set forth in SEQ ID NO: 674.

134. The fusion protein of any one of claims 128-133, wherein the Cas9 nickase domain comprises one or more of a N497X, R661X, Q695X, and Q926X mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260, wherein X is any amino acid.

135. The fusion protein of any one of claims 128-134, wherein the Cas9 nickase domain comprises one or more of a N497A, R661A, Q695A, and Q926A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.

136. The fusion protein of any one of claims 128-135, wherein the Cas9 nickase domain comprises a N497A, R661A, Q695A, and Q926A mutation of the amino acid sequence provided in SEQ ID NO: 10, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.

137. The fusion protein of any one of claims 128-136, wherein the Cas9 nickase domain comprises a *Staphylococcus aureus* (SaCas9).

138. The fusion protein of claim 137, wherein the SaCas9 comprises the amino acid sequence SEQ ID NO: 4273.
139. The fusion protein of claim 137 or 138, wherein the SaCas9 comprises one or more of a E781K, N967K, or R1014H mutation of SEQ ID NO: 4273, or one or more corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
140. The fusion protein of any one of claims 128-136, wherein the dCas9 domain comprises one or more of a D1134E, R1334Q, and T1336R mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
141. The fusion protein of any one of claims 128-136, wherein the dCas9 domain comprises one or more of a D1134V, R1334Q, and T1336R mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
142. The fusion protein of any one of claims 128-136, wherein the dCas9 domain comprises one or more of a D1134V, G1217R, R1334Q, and T1336R mutation of SEQ ID NO: 4276, or a corresponding mutation in any of the amino acid sequences provided in SEQ ID NOs: 11-260.
143. The fusion protein of any of claims 128-142, wherein the nucleic acid editing domain is fused to the N-terminus of the Cas9 nickase domain.
144. The fusion protein of any one of claims 128-143, wherein the UGI domain is fused to the C-terminus of the Cas9 nickase domain.
145. The fusion protein of any one of claims 128-144, wherein the Cas9 nickase domain and the nucleic acid editing domain are fused via a linker.
146. The fusion protein of any one of claims 128-145, wherein the Cas9 nickase domain and the UGI domain are fused via a linker.

147. The fusion protein of claims 145 or 146, wherein the linker comprises the amino acid sequence (GGGS)_n (SEQ ID NO: 5), (G)_n, (EAAAK)_n (SEQ ID NO: 6), (GGS)_n, SGSETPGTSESATPES (SEQ ID NO: 7), SGGS (SEQ ID NO: 4288), (XP)_n, or any combination thereof, wherein n is independently an integer between 1 and 30, and wherein X is any amino acid.
148. The fusion protein of claim 145 or 146, wherein the linker comprises a covalent bond.
149. The fusion protein of claim 147, wherein the linker comprises the amino acid sequence (GGS)_n, wherein n is 1, 3, or 7.
150. The fusion protein of claim 147, wherein the linker comprises the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).
151. The fusion protein of claim 145, wherein the nCas9 domain and the nucleic acid editing domain are fused via a linker comprising the amino acid sequence SGSETPGTSESATPES (SEQ ID NO: 7).
152. The fusion protein of claim 145, wherein the nCas9 domain and the nucleic acid editing domain are fused via a linker comprising the amino acid sequence (GGS)_n, wherein n is 1, 3, or 7.
153. The fusion protein of claim 146, wherein the nCas9 domain and the UGI domain are fused via a linker comprising the amino acid sequence (GGGS)_n (SEQ ID NO: 5), (G)_n, (EAAAK)_n (SEQ ID NO: 6), (GGS)_n, SGSETPGTSESATPES (SEQ ID NO: 7), SGGS (SEQ ID NO: 4288), (XP)_n, or any combination thereof, wherein n is independently an integer between 1 and 30, and wherein X is any amino acid.
154. The fusion protein of claim 146, wherein the nCas9 domain and the UGI domain are fused via a linker comprising the amino acid sequence SGGS (SEQ ID NO: 4288).
155. The fusion protein of any one of claims 128-154 wherein the fusion protein comprises the structure [nucleic acid editing domain]-[optional linker]-[Cas9 nickase]-[optional linker]-[UGI domain].

156. The fusion protein of any one of claims 128-154 wherein the fusion protein comprises the structure [nucleic acid editing domain]-[optional linker]-[UGI domain]-[optional linker]-[Cas9 nickase]; [UGI domain]-[optional linker]-[nucleic acid editing domain]-[optional linker]-[Cas9 nickase]; [UGI domain]-[optional linker]-[Cas9 nickase]-[optional linker]-[nucleic acid editing domain]; [Cas9 nickase]-[optional linker]-[UGI domain]-[optional linker]-[nucleic acid editing domain]; or [Cas9 nickase]-[optional linker]-[nucleic acid editing domain]-[optional linker]-[UGI domain].

157. The fusion protein of any one of claims 128-156, wherein the nucleic acid editing domain comprises a deaminase.

158. The fusion protein of claim 157 wherein the deaminase is a cytidine deaminase.

159. The fusion protein of claim 157 or 158, wherein the deaminase is an apolipoprotein B mRNA-editing complex (APOBEC) family deaminase.

160. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC1 deaminase.

161. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC2 deaminase.

162. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC3A deaminase.

163. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC3B deaminase.

164. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC3C deaminase.

165. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC3D deaminase.

166. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC3F deaminase.
167. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC3G deaminase.
168. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC3H deaminase.
169. The fusion protein of any one of claims 157-159, wherein the deaminase is an APOBEC4 deaminase.
170. The fusion protein of claim 157 or 158, wherein the deaminase is an activation-induced deaminase (AID).
171. The fusion protein of claim 157 or 158, wherein the deaminase is an APOBEC deaminase comprising one or more mutations selected from the group consisting of H121R, H122R, R126A, R126E, R118A, W90A, W90Y, and R132E of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase.
172. The fusion protein of claim 157 or 158, wherein the deaminase is an APOBEC deaminase comprising a W90Y, a R126E, and a R132E mutation of rAPOBEC1 (SEQ ID NO: 284), or one or more corresponding mutations in another APOBEC deaminase.
173. The fusion protein of claim 157 or 158, wherein the deaminase comprises one or more mutations selected from the group consisting of D316R, D317R, R320A, R320E, R313A, W285A, W285Y, R326E of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase.
174. The fusion protein of claim 157 or 158, wherein the deaminase is an APOBEC deaminase comprising a W285Y, a R320E, and a R326E mutation of hAPOBEC3G (SEQ ID NO: 275), or one or more corresponding mutations in another APOBEC deaminase.

175. The fusion protein of any one of claims 157-174, wherein the deaminase is from a human, chimpanzee, gorilla, monkey, cow, dog, rat, or mouse.
176. The fusion protein of any one of claims 157-175, wherein the deaminase is from a human.
177. The fusion protein of any one of claims 157-175, wherein the deaminase is from a rat.
178. The fusion protein of claim 157 or 158, wherein the deaminase is an cytidine deaminase 1 from *Petromyzon marinus* (pmCDA1).
179. The fusion protein of any one of claims 157-159, wherein the deaminase is a rat APOBEC1 deaminase comprising the amino acid sequence set forth in (SEQ ID NO: 284).
180. The fusion protein of any one of claims 157-159, wherein the deaminase is a human APOBEC1 deaminase comprising the amino acid sequence set forth in (SEQ ID NO: 282).
181. The fusion protein of claim 178, wherein the pmCDA1 comprises an amino acid sequence set forth in (SEQ ID NO: 5738).
182. The fusion protein of claim 167, wherein the APOBEC3G is a human APOBEC3G comprising the amino acid sequence set forth in (SEQ ID NO: 275).
183. The fusion protein of claim 167, wherein the APOBEC3G is a human APOBEC3G variant comprising the amino acid sequence set forth in any one of (SEQ ID NO: 5739-5741).
184. The fusion protein of claim 157 or 158, wherein the deaminase is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of the amino acid sequences set forth in SEQ ID NOs: 266-284, 607-610, 5724-5736 and 5738-5741.
185. The fusion protein of claim 157 or 158, wherein the deaminase comprises the amino acid sequence set forth in any one of SEQ ID NOs: 266-284, 607-610, 5724-5736 and 5738-5741.

186. The fusion protein of any one of claims 128-185, wherein the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to SEQ ID NO: 600.

187. The fusion protein of any one of claims 128-186, wherein the UGI domain comprises the amino acid sequence as set forth in SEQ ID NO: 600.

188. The fusion protein of any one of claims 128-185, wherein the UGI domain comprises an amino acid sequence that is at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or at least 99.5% identical to any one of SEQ ID NOs: 322-324.

189. The fusion protein of any one of claims 128- 185, wherein the UGI domain comprises the amino acid sequence as set forth in any one of SEQ ID NOs: 322-324.

190. A complex comprising the fusion protein of anyone of claims 1-30, and a guide RNA (gRNA) bound to the Cas9 domain of the fusion protein.

191. A complex comprising the fusion protein of any one of claims 31-106, and a guide RNA (gRNA) bound to the dCas9 domain of the fusion protein.

192. A complex comprising the fusion protein of any one of claims 107-189 and a guide RNA (gRNA) bound to the Cas9 nickase (nCas9) domain of the fusion protein.

193. The complex of any one of claims 190-192, wherein the guide RNA is from 15-100 nucleotides long and comprises a sequence of at least 10 contiguous nucleotides that is complementary to a target sequence.

194. The complex of claim 193, wherein the guide RNA is 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, or 50 nucleotides long.

195. The complex of any one of claims 190-194, wherein the guide RNA comprises a sequence of 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, or 40 contiguous nucleotides that is complementary to a target sequence.
196. The complex of any one of claims 190-195, wherein the target sequence is a DNA sequence.
197. The complex of claim 196, wherein the target sequence is in the genome of an organism.
198. The complex of claim 197, wherein the organism is a prokaryote.
199. The complex of claim 198, wherein the prokaryote is bacteria.
200. The complex of claim 197, wherein the organism is a eukaryote.
201. The complex of claim 200, wherein the organism is a plant.
202. The complex of claim 200, wherein the organism is a vertebrate.
203. The complex of claim 202, wherein the vertebrate is a mammal.
204. The complex of claim 203, wherein the mammal is a mouse or rat.
205. The complex of claim 203, wherein the mammal is human.
206. A method comprising contacting a nucleic acid molecule with the fusion protein of any one of claims 1-189 and a guide RNA, wherein the guide RNA is from 15-100 nucleotides long and comprises a sequence of at least 10 contiguous nucleotides that is complementary to a target sequence.
207. A method comprising contacting a nucleic acid molecule with the complex of any one of claims 190-205.

208. The method of claim 206 or 207, wherein the nucleic acid is DNA.
209. The method of claim 208, wherein the nucleic acid is double-stranded DNA.
210. The method of any one of claims 206-209, wherein the target sequence comprises a sequence associated with a disease or disorder.
211. The method of claim 210, wherein the target sequence comprises a point mutation associated with a disease or disorder.
212. The method of claim 211, wherein the activity of the fusion protein, or the complex results in a correction of the point mutation.
213. The method of any one of claims 206-212, wherein the target sequence comprises a T to C point mutation associated with a disease or disorder, and wherein the deamination of the mutant C base results in a sequence that is not associated with a disease or disorder.
214. The method of claim 213, wherein the target sequence encodes a protein, and wherein the point mutation is in a codon and results in a change in the amino acid encoded by the mutant codon as compared to a wild-type codon.
215. The method of claim 214, wherein the deamination of the mutant C results in a change of the amino acid encoded by the mutant codon.
216. The method of claim 215, wherein the deamination of the mutant C results in the codon encoding a wild-type amino acid.
217. The method of any one of claims 206-216, wherein the contacting is performed *in vivo* in a subject.
218. The method of any one of claims 206-216, wherein the contacting is performed *in vitro*.

219. The method of claim 217, wherein the subject has been diagnosed with a disease or disorder.

220. The method of any one of claims 210-219, wherein the disease or disorder is cystic fibrosis, phenylketonuria, epidermolytic hyperkeratosis (EHK), Charcot-Marie-Toot disease type 4J, neuroblastoma (NB), von Willebrand disease (vWD), myotonia congenital, hereditary renal amyloidosis, dilated cardiomyopathy (DCM), hereditary lymphedema, familial Alzheimer's disease, HIV, Prion disease, chronic infantile neurologic cutaneous articular syndrome (CINCA), desmin-related myopathy (DRM), a neoplastic disease associated with a mutant PI3KCA protein, a mutant CTNNB1 protein, a mutant HRAS protein, or a mutant p53 protein.

221. The method of any one of claims 211-220, wherein the disease or disorder is associated with a T>C or A>G mutation in a gene selected from the genes disclosed in Table 1.

222. The method of any one of claims 211-220, wherein the disease or disorder is associated with a T>C or A>G mutation in a gene selected from the genes disclosed in Table 2 or 3.

223. The method of any one of claims 206-222, wherein the guide RNA comprises a nucleotide sequence of any one of the protospacer sequences in Table 2 or Table 3.

224. A method for editing a nucleobase pair of a double-stranded DNA sequence, the method comprising:

- a. contacting a target region of the double-stranded DNA sequence with a complex comprising a nucleobase editor and a guide nucleic acid, wherein the target region comprises a target nucleobase pair;
- b. inducing strand separation of said target region;
- c. converting a first nucleobase of said target nucleobase pair in a single strand of the target region to a second nucleobase; and
- d. cutting no more than one strand of said target region;

wherein a third nucleobase complementary to the first nucleobase base is replaced by a fourth nucleobase complementary to the second nucleobase and the method causes less than 20% indel formation in the double-stranded DNA sequence.

225. The method of claim 224, wherein the method causes less than 20%, 19%, 18%, 16%, 14%, 12%, 10%, 8%, 6%, 4%, 2%, or 1% indel formation.

226. The method of claim 224 or 225, further comprising replacing the second nucleobase with a fifth nucleobase that is complementary to the fourth nucleobase, thereby generating an intended edited basepair.

227. The method of any one of claims 224-226, wherein the efficiency of generating the intended edited basepair is at least 5%.

228. The method of claim 227, wherein the efficiency is at least 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, or 50%.

229. The method of claim 226, wherein the ratio of intended products to unintended products at the target nucleotide is at least 2:1, 5:1, 10:1, 20:1, 30:1, 40:1, 50:1, 60:1, 70:1, 80:1, 90:1, 100:1, or 200:1.

230. The method of claim 226, wherein the ratio of intended point mutation to indel formation is greater than 1:1, 10:1, 50:1, 100:1, 500:1, or 1000:1.

231. The method of any one of claims 224-230, wherein the cut single strand is hybridized to the guide nucleic acid.

232. The method of any one of claims 224-231, wherein the cut single strand is opposite to the strand comprising the first nucleobase.

233. The method of any one of claims 224-232, wherein said first base is cytosine.

234. The method of any one of claims 224-233, wherein the second nucleobase is not a G, C, A, or T.

235. The method of any one of claims 224-234, wherein said second base is uracil.
236. The method of any one of claims 224-235, wherein the nucleobase editor comprises UGI activity.
237. The method of any one of claims 224-236, wherein the nucleobase editor comprises nickase activity.
238. The method of any one of claims 226-237, wherein the intended edited basepair is upstream of a PAM site.
239. The method of claim 238, wherein the intended edited base pair is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides upstream of the PAM site.
240. The method of claim 239, wherein the intended edited basepair is downstream of a PAM site.
241. The method of claim 240, wherein the intended edited base pair is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides downstream stream of the PAM site.
242. The method of any one of claims 224-241, wherein the method does not require a canonical PAM site.
243. The method of claim 242, wherein the canonical PAM sit comprises NGG, wherein N is A, T, C, or G.
244. The method of any one of claims 224-243, wherein the nucleobase editor comprises a linker.
245. The method of claim 244, wherein the linker is 1-25 amino acids in length.
246. The method of claim 244 or 245, wherein the linker is 5-20 amino acids in length.

247. The method of any one of claims 244-246, wherein the linker is 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 amino acids in length.
248. The method of any one of claims 224-247, wherein the target region comprises a target window, wherein the target window comprises the target nucleobase pair.
249. The method of claim 248, wherein the target window comprises 1-10 nucleotides.
250. The method of claim 248, wherein the target window is 1-9, 1-8, 1-7, 1-6, 1-5, 1-4, 1-3, 1-2, or 1 nucleotides in length.
251. The method of claim 248, wherein the target window is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides in length.
252. The method of claim any one of claims 224-251, wherein the intended edited base pair occurs within the target window.
253. The method of claim any one of claims 224-252, wherein the target window comprises the intended edited base pair.
254. The method of any one of claims 224-253, wherein the nucleobase editor comprises any one of the fusion proteins of claims 1-189
255. A method for editing a nucleobase pair of a double-stranded DNA sequence, the method comprising:
- a. contacting a target region of the double-stranded DNA sequence with a complex comprising a nucleobase editor and a guide nucleic acid, wherein the target region comprises a target nucleobase pair;
 - b. inducing strand separation of said target region;
 - c. converting a first nucleobase of said target nucleobase pair in a single strand of the target region to a second nucleobase;
 - d. cutting no more than one strand of said target region;

wherein a third nucleobase complementary to the first nucleobase base is replaced by a fourth nucleobase complementary to the second nucleobase; and

- e. replacing the second nucleobase with a fifth nucleobase that is complementary to the fourth nucleobase, thereby generating an intended edited basepair,

wherein the efficiency of generating the intended edited basepair is at least 5%.

256. The method of claim 255, wherein the efficiency is at least 5%, 10%, 15%, 20%, 25%, 30%, 35%, 40%, 45%, or 50%..

257. The method of claim 255 or 256, wherein the method causes less than 19%, 18%, 16%, 14%, 12%, 10%, 8%, 6%, 4%, 2%, or 1% indel formation.

258. The method of any one of claims 255-257, wherein the ratio of intended product to unintended products at the target nucleotide is at least 2:1, 5:1, 10:1, 20:1, 30:1, 40:1, 50:1, 60:1, 70:1, 80:1, 90:1, 100:1, or 200:1.

259. The method of any one of claims 255-258, wherein the ratio of intended point mutation to indel formation is greater than 1:1, 10:1, 50:1, 100:1, 500:1, or 1000:1.

260. The method of any one of claims 255-259, wherein the cut single strand is hybridized to the guide nucleic acid.

261. The method of claim any one of claims 255-260, wherein the cut single strand is opposite to the strand comprising the first nucleobase.

262. The method of any one of claims 255-261, wherein said first base is cytosine.

263. The method of any one of claims 255-262, wherein the second nucleobase is not G, C, A, or T.

264. The method of any one of claims 255-263, wherein said second base is uracil.

265. The method of any one of claims 255-264, wherein the nucleobase editor comprises UGI activity.

266. The method of any one of claims 255-265, wherein the nucleobase edit comprises nickase activity.
267. The method of any one of claims 255-266, wherein the intended edited basepair is upstream of a PAM site.
268. The method of claim 267, wherein the intended edited base pair is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides upstream of the PAM site.
269. The method of any one of claims 255-266, wherein the intended edited basepair is downstream of a PAM site.
270. The method of claim 269, wherein the intended edited base pair is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides downstream stream of the PAM site.
271. The method of any one of claims 255-270, wherein the method does not require a canonical PAM site.
272. The method of claim 271, wherein the canonical PAM site comprises NGG, wherein N is A, T, C, or G.
273. The method of any one of claims 255-272, wherein the nucleobase editor comprises a linker.
274. The method of claim 273, wherein the linker is 1-25 amino acids in length.
275. The method of claim 274 or 275, wherein the linker is 5-20 amino acids in length.
276. The method of any one of claims 274-275, wherein the linker is 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 amino acids in length.

277. The method of any one of claims 274-27, wherein the target region comprises a target window, wherein the target window comprises the target nucleobase pair.
278. The method of claim 277 wherein the target window comprises 1-10 nucleotides.
279. The method of claim 277, wherein the target window is 1-9, 1-8, 1-7, 1-6, 1-5, 1-4, 1-3, 1-2, or 1 nucleotides in length.
280. The method of claim 277, wherein the target window is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, or 20 nucleotides in length.
281. The method of any one of claims 277- 280, wherein the intended edited base pair occurs within the target window.
282. The method of any one of claims 277- 281, wherein the target window comprises the intended edited base pair.
283. The method of any one of claims 255-282, wherein the nucleobase editor comprises any one of the fusion proteins of claims 1-189
284. A nucleic acid-guided deaminase coupled to an inhibitor of base excision repair.
285. The nucleic acid-guided deaminase of claim 284 comprising an initiator of mismatch repair.
286. The nucleic acid-guided deaminase of claim 284 comprising a nickase.
287. A method for editing a nucleobase pair of a double-stranded DNA sequence, the method comprising:
- contacting a target region of the double-stranded DNA sequence with a nucleic acid-guided deaminase, wherein the target region comprises a target nucleobase pair;
 - converting a first nucleobase of said target nucleobase pair of the target region to a second nucleobase; and
 - inhibiting base excision repair of the second nucleobase.

288. The method of claim 287 further comprising nicking the non-edited strand of the target double-stranded DNA sequence.

289. The method of claim 287 further comprising initiating mismatch repair to convert the nucleobase complementary to the first nucleobase on the non-edited strand to a nucleobase complementary to the second nucleobase.

290. The method of claim 287 further comprising inducing strand separation in the target region.

291. A method for editing a nucleobase pair of a double-stranded DNA sequence, the method comprising:

- a. contacting a target region of the double-stranded DNA sequence with a nucleic acid-guided deaminase, wherein the target region comprises a target nucleobase pair;
- b. converting a first nucleobase of said target nucleobase pair in the target region to a second nucleobase; and
- c. initiating mismatch repair to convert the nucleobase complementary to the first nucleobase on the non-edited strand to a nucleobase complementary to the second nucleobase.

292. The method of claim 291 further comprising inhibiting base excision repair of the second nucleobase.

293. The method of claim 291 further comprising inducing strand separation in the target region.

294. The method of claim 287 or 291, wherein the nucleic acid-guided deaminase is a nucleic acid-guided cytidine deaminase.

295. A kit comprising a nucleic acid construct, comprising
(a) a nucleic acid sequence encoding the fusion protein of any one of claims 1-189;
and
(b) a heterologous promoter that drives expression of the sequence of (a).

296. The kit of claim 256, further comprising an expression construct encoding a guide RNA backbone, wherein the construct comprises a cloning site positioned to allow the cloning of a nucleic acid sequence identical or complementary to a target sequence into the guide RNA backbone.
297. A polynucleotide encoding the fusion protein of any one of claims 1-189.
298. A vector comprising a polynucleotide of claim 258.
299. The vector of claim 259, wherein the vector comprises a heterologous promoter driving expression of the polynucleotide.
300. A cell comprising the fusion protein of any one of claims 1-189.
301. A cell comprising the complex of any of claims 190-205.
302. A cell comprising the nucleic acid molecule encoding the fusion protein of any one of claims 1-189.

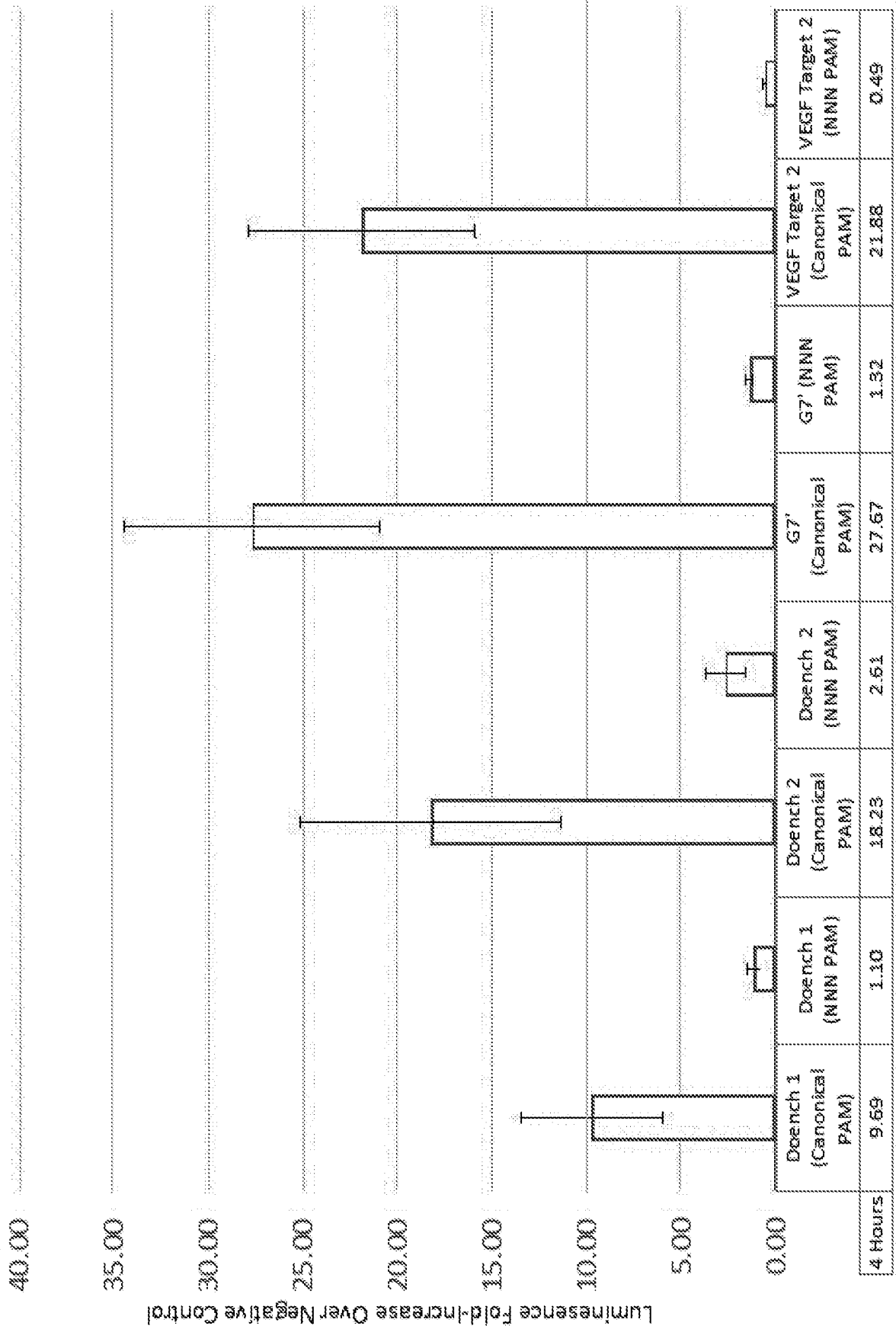


FIGURE 1

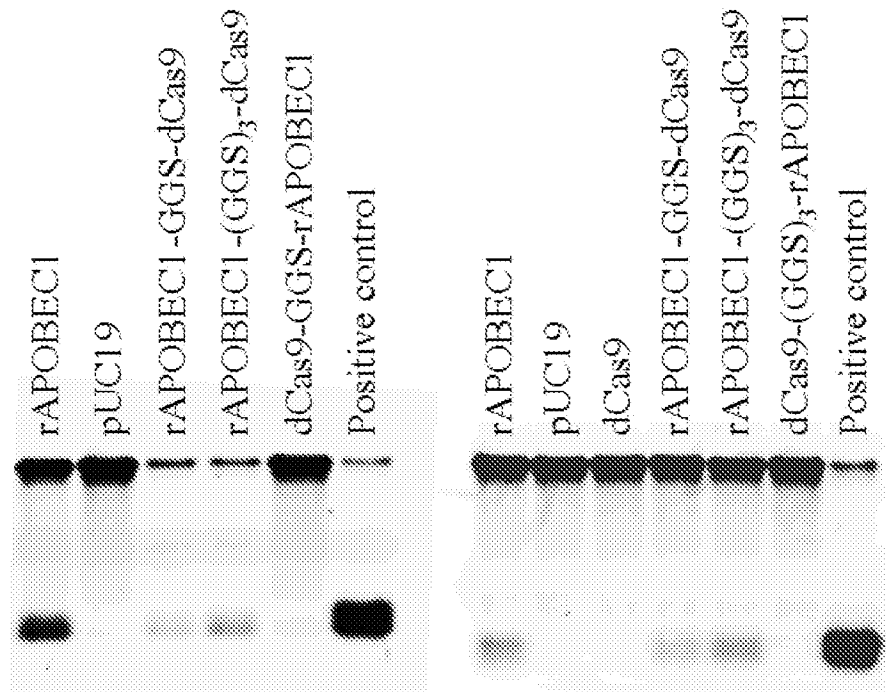


FIGURE 2

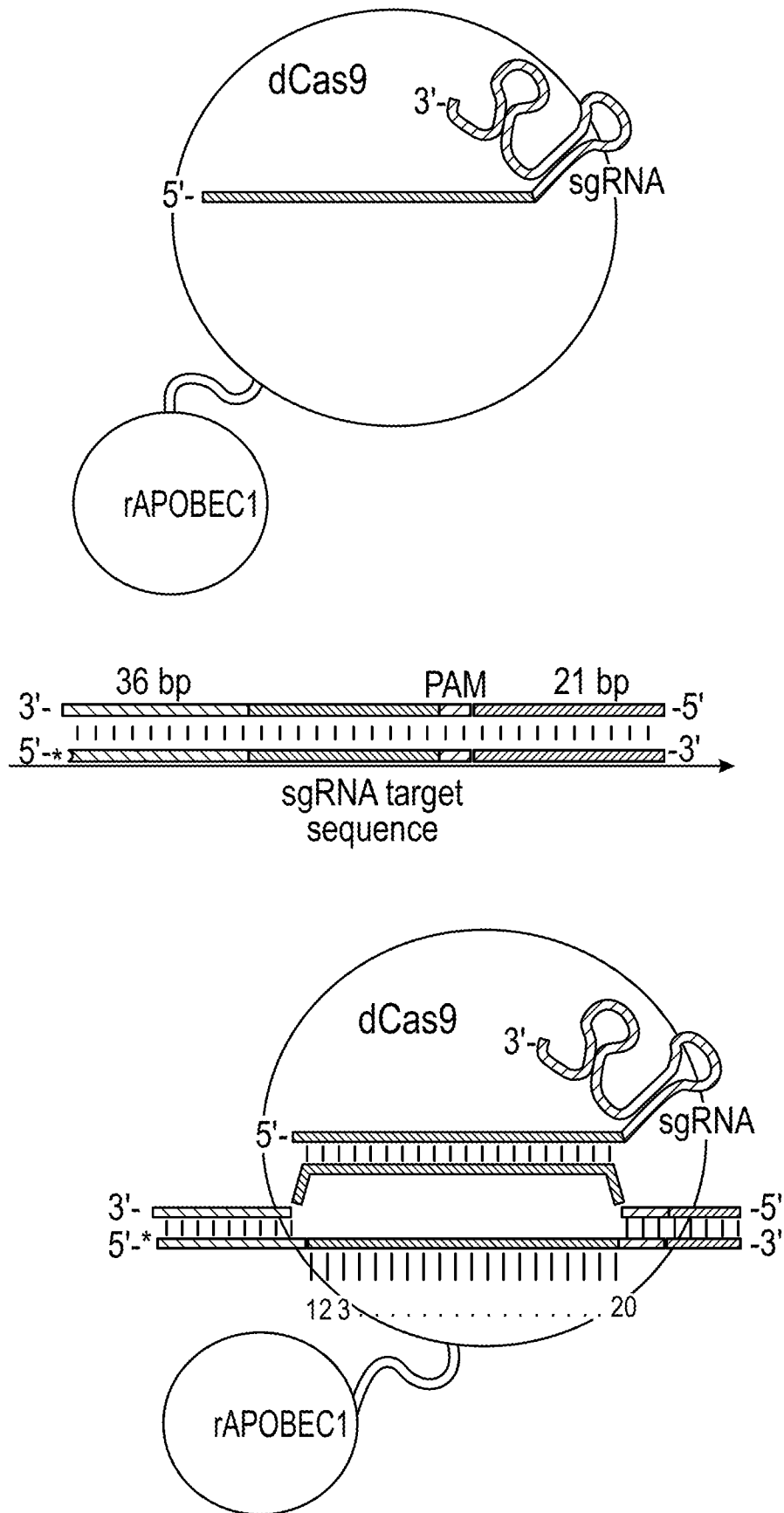


FIGURE 3

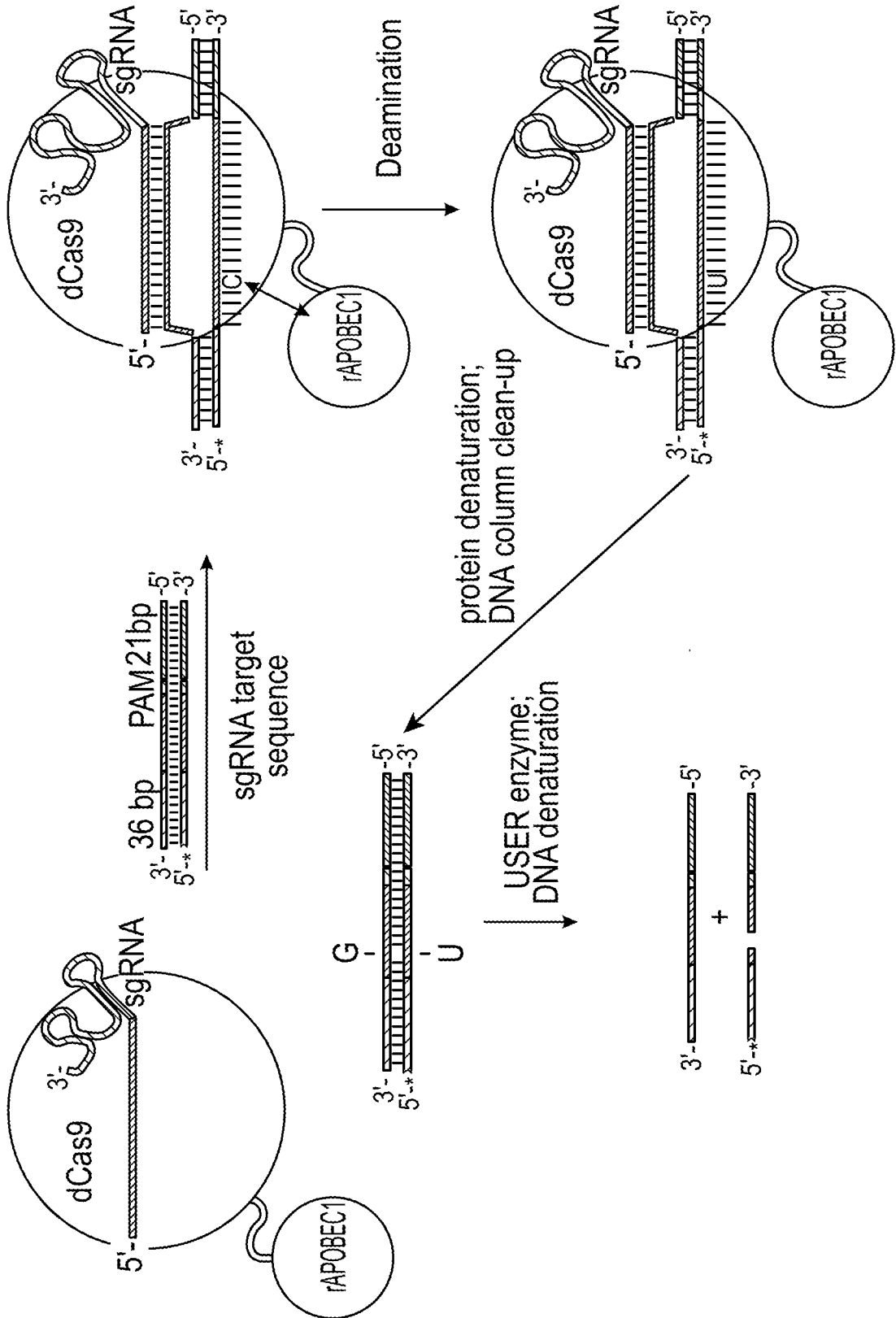


FIGURE 4

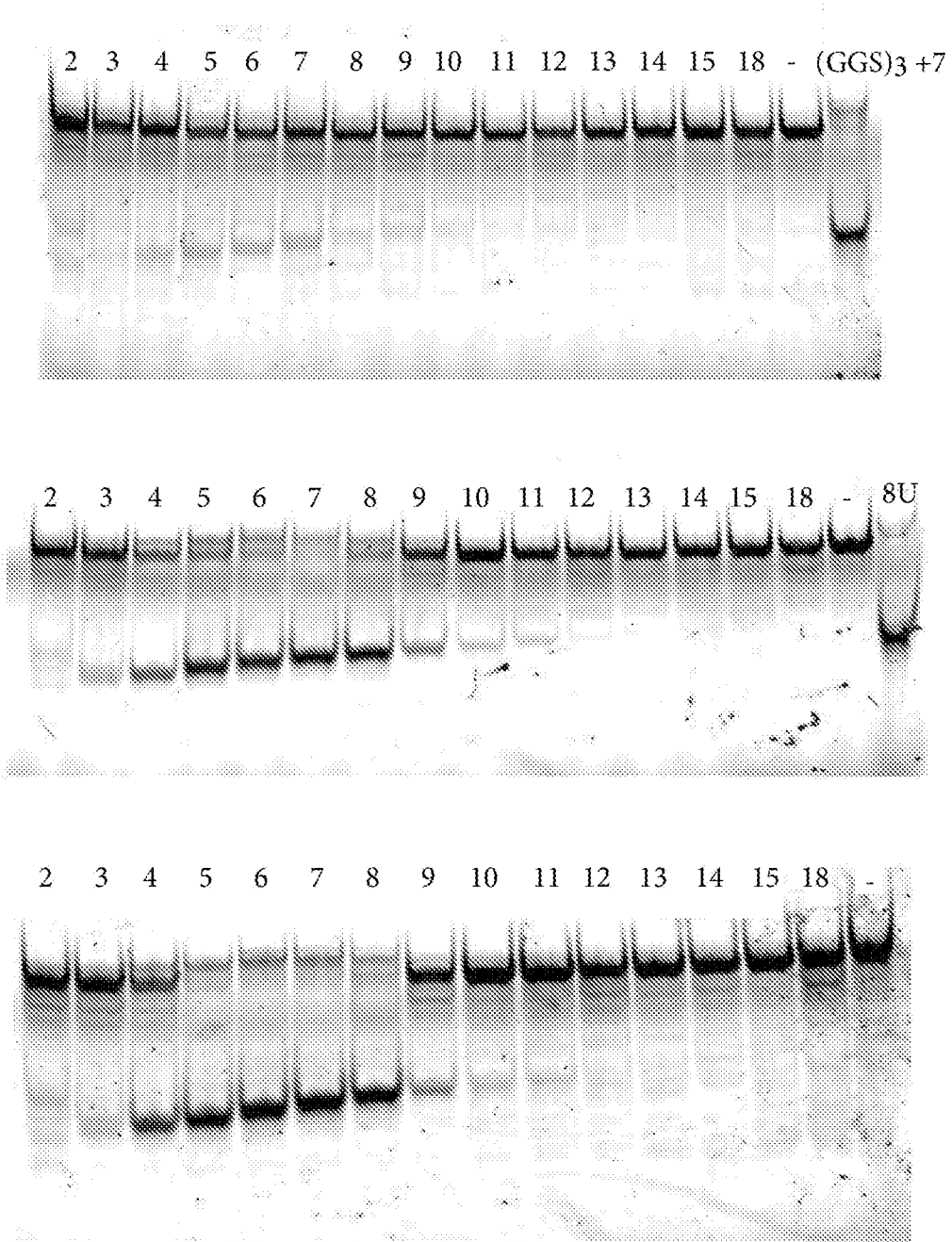


FIGURE 5

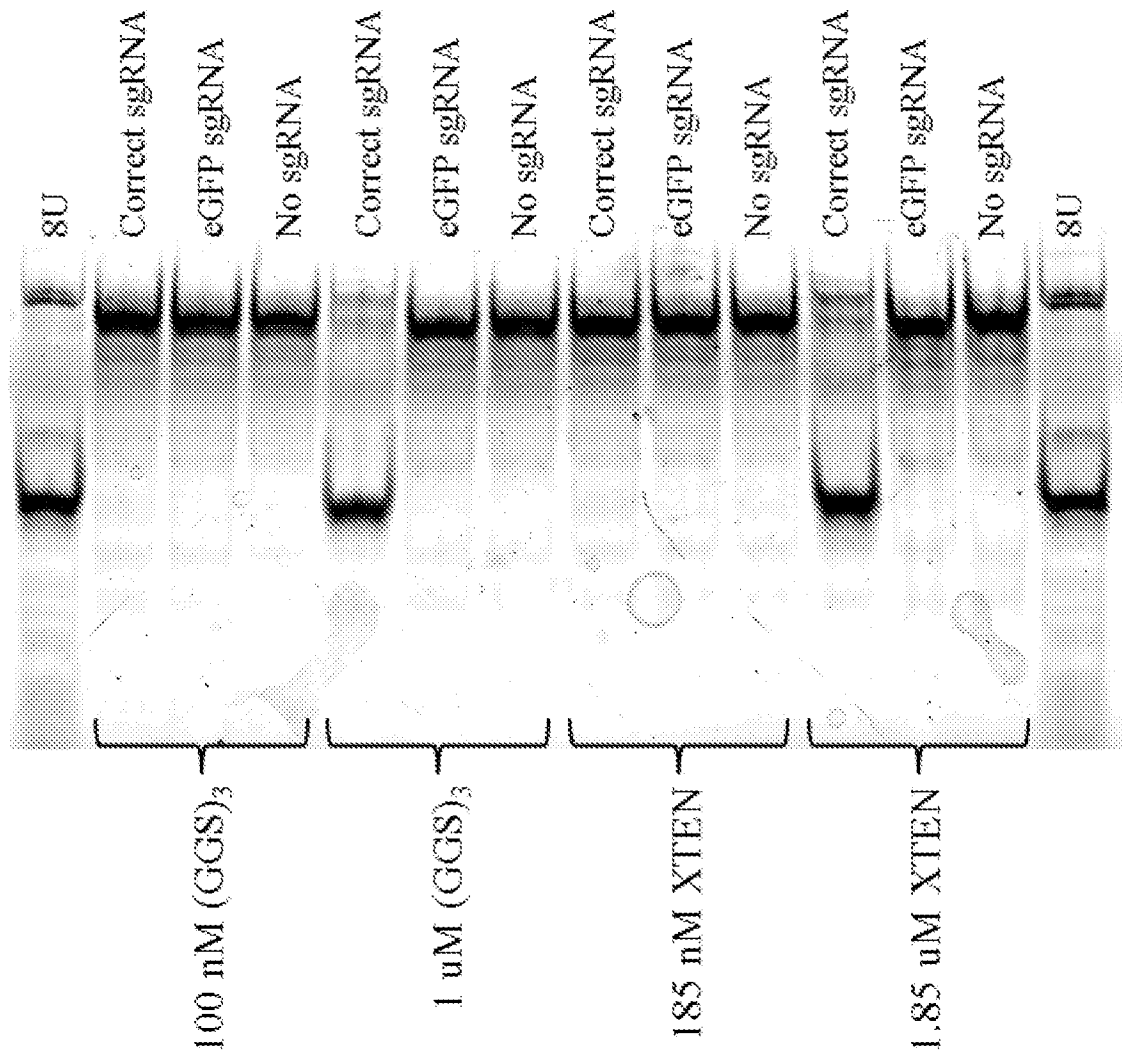


FIGURE 6

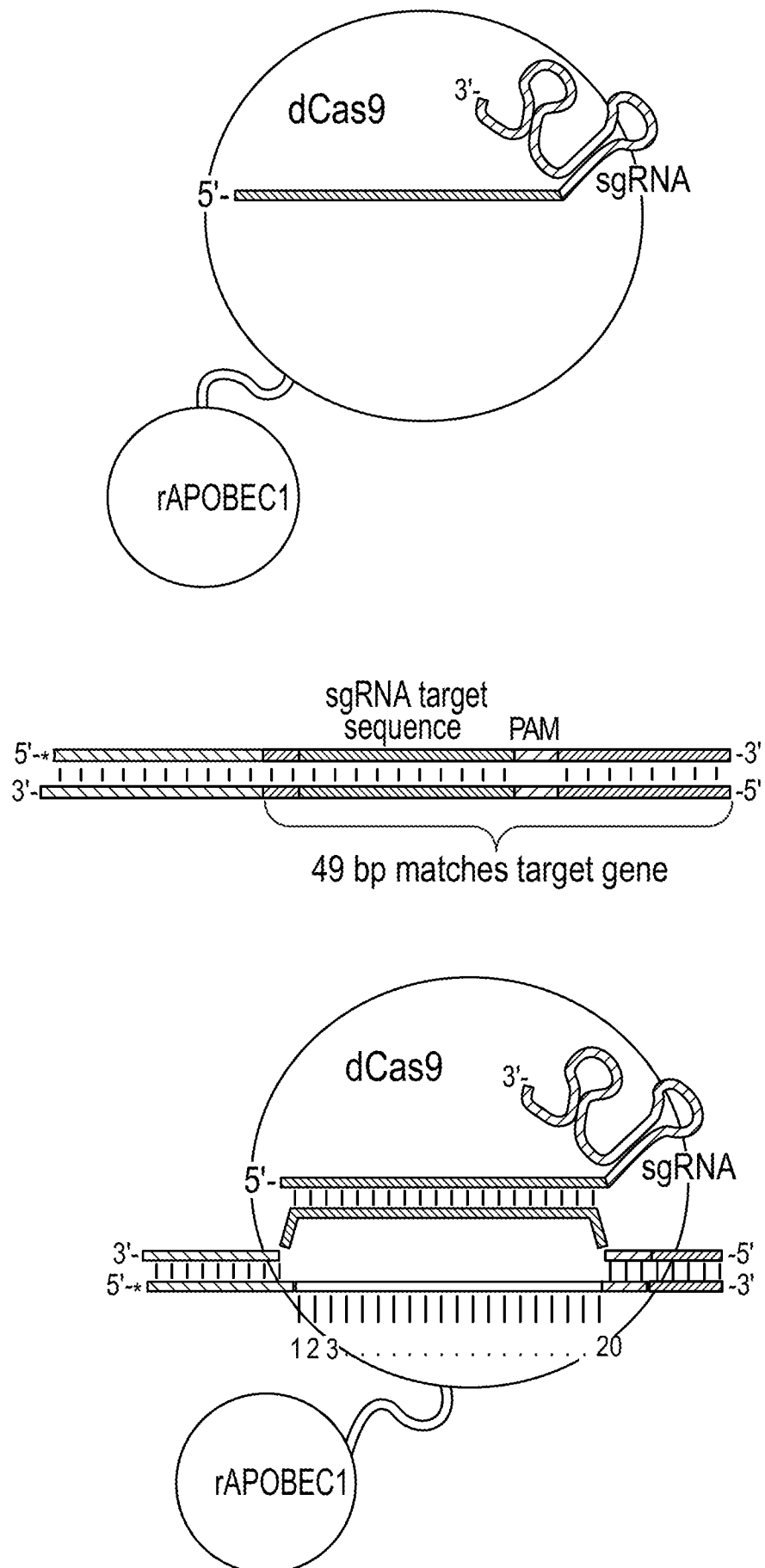


FIGURE 7

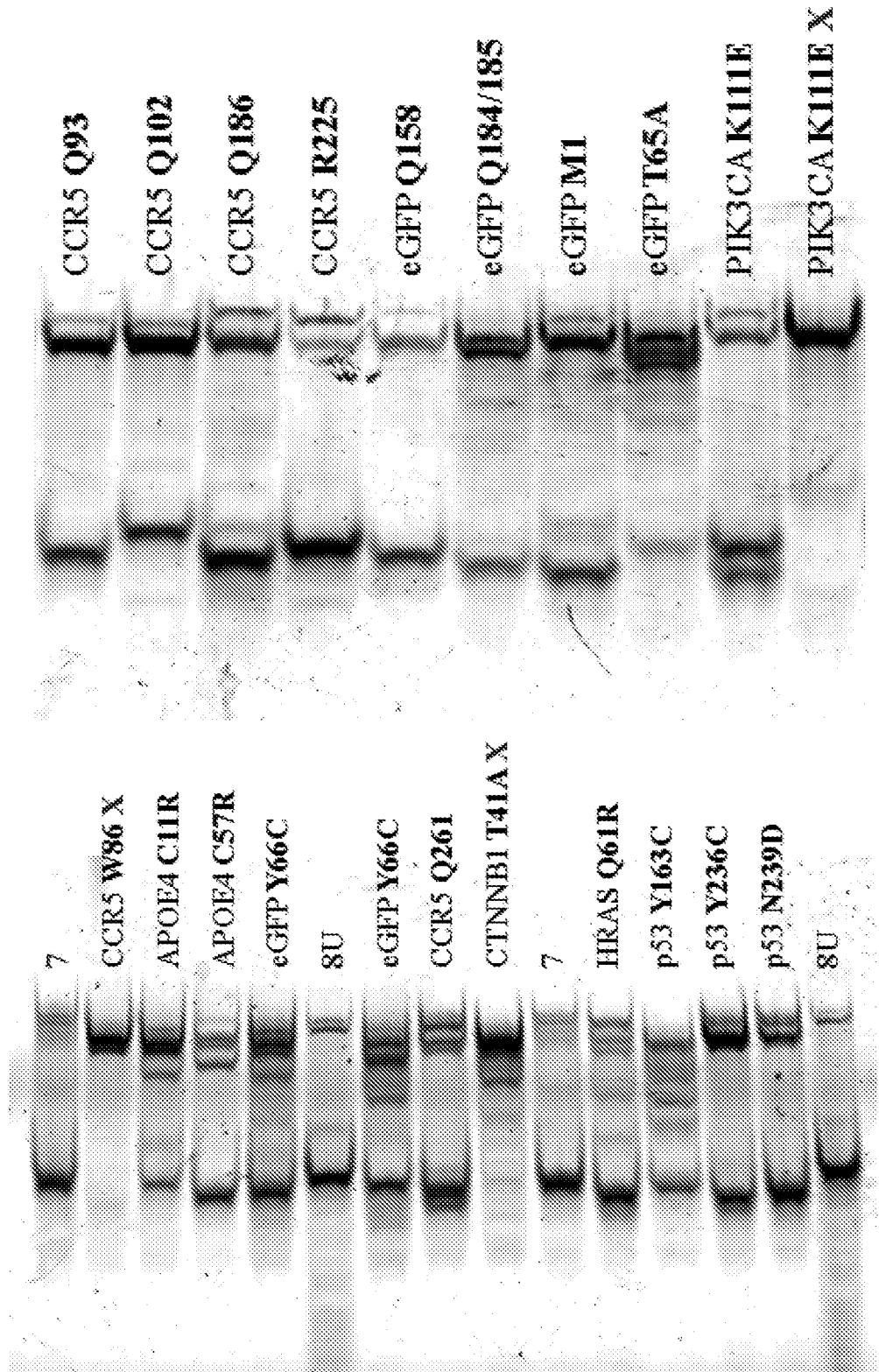


FIGURE 8

EMX1		C ₅	C ₆	C ₁₀
0 μ M XTEN	A	0.1%	0.1%	0.1%
	C	99.8%	99.8%	99.8%
	G	0.0%	0.0%	0.1%
	T	0.0%	0.1%	0.0%
1.85 μ M XTEN	A	0.1%	0.0%	0.1%
	C	60.4%	61.0%	99.1%
	G	0.0%	0.0%	0.1%
	T	39.5%	39.0%	0.7%

FANCF		C ₆	C ₇	C ₈	C ₁₁
0 μ M XTEN	A	0.1%	0.1%	0.1%	0.1%
	C	99.8%	99.8%	99.9%	99.9%
	G	0.0%	0.1%	0.0%	0.0%
	T	0.0%	0.0%	0.0%	0.0%
1.85 μ M XTEN	A	0.1%	0.1%	0.1%	0.1%
	C	63.9%	64.7%	65.0%	72.6%
	G	0.0%	0.0%	0.0%	0.0%
	T	36.0%	35.1%	34.9%	27.3%

HEK293 site 2		C ₄	C ₆	C ₁₁
0 μ M XTEN	A	0.1%	0.1%	0.1%
	C	99.9%	99.9%	99.9%
	G	0.0%	0.0%	0.0%
	T	0.0%	0.0%	0.1%
1.85 μ M XTEN	A	0.1%	0.1%	0.1%
	C	80.6%	76.9%	99.6%
	G	0.0%	0.0%	0.0%
	T	19.3%	22.9%	0.3%

HEK293 site 3		C ₃	C ₄	C ₅	C ₉
0 μ M XTEN	A	0.1%	0.1%	0.0%	0.1%
	C	99.8%	99.9%	99.9%	99.9%
	G	0.0%	0.0%	0.0%	0.0%
	T	0.1%	0.0%	0.0%	0.0%
1.85 μ M XTEN	A	0.1%	0.1%	0.0%	0.1%
	C	92.2%	74.8%	71.5%	96.6%
	G	0.0%	0.0%	0.0%	0.0%
	T	7.7%	25.1%	28.5%	3.3%

HEK293 site 4		C ₃	C ₅	C ₈	C ₁₁
0 μ M XTEN	A	0.1%	0.0%	0.1%	0.0%
	C	99.8%	99.9%	99.8%	99.9%
	G	0.0%	0.0%	0.0%	0.0%
	T	0.0%	0.0%	0.1%	0.0%
1.85 μ M XTEN	A	0.1%	0.1%	0.1%	0.1%
	C	98.8%	60.1%	97.0%	99.4%
	G	0.0%	0.0%	0.0%	0.0%
	T	1.1%	39.8%	2.9%	0.5%

RNF2		C ₃	C ₆
0 μ M XTEN	A	0.1%	0.0%
	C	99.9%	99.9%
	G	0.0%	0.0%
	T	0.0%	0.0%
1.85 μ M XTEN	A	0.1%	0.0%
	C	59.1%	57.8%
	G	0.0%	0.0%
	T	40.8%	42.1%

FIGURE 9

EMX1		C ₅	C ₆	C ₁₀
untreated	A	0.0%	0.0%	0.0%
	C	99.5%	99.7%	100.0%
	G	0.0%	0.1%	0.0%
	T	0.5%	0.2%	0.0%
XTEN	A	0.7%	0.5%	0.0%
	C	93.5%	95.8%	100.0%
	G	2.1%	0.3%	0.0%
	T	3.6%	3.3%	0.0%
XTEN-UGI	A	0.2%	0.0%	0.0%
	C	81.8%	82.5%	100.0%
	G	0.6%	0.3%	0.0%
	T	17.4%	17.1%	0.0%

FANCF		C ₆	C ₇	C ₈	C ₁₁
untreated	A	0.0%	0.0%	0.2%	0.1%
	C	99.9%	99.8%	99.8%	99.9%
	G	0.0%	0.0%	0.0%	0.0%
	T	0.1%	0.1%	0.0%	0.0%
XTEN	A	0.3%	0.1%	0.0%	0.0%
	C	98.1%	99.2%	99.0%	99.8%
	G	0.4%	0.0%	0.0%	0.0%
	T	1.2%	0.7%	1.0%	0.2%
XTEN-UGI	A	0.0%	0.0%	0.1%	0.0%
	C	93.2%	93.5%	93.4%	98.2%
	G	0.0%	0.0%	0.0%	0.0%
	T	6.7%	6.5%	6.5%	1.8%

HEK293 site 2		C ₄	C ₆	C ₁₁
untreated	A	0.3%	0.2%	0.2%
	C	99.7%	99.7%	99.7%
	G	0.0%	0.0%	0.0%
	T	0.0%	0.0%	0.0%
XTEN	A	0.3%	0.3%	0.3%
	C	99.7%	99.4%	99.7%
	G	0.0%	0.3%	0.0%
	T	0.0%	0.0%	0.0%
XTEN-UGI	A	0.3%	0.2%	0.2%
	C	98.8%	98.2%	99.8%
	G	0.0%	0.3%	0.0%
	T	0.9%	1.3%	0.0%

HEK293 site 2		C ₃	C ₄	C ₅	C ₉
untreated	A	0.0%	0.0%	0.0%	0.0%
	C	100.0%	100.0%	100.0%	99.9%
	G	0.0%	0.0%	0.0%	0.0%
	T	0.0%	0.0%	0.0%	0.1%
XTEN	A	0.0%	0.6%	0.3%	0.1%
	C	100.0%	95.8%	95.8%	99.2%
	G	0.0%	0.2%	0.7%	0.4%
	T	0.0%	3.4%	3.2%	0.3%
XTEN-UGI	A	0.0%	0.3%	0.3%	0.0%
	C	96.8%	83.0%	79.2%	98.5%
	G	0.0%	0.0%	1.1%	0.2%
	T	3.2%	16.8%	19.4%	1.3%

HEK293 site 4		C ₃	C ₅	C ₈	C ₁₁
untreated	A	0.0%	0.4%	0.0%	0.0%
	C	99.8%	97.6%	99.9%	100.0%
	G	0.0%	1.0%	0.0%	0.0%
	T	0.2%	1.0%	0.0%	0.0%
XTEN	A	0.0%	1.1%	0.0%	0.0%
	C	99.6%	92.2%	99.9%	100.0%
	G	0.0%	2.2%	0.0%	0.0%
	T	0.4%	4.5%	0.0%	0.0%
XTEN-UGI	A	0.0%	0.5%	0.0%	0.0%
	C	99.4%	86.7%	99.1%	100.0%
	G	0.0%	1.8%	0.0%	0.0%
	T	0.6%	11.0%	0.9%	0.0%

RNF2		C ₃	C ₆
untreated	A	0.0%	0.0%
	C	99.9%	99.5%
	G	0.0%	0.2%
	T	0.0%	0.3%
XTEN	A	0.0%	0.0%
	C	99.8%	99.3%
	G	0.0%	0.2%
	T	0.2%	0.5%
XTEN-UGI	A	0.0%	0.0%
	C	99.6%	99.1%
	G	0.0%	0.4%
	T	0.4%	0.5%

FIGURE 10

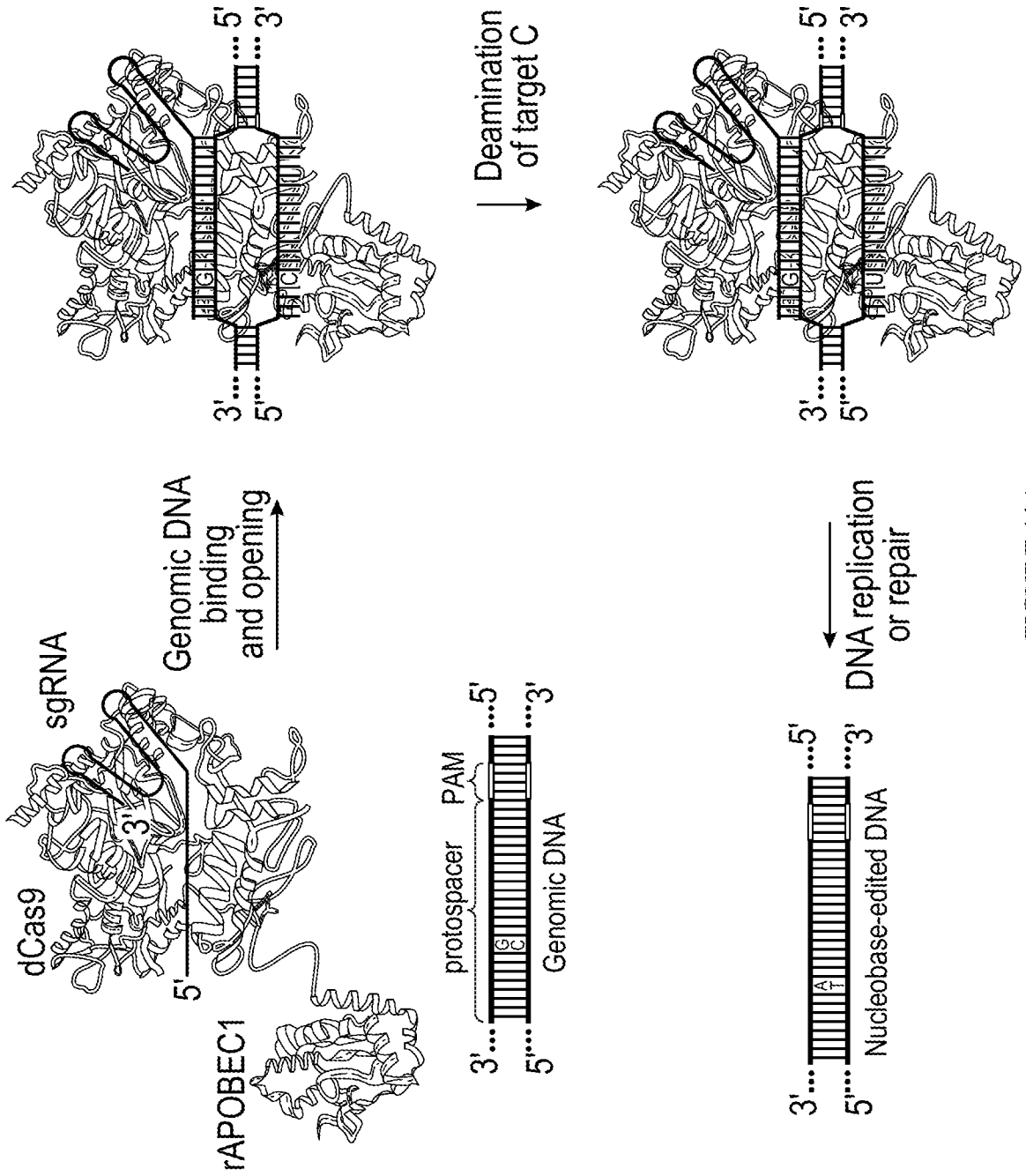


FIGURE 11A

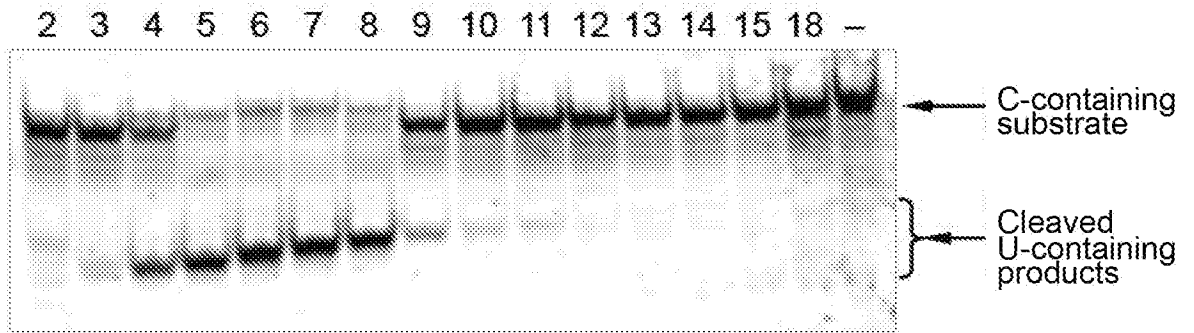


FIGURE 11B

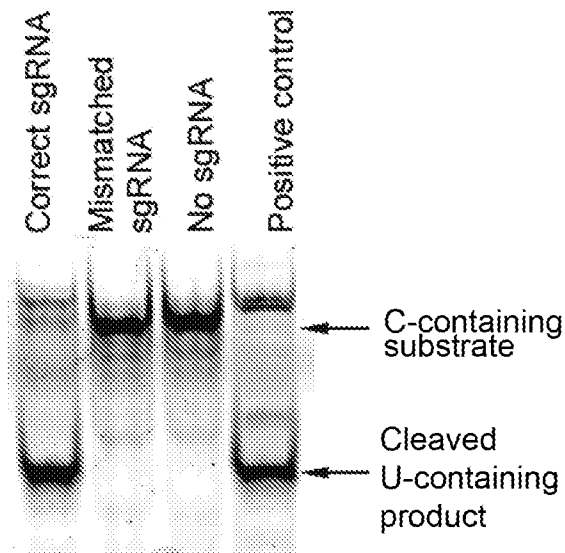


FIGURE 11C

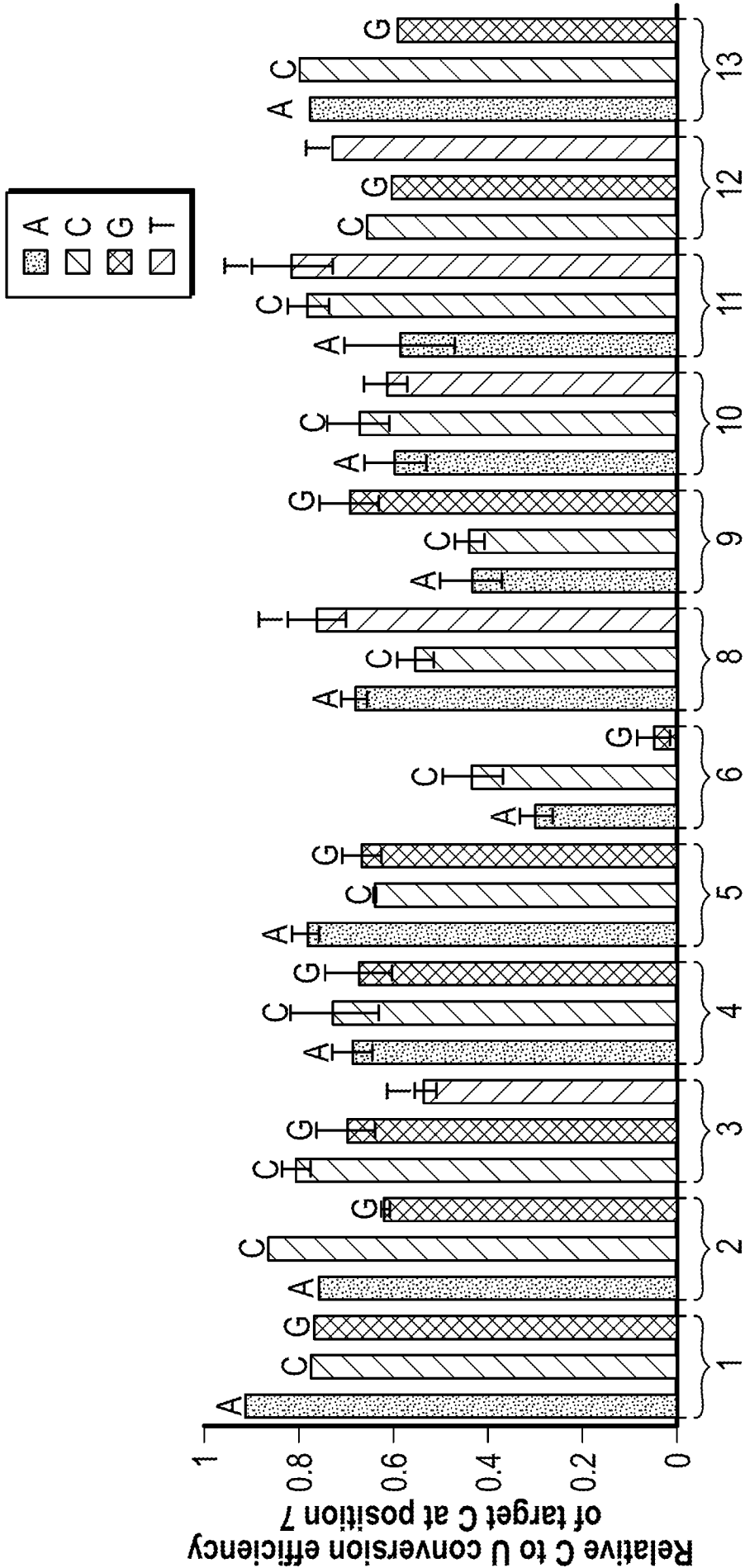


FIGURE 12A

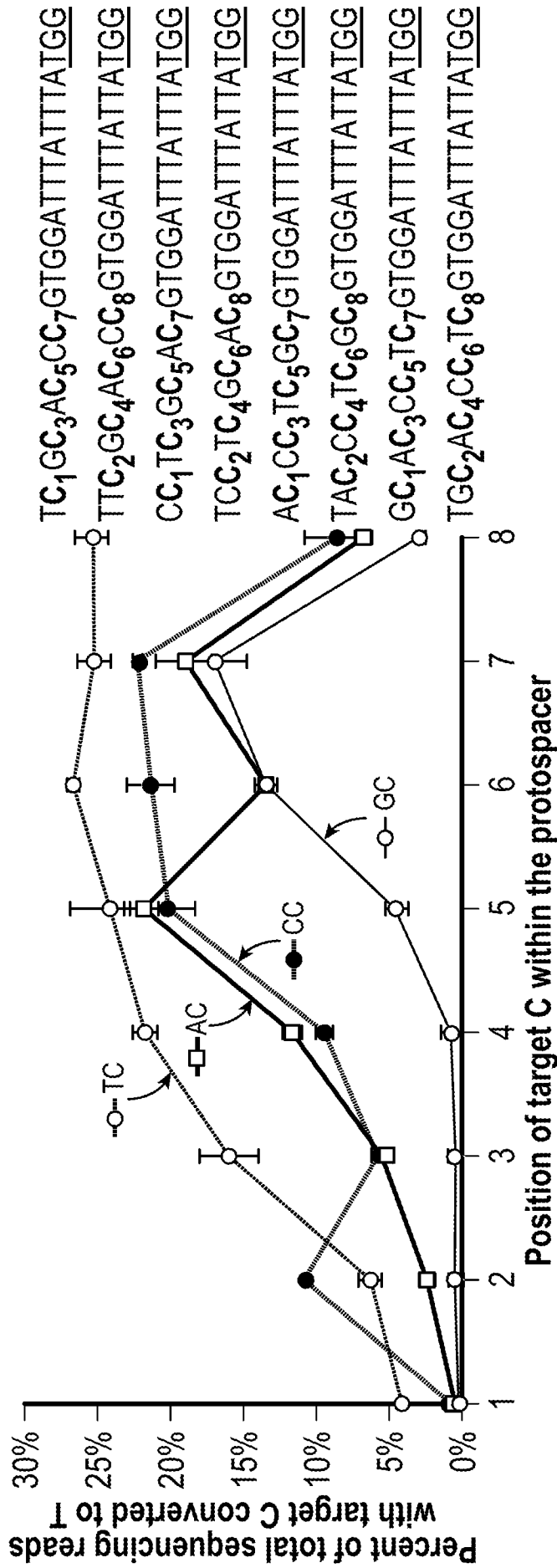


FIGURE 12B

EMX1: GAGTC₅C₆GAGCAGAAGAAGAAGGG

FANCF: GGAATC₆C₇C₈TTC₁₁TGCAGCACCTGG

HEK293 site 2: GAAC₄AC₆AAAGCATAGACTGCGGG

HEK293 site 3: GGCC₄C₅AGACTGAGCACGTGATGG

HEK293 site 4: GGCAC₅TGCGGCTGGAGGTCCGGG

RNF2: GTC₃ATC₆TTAGTC₁₂ATTACCTGAGG

FIGURE 13A

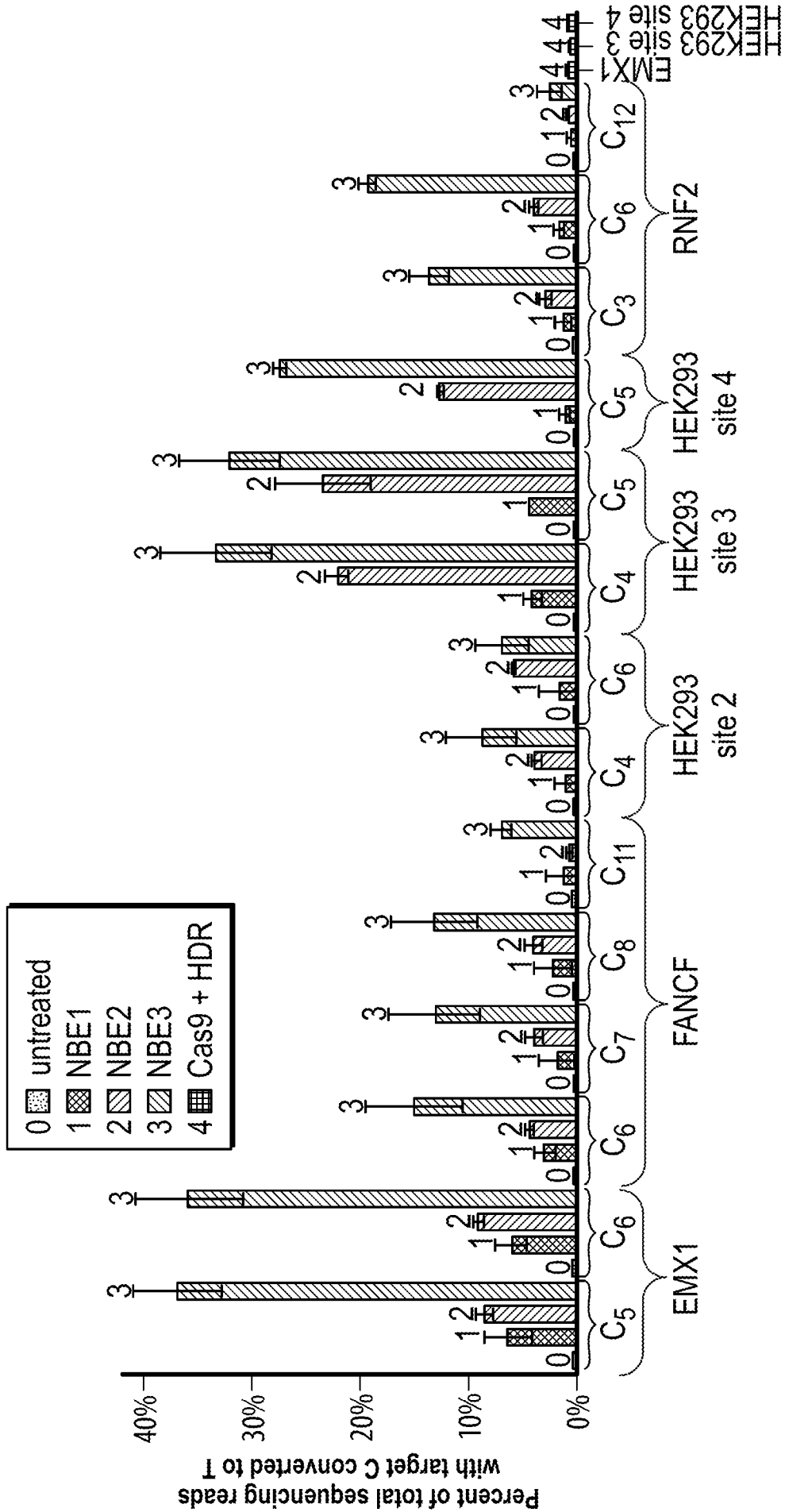


FIGURE 13B

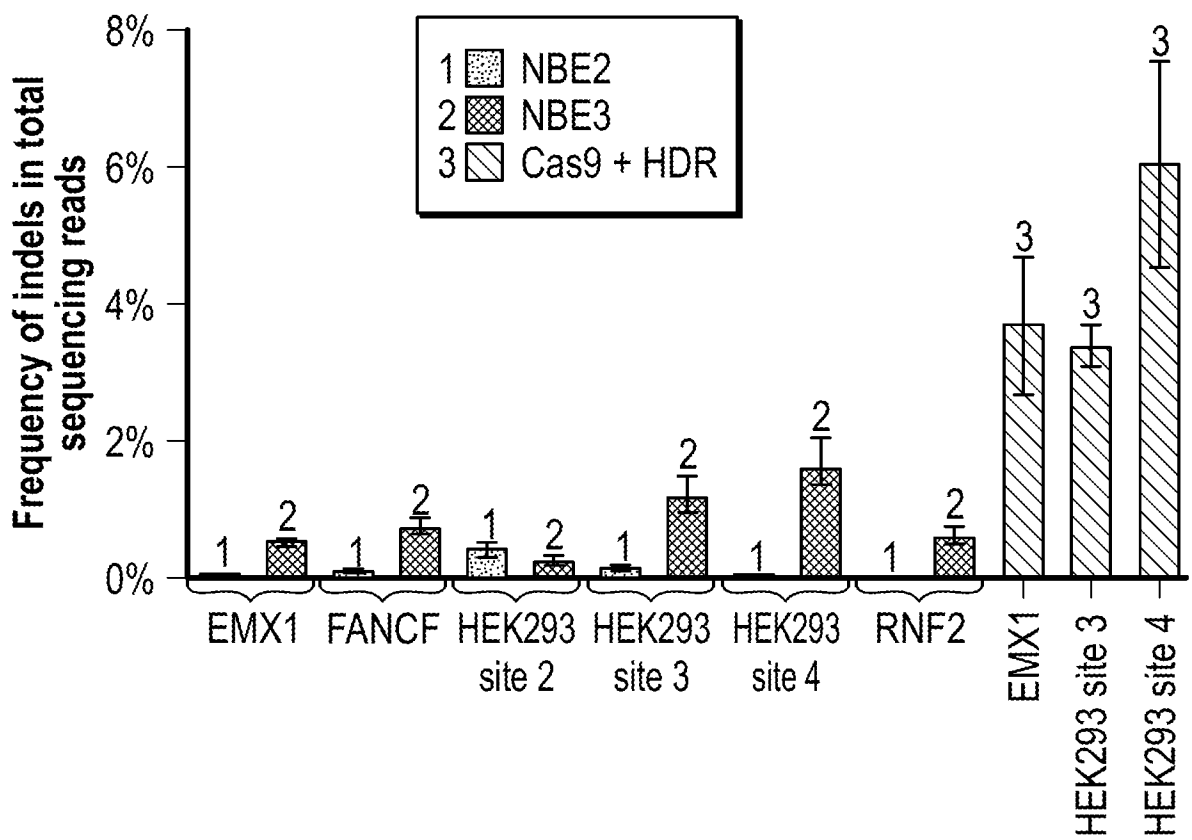


FIGURE 13C

	Lys		Arg → Cys		Leu → Leu			Ala			Val			Tyr			Gln		
	G	A	G	C	C	T	G	C	A	G	T	G	T	A	C	A	C	A	
APOE4C158R																			
A	0.0	100.0	0.0	0.2	0.0	0.1	0.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
C	0.0	0.0	0.0	86.7	0.0	91.3	91.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
G	100.0	0.0	0.0	100.0	0.2	100.0	0.4	0.1	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
T	0.0	0.0	0.0	10.9	0.0	8.2	8.4	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FIGURE 14A

	His		Tyr		Asn			Tyr			Met			Cys			Asp → Asn			
	C	A	T	A	A	C	T	A	C	A	T	G	T	G	T	G	T	A	C	
TP53N239D																				
A	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
G	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

FIGURE 14B

	Arg		Ala		Met			Ile			Cys → Tyr			Lys				
	C	G	C	G	A	T	G	C	A	T	C	T	G	A	C	A	C	
TP53Y163C																		
A	0.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	100.0	100.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
G	0.0	0.0	0.0	99.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

FIGURE 14C

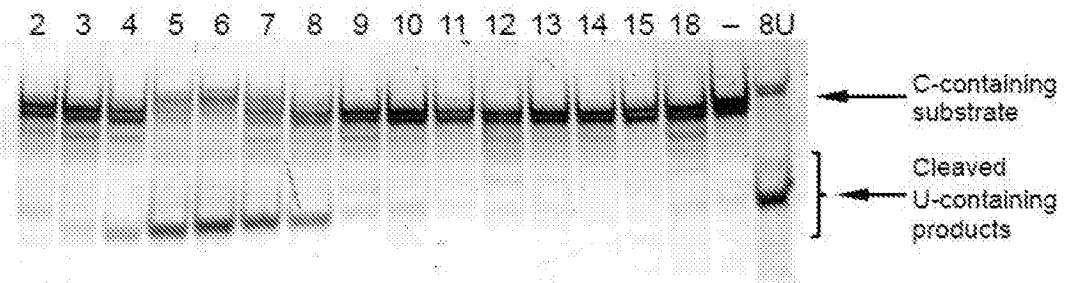


FIGURE 15A

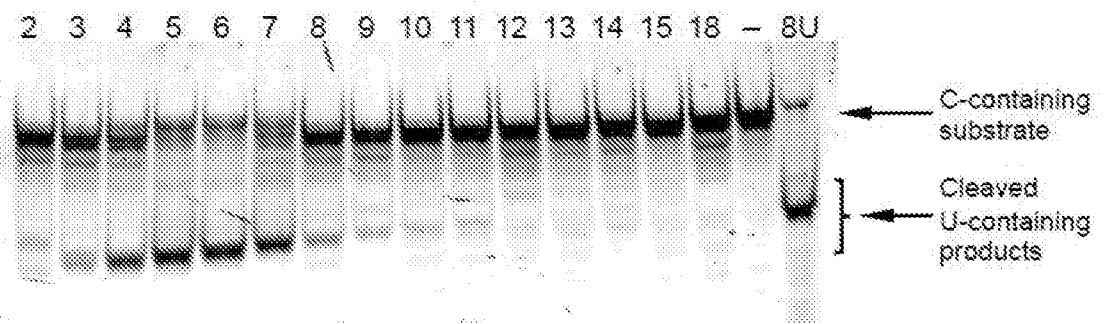


FIGURE 15B

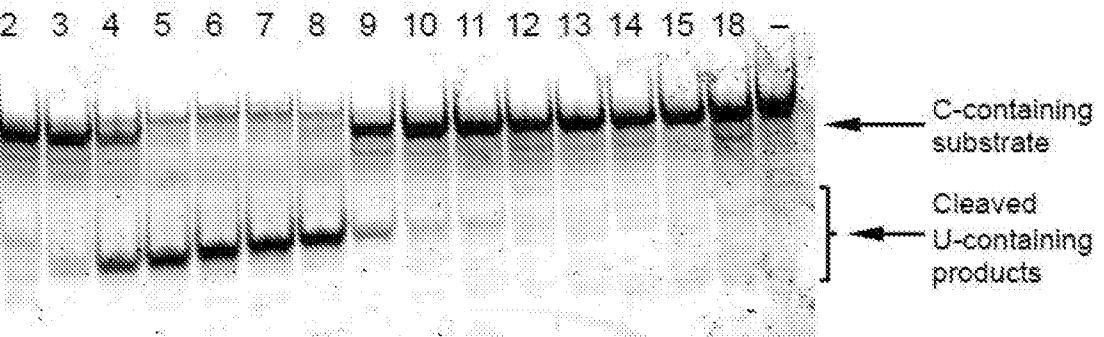


FIGURE 15C

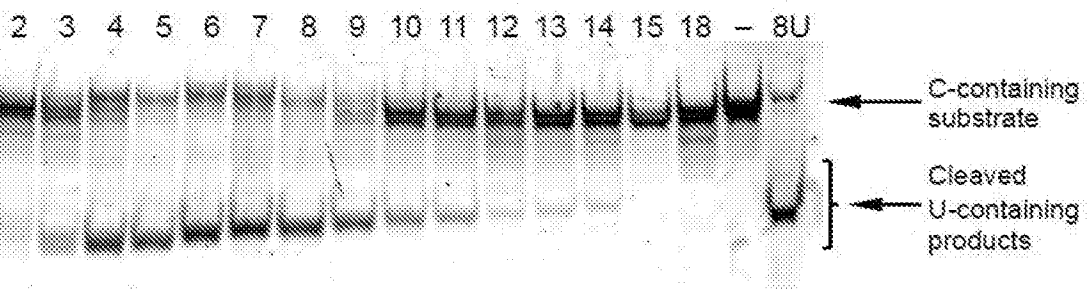


FIGURE 15D

APOE4 Cys112Arg: 5'-GGAGGACGTGC₁₁GCGGCCGCCTGG
 APOE4 Cys158Arg: 5'-GAAGC₈GCCTGGCAGTGTACCAGG
 CTNNB1 Thr41Ala: 5'-CTGTGGC₇AGTGGCACCAGAATGG
 HRAS Gln61Arg: 5'-CCTCCC₅GGCCGGCGGTATCCAGG
 p53 Tyr163Cys: 5'-GCTTGC₉AGATGGCCATGGCGCGG
 p53 Tyr236Cys: 5'-ACACATGC₈AGTTGTAGTGGATGG
 p53 Asn239Asp: 5'-TGTC₄ACACATGTAGTTGTAGTGG

FIGURE 16A

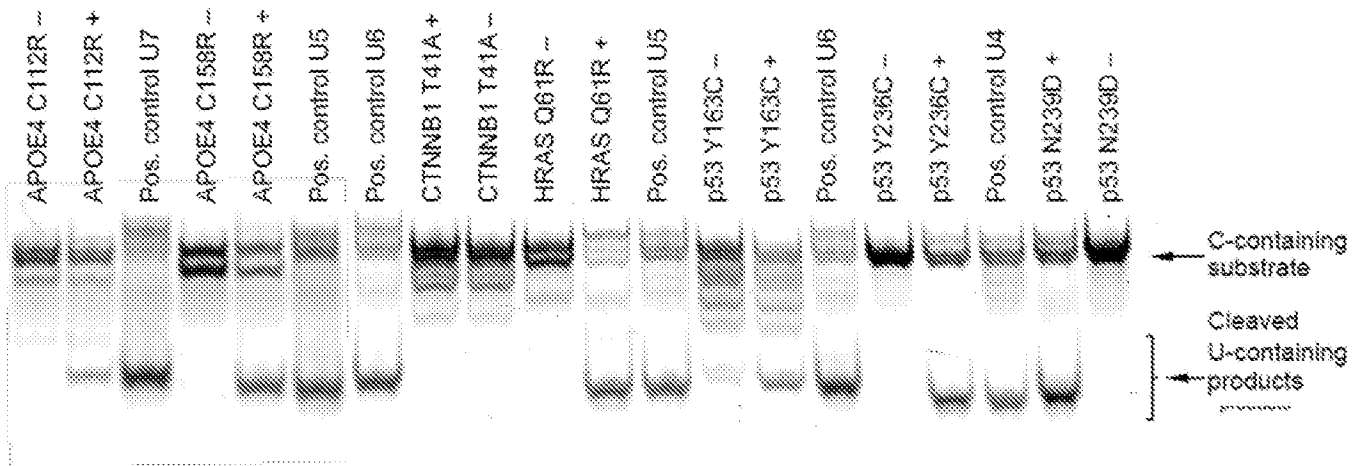


FIGURE 16B

Protospacer and PAM sequence: 5'-TTCCCCCCCCGATTTATTTATGG-3'

Sequence	% of total reads
CCCCCCCC	62.4
TTTTTTCC	18.2
TTTTTTTC	13.4
TTTTTTTT	3.3
TCCCCCCC	0.8
CCCCTTCC	0.3
CCCTTTCC	0.3
TTTTTCCC	0.3
CCCCTCCC	0.3

FIGURE 17

EMX1: GAGTC₅C₆GAGCAGAAGAAGAAGGG
 FANCF: GGAATC₆C₇C₈TTC₁₁TGCAGCACCTGG
 HEK293 site 2: GAAC₄AC₆AAAGCATAGACTGCGGG
 HEK293 site 3: GGCC₄C₅AGACTGAGCACGTGATGG
 HEK293 site 4: GGCAC₅TGCGGCTGGAGGTCCGGG
 RNF2: GTC₃ATC₆TTAGTCATTACCTGAGG

FIGURE 18A

EMX1	C ₅	C ₆
NBE1	6.2%	6.5%
NBE1 + UGI	9.7%	10.1%
NBE2	8.0%	8.7%

FIGURE 18B

FANCF	C ₆	C ₇	C ₈	C ₁₁
NBE1	3.7%	3.2%	3.4%	2.4%
NBE1 + UGI	7.5%	7.6%	7.5%	1.6%
NBE2	4.7%	4.6%	4.6%	0.8%

FIGURE 18C

HEK293 site 2	C ₄	C ₆
NBE1	0.4%	0.4%
NBE1 + UGI	1.6%	2.6%
NBE2	3.4%	5.9%

FIGURE 18D

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<u>HEK293 site 3</u>	<u>C₄</u>	<u>C₅</u>
NBE1	2.0%	1.9%
NBE1 + UGI	6.5%	6.7%
NBE2	10.0%	12.5%

FIGURE 18E

<u>HEK293 site 4</u>	<u>C₅</u>
NBE1	1.4%
NBE1 + UGI	5.4%
NBE2	8.2%

FIGURE 18F

<u>RNF2</u>	<u>C₃</u>	<u>C₆</u>
NBE1	0.7%	1.4%
NBE1 + UGI	3.4%	3.9%
NBE2	2.5%	3.7%

FIGURE 18G

<u>Non-protospacer Cs</u>	<u>C</u>	<u>T</u>
untreated	99.93%	0.03%
NBE1	99.95%	0.03%
NBE1 + UGI	99.91%	0.06%
NBE2	99.92%	0.04%

FIGURE 18H

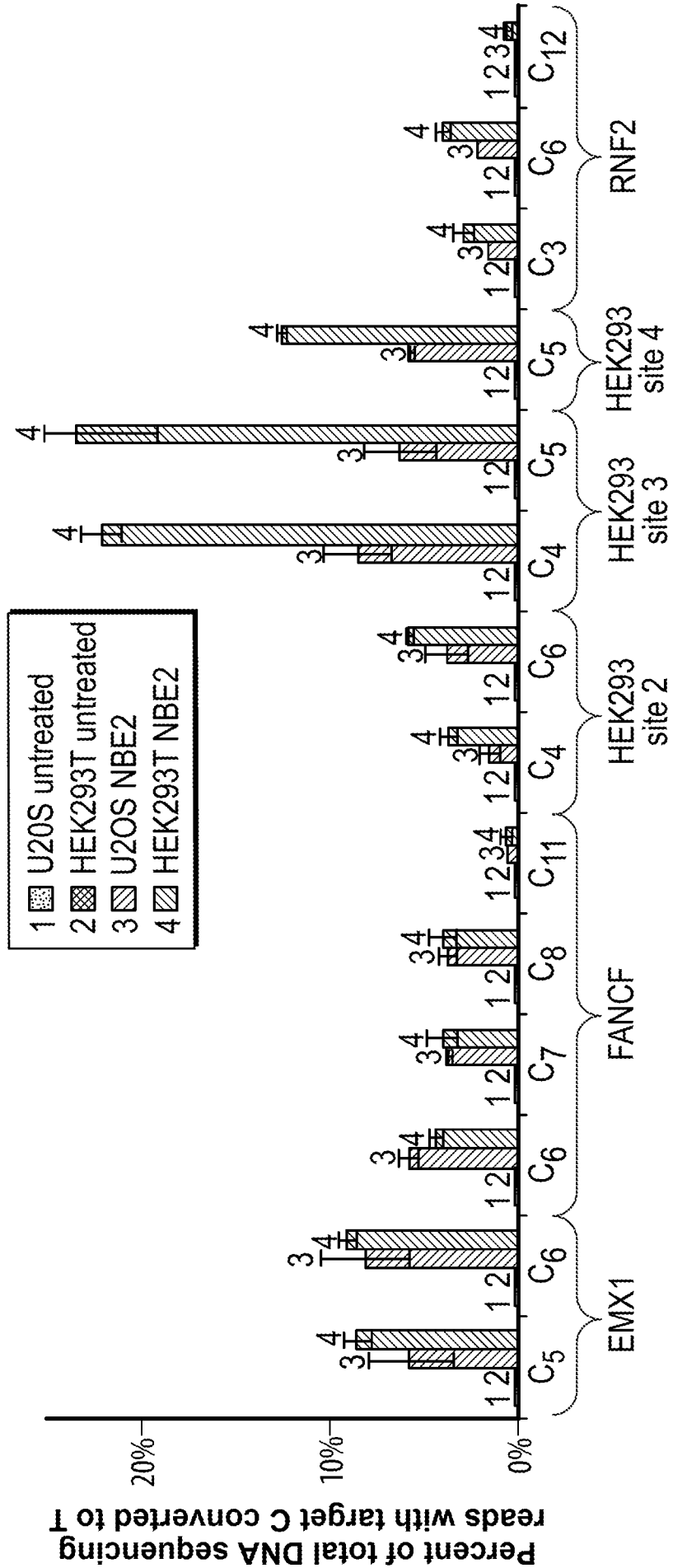


FIGURE 19

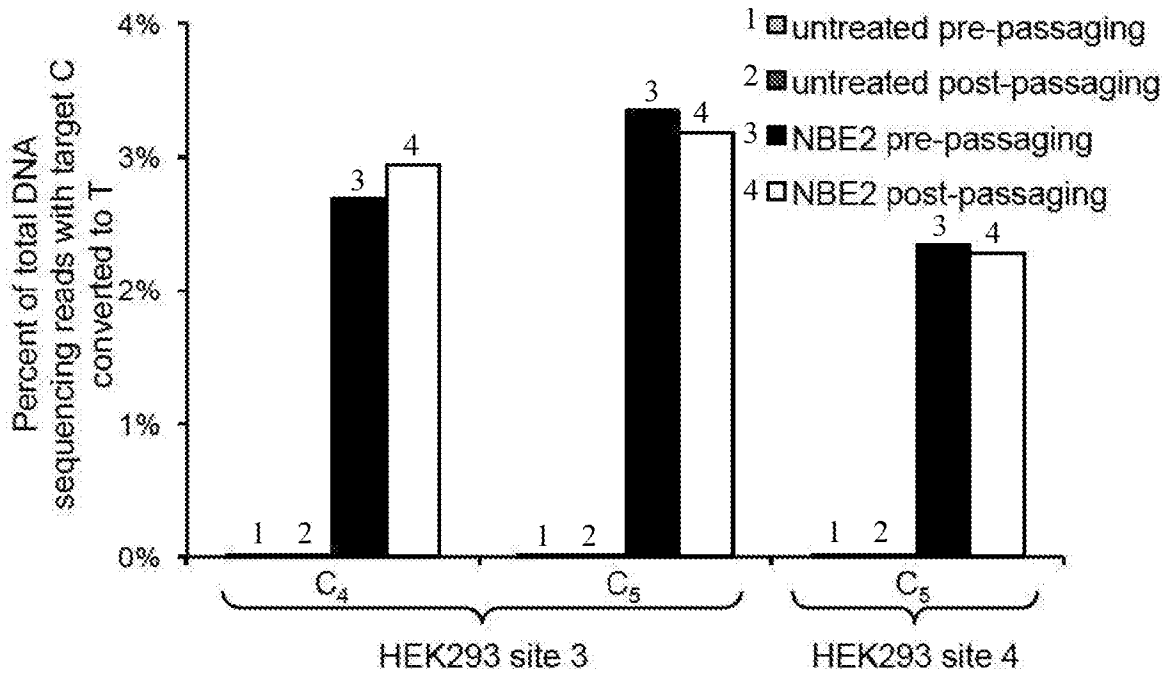


FIGURE 20

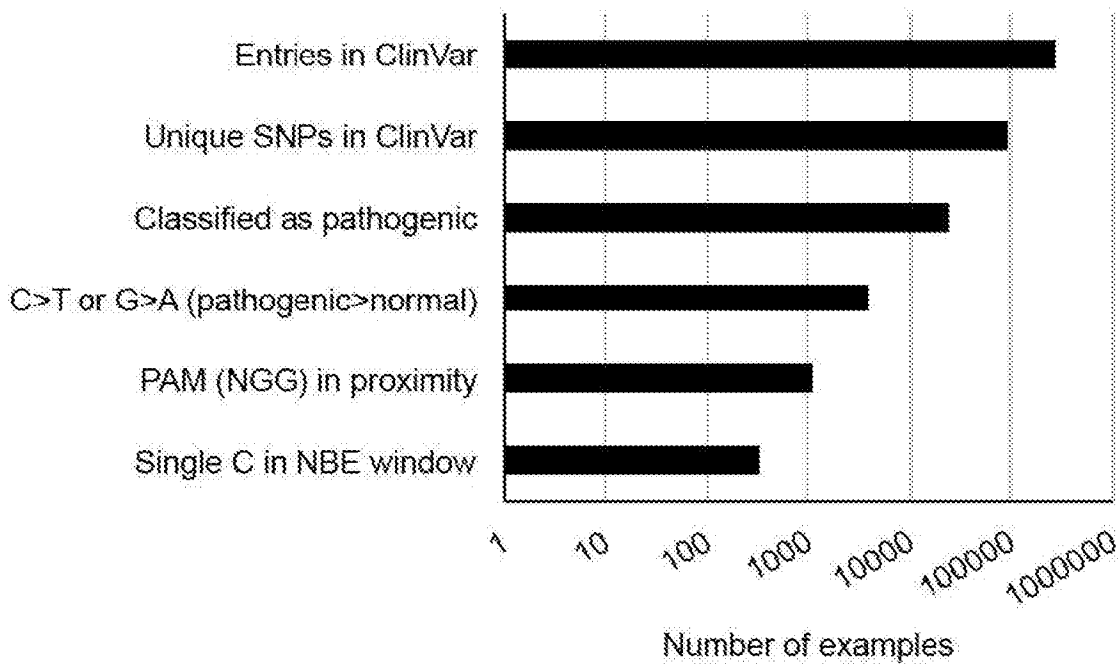


FIGURE 21

HEK293 site 3		G	G	C	C4	C5	A	G	A	C	T	G	A	G	C	A	C	G	T	G	A	T	G	G	
1.85 uM NBE1	A	0.2	0.1	0.1	0.1	0.0	99.9	0.2	99.9	0.1	0.0	0.2	99.9	0.1	0.1	100.0	0.1	0.1	0.0	0.2	99.6	0.0	0.1	0.1	
	C	0.0	0.0	92.2	74.8	71.5	0.0	0.0	96.6	0.0	0.0	0.0	99.8	0.0	99.7	0.0	99.7	0.0	0.0	0.0	1.3	0.0	0.0	0.0	
	G	99.7	99.8	0.0	0.0	0.0	0.0	99.8	0.0	0.0	0.0	99.8	0.0	99.8	0.0	0.0	99.8	0.0	99.8	0.0	0.0	0.0	99.9	99.8	
	T	0.1	0.1	7.7	25.1	28.5	0.0	0.0	3.3	99.9	0.1	0.0	0.1	0.0	0.2	0.0	0.2	0.0	99.9	0.0	0.0	100.0	0.0	0.0	
HEK293 site 4		G	G	C	A	C5	T	G	C	G	C	T	G	G	A	G	A	G	T	G	G	T	G	G	
1.85 uM NBE1	A	0.3	0.2	0.1	100.0	0.1	0.0	0.2	0.1	0.2	0.2	0.1	0.0	0.3	0.5	100.0	0.1	0.1	0.0	0.1	0.1	0.1	0.1	0.2	0.2
	C	0.0	0.0	98.8	0.0	60.1	0.0	0.0	97.0	0.0	0.0	99.4	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
	G	99.7	99.7	0.0	0.0	0.0	0.0	99.8	0.0	99.8	99.8	0.0	0.0	99.7	98.4	0.0	99.8	99.8	0.1	99.8	99.8	99.8	99.8	99.7	
	T	0.0	0.1	1.1	0.0	39.3	100.0	0.1	2.9	0.0	0.1	0.5	100.0	0.0	0.1	0.0	0.0	0.0	99.9	0.0	0.1	0.1	0.0	0.0	
RNF2		G	T	C3	A	T	C6	T	T	A	G	T	C12	A	T	T	A	C	C	T	G	A	G	G	
1.85 uM NBE1	A	0.1	0.0	0.1	99.9	0.0	0.0	0.0	0.0	99.9	0.1	0.0	0.1	99.9	0.0	0.0	99.9	0.1	0.0	0.0	0.1	98.6	0.0	0.0	
	C	0.0	0.0	99.1	0.0	0.0	57.8	0.0	0.0	0.1	0.0	0.0	86.4	0.0	0.0	0.1	99.9	99.9	0.0	0.0	1.4	0.0	0.0		
	G	99.9	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.8	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.8	0.0	99.9	99.9	
	T	0.0	99.9	40.8	0.0	100.0	42.1	100.0	100.0	0.0	0.1	99.9	13.4	0.0	100.0	99.9	0.0	0.0	0.0	99.9	0.1	0.0	0.1	0.0	

FIGURE 22 (CONTINUED)

GUIDE-seq counts

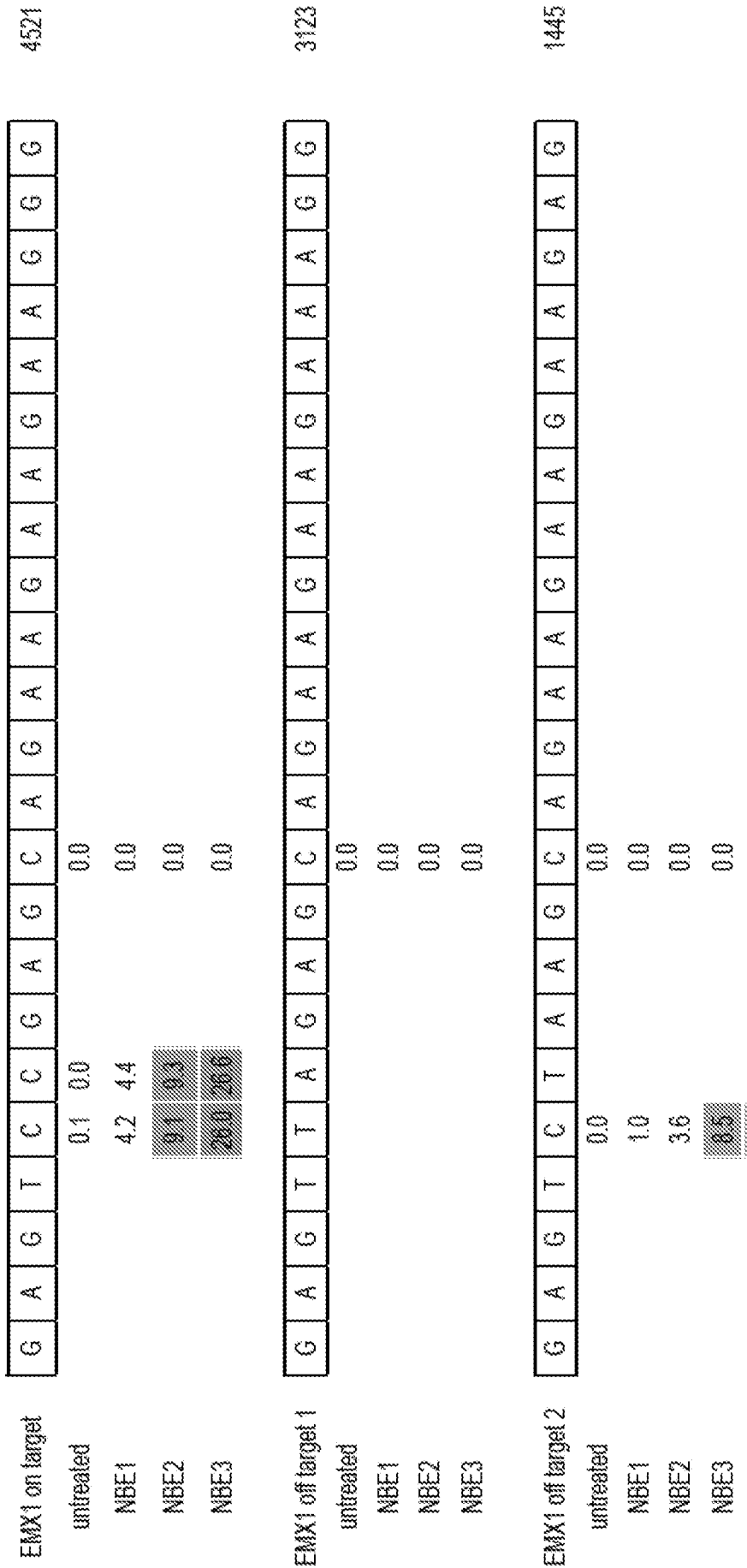


FIGURE 23

GUIDE-seq counts

EMX1 off target 3	G	A	G	G	C	C	C	G	A	G	A	A	G	A	A	A	G	A	C	G	G	700
untreated	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE2	0.8	0.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE3	5.1	5.2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

EMX1 off target 4	G	A	G	T	C	C	C	T	A	G	G	A	G	A	A	G	A	A	G	A	G	300
untreated	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE1	0.2	0.2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE2	0.5	0.5	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE3	2.2	2.2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

EMX1 off target 6	G	A	G	T	C	C	C	G	G	G	A	A	G	A	A	G	A	A	A	G	G	216
untreated	0.0	0.1	0.0	0.1	0.0	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE1	0.1	0.1	0.0	0.1	0.0	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE2	0.2	0.3	0.0	0.3	0.0	0.3	0.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE3	1.0	1.0	0.0	1.0	0.0	1.0	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

FIGURE 23 (CONTINUED)

GUIDE-seq counts

	G	A	G	C	C	C	G	G	A	G	C	A	G	A	A	G	A	G	G	A	G	G	G	143
EMX1 off target 7																								
untreated	0.0			0.0							0.0													
NBE1	0.0			0.0							0.0													
NBE2	0.1			0.0							0.0													
NBE3	0.1			0.1							0.0													
EMX1 off target 8																								
untreated							0.0	0.0																102
NBE1							0.0	0.0																
NBE2							0.0	0.0																
NBE3							0.1	0.1																
EMX1 off target 9																								
untreated							0.0	0.1																67
NBE1							0.0	0.1																
NBE2							0.0	0.1																
NBE3							0.3	0.3																
EMX1 off target 10																								
untreated							0.0																	38
NBE1							0.0																	
NBE2							0.3																	
NBE3							0.5																	
							2.1																	

FIGURE 23 (CONTINUED)

GUIDE-seq counts

		G	G	A	A	T	C	C	C	C	T	T	C	T	G	C	A	G	C	A	C	C	A	C	C	T	G	G	
FANCF on target		4816																											
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		37	32	34									24																
NBE2		41	35	35									0.4																
NBE3		191	160	167									0.8																
FANCF off target 1		2099																											
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.4	0.0	0.1								0.0																
NBE2		0.2	0.2	0.3	0.3								0.0																
NBE3		1.9	2.1	1.5	1.3								1.3																
FANCF off target 2		524																											
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1
NBE3		0.2	0.2	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1
FANCF off target 3		150																											
untreated		0.0	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		0.0	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
FANCF off target 4		125																											
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		0.0	0.0	0.0	0.0	0.1	0.1	0.1	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3		0.0	0.0	0.0	0.0	0.7	0.4	0.3	0.3	0.3	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FIGURE 24

GUIDE-seq counts

		G	G	A	T	T	G	C	C	C	A	A	T	C	C	G	C	C	A	G	C	A	C	C	T	G	G	101
FANCF off target 5																												
untreated							0.1	0.0						0.0	0.0													
NBE1							0.1	0.0						0.0	0.1													
NBE2							0.1	0.0						0.0	0.0													
NBE3							0.1	0.0						0.0	0.0													
FANCF off target 6																												
untreated							0.0	0.0	0.0					0.0	0.1													
NBE1							0.0	0.0	0.0					0.0	0.0													
NBE2							0.2	0.2	0.2					0.2	0.1	0.0												
NBE3							0.3	0.3	0.3					0.3	0.1	0.0												
FANCF off target 7																												
untreated							0.0	0.0	0.0					0.0	0.0													
NBE1							0.0	0.0	0.0					0.0	0.0													
NBE2							0.0	0.0	0.0					0.0	0.0													
NBE3							0.5	0.5	0.5					0.3	0.1													
FANCF off target 8																												
untreated							0.0	0.0	0.0					0.0	0.0													
NBE1							0.0	0.0	0.0					0.0	0.0													
NBE2							1.4	0.5	0.5					0.4	0.5													
NBE3							0.1	0.1	0.0					0.0	0.0													

FIGURE 24 (CONTINUED)

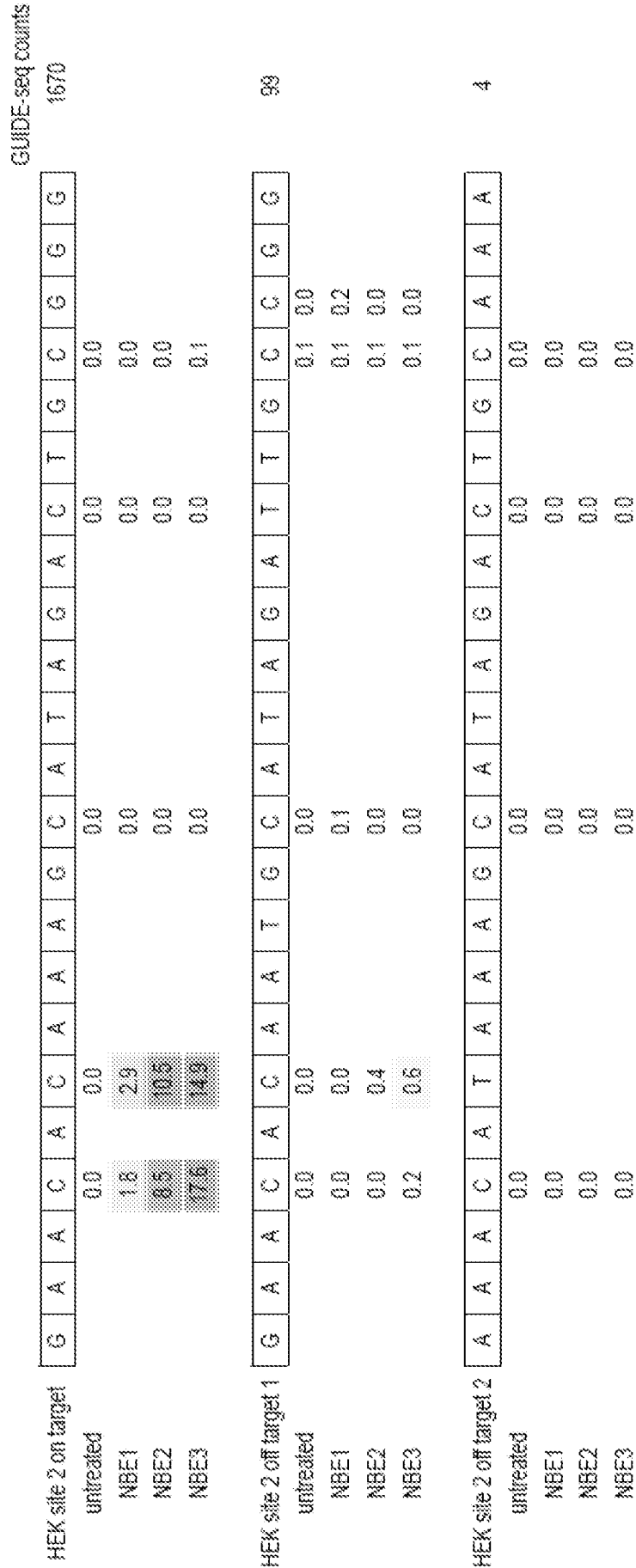


FIGURE 25

GUIDE-seq counts

		G	G	C	C	C	A	A	G	A	C	T	G	A	G	C	A	C	A	G	T	G	A	T	G	G	
HEK site 3 on target		2074																									
untreated		0.1	0.1	0.1	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.1	4.6	4.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		5.6	27.3	20.4	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3		0.5	33.4	3.5	0.0	0.0	0.5	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
HEK site 3 off target 1		327																									
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
HEK site 3 off target 2		306																									
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
HEK site 3 off target 3		196																									
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
HEK site 3 off target 4		10																									
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
HEK site 3 off target 5		2																									
untreated		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE1		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3		0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FIGURE 26

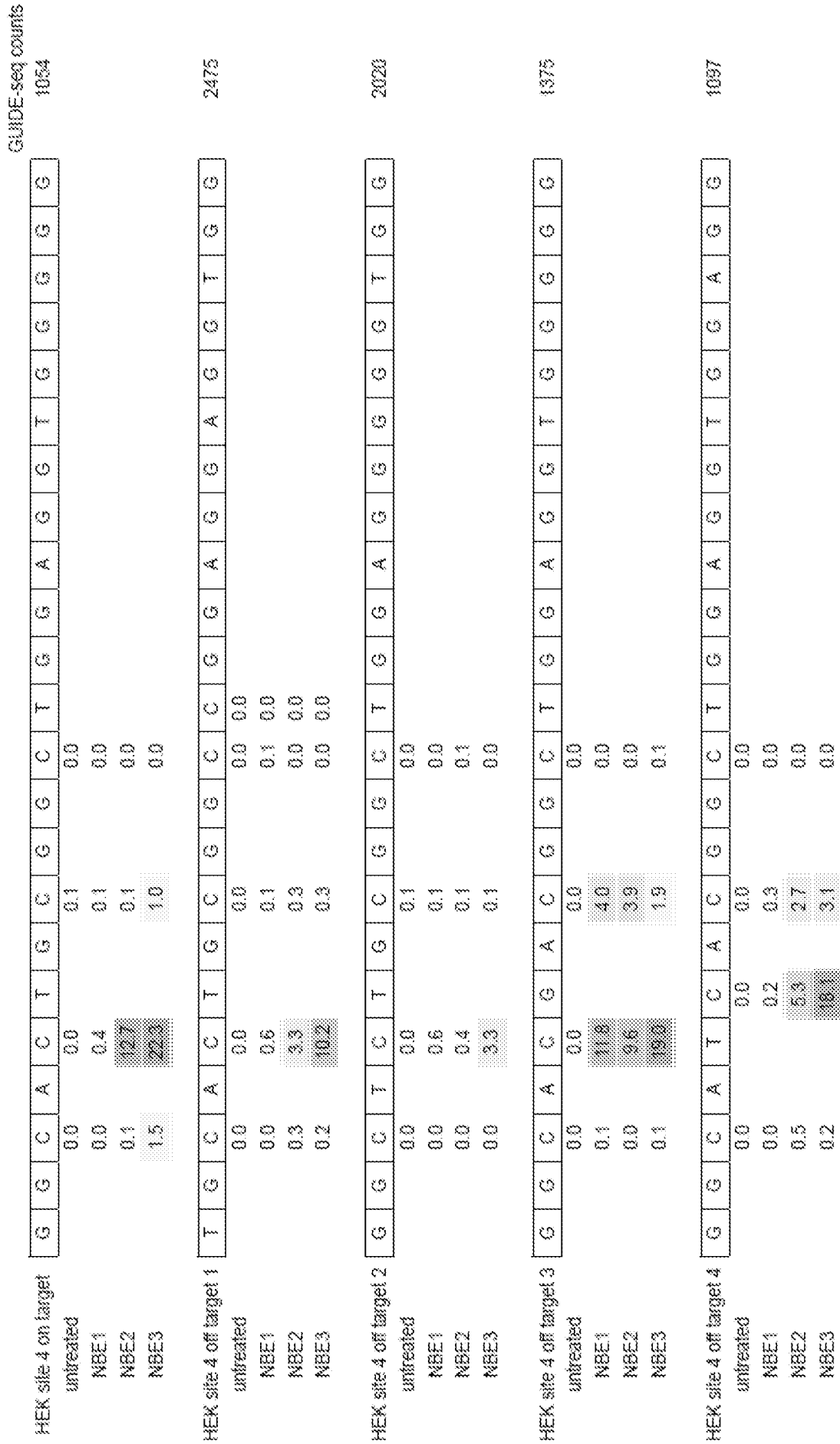


FIGURE 27

GUIDE-seq counts

1002

HEK site 4 off target 5

untreated	G	G	C	G	C	G	C	T	G	C	G	G	C	G	C	G	G	A	G	G	T	G	G	A	G	G
NBE1	0.0	0.0	0.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1

982

HEK site 4 off target 6

untreated	G	G	C	A	C	T	G	A	G	A	G	G	C	T	G	G	G	G	G	G	T	G	G	G	G	G
NBE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2	0.6	0.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6	3.6
NBE3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

981

HEK site 4 off target 7

untreated	A	G	C	A	G	T	G	C	T	A	G	C	T	A	G	A	G	G	T	G	G	T	G	G	T	G	G
NBE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

789

HEK site 4 off target 8

untreated	G	G	C	A	C	T	G	C	T	A	C	T	G	G	G	G	G	G	T	G	G	T	G	G	T	G	G
NBE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
NBE2	0.2	0.2	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	
NBE3	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	0.2	

712

HEK site 4 off target 9

untreated	G	G	C	A	C	T	G	G	G	G	C	T	G	G	G	G	G	A	G	G	G	A	G	G	G	G
NBE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

664

HEK site 4 off target 10

untreated	G	G	C	A	C	T	G	G	G	G	T	T	G	G	A	G	G	A	G	G	T	G	G	G	G	G
NBE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
NBE2	0.6	0.6	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5
NBE3	0.9	0.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FIGURE 27 (CONTINUED)

Non-protospacer Cs	C (%)	T (%)
untreated	99.94	0.04
NBE1	99.92	0.05
NBE2	99.92	0.05
NBE3	99.94	0.03

FIGURE 28

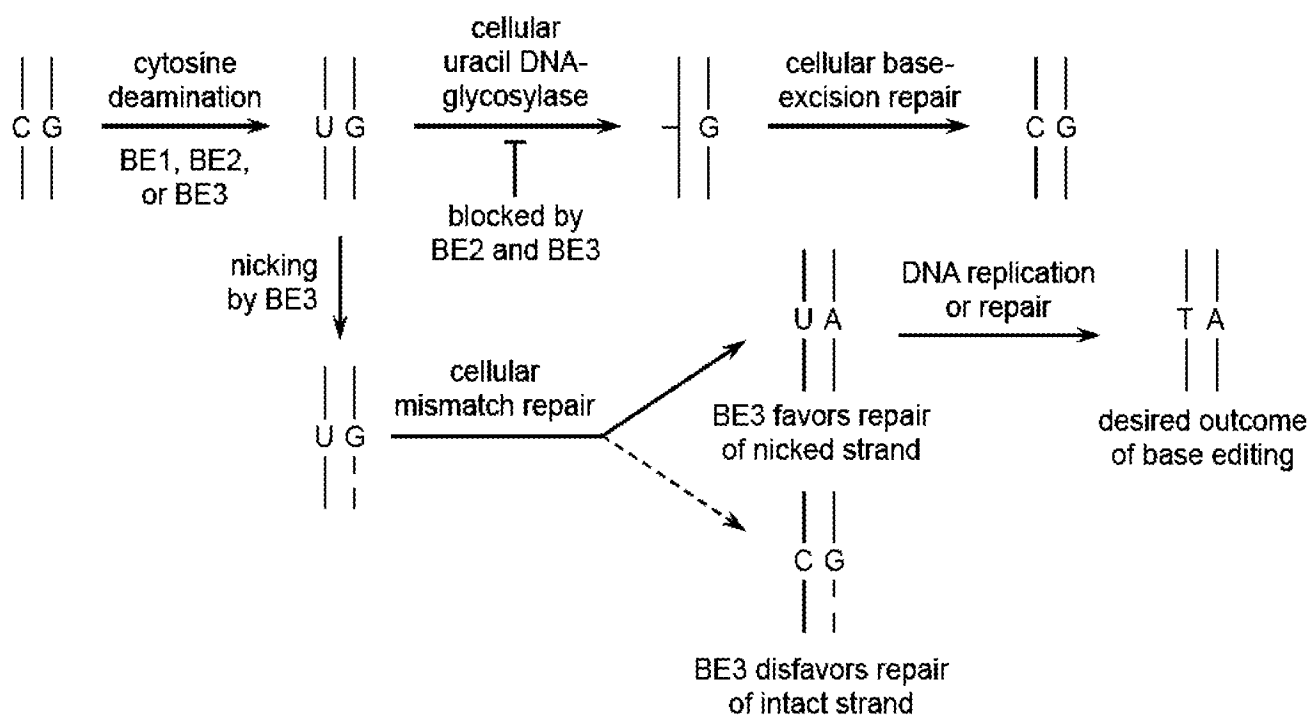


FIGURE 29A

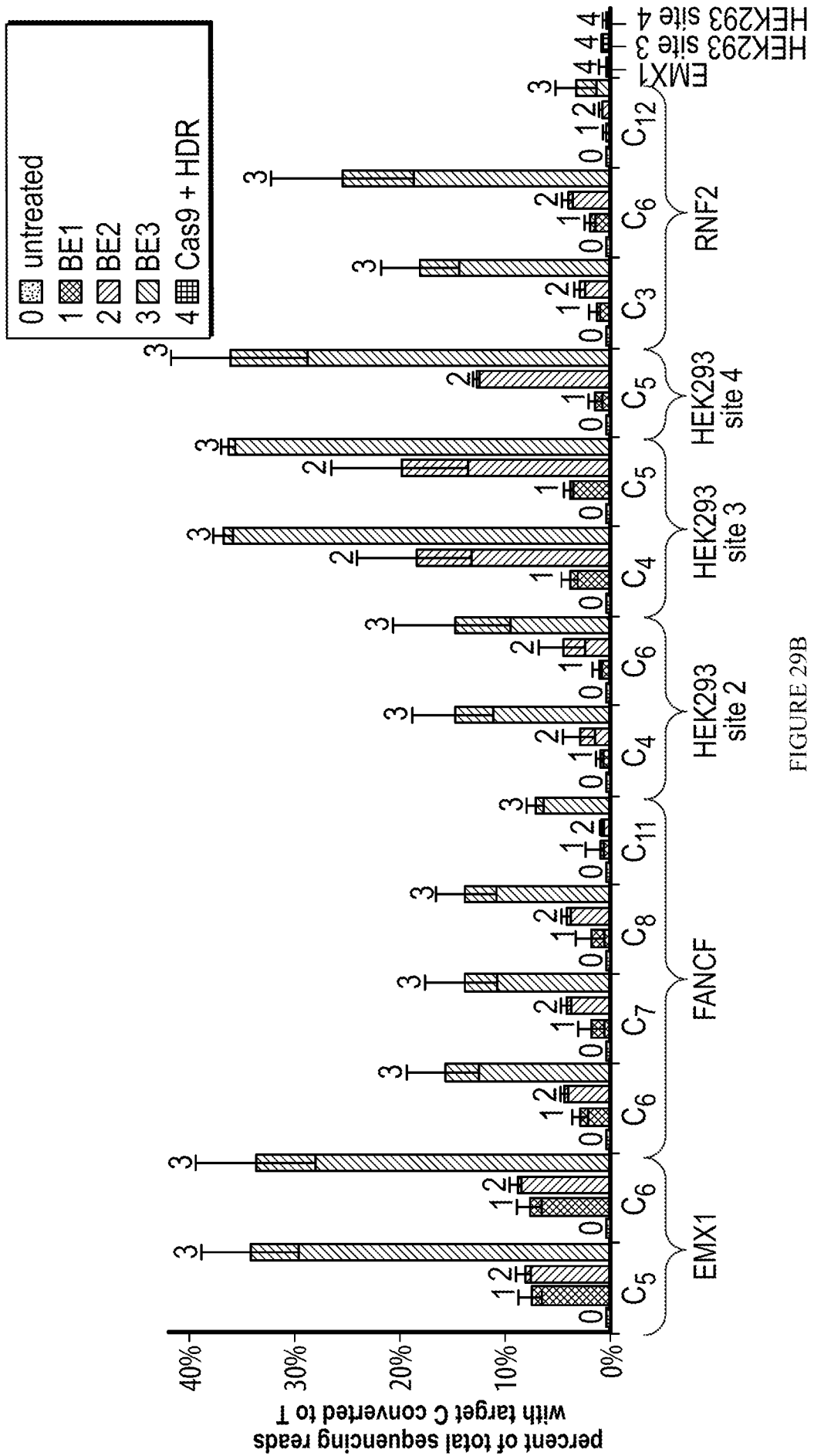


FIGURE 29B

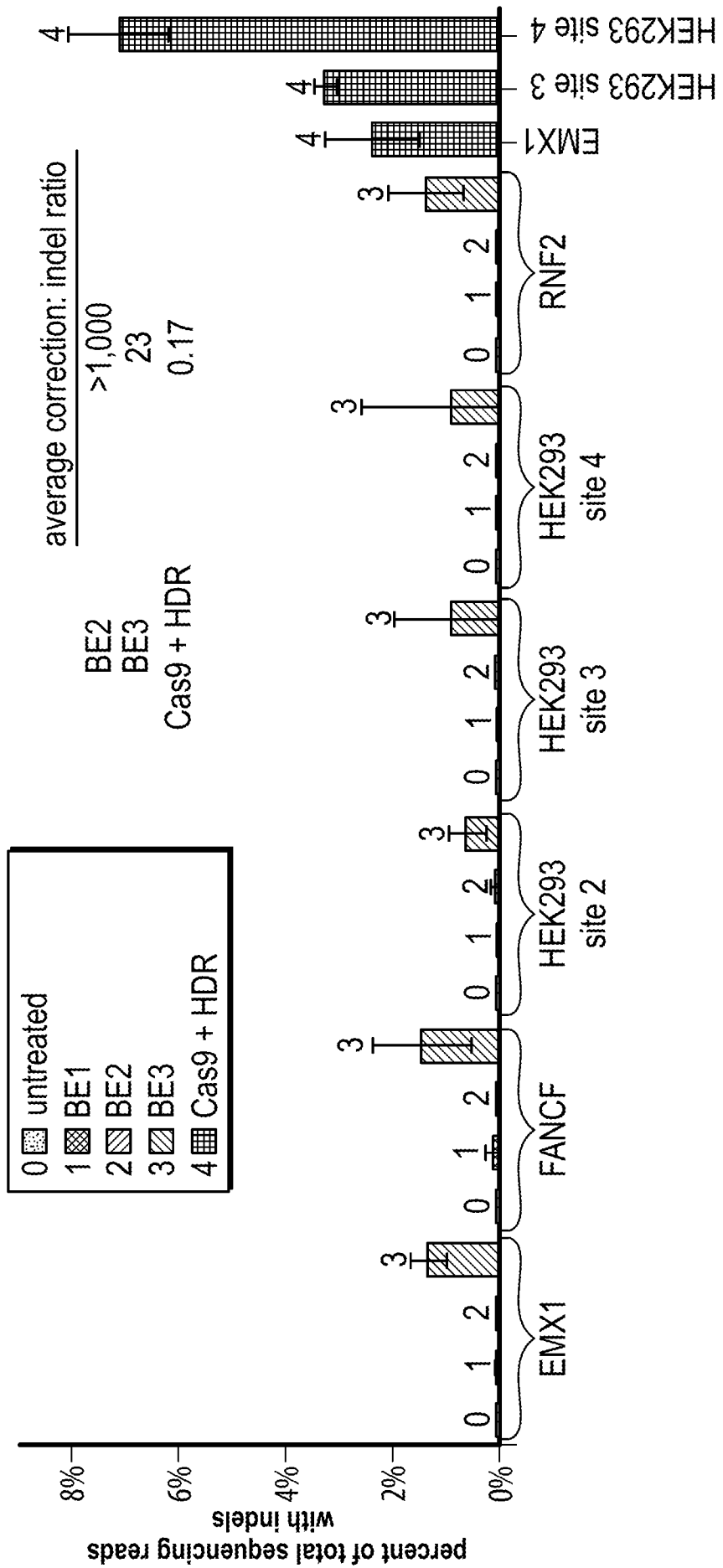


FIGURE 29C

untreated	Lys			Arg			Leu			Ala			Val			Tyr			Gln			indel %			
	G	A	A	G	C	G	G	T	G	G	C	A	G	T	G	T	A	A	C	C	A		G	A	G
APOE4C158R	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
A	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
C	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
G	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.9
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3 treated	Lys			Arg → Cys			Leu → Leu			Ala			Val			Tyr			Gln			indel %			
APOE4C158R	G	A	A	G	C	G	G	T	G	G	C	A	G	T	G	T	A	A	C	C	A		G	A	G
A	0.1	100.0	0.0	0.5	0.0	1.3	0.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1
C	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
G	99.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
Cas9 + HDR	Lys			Arg → Cys			Leu			Ala			Val			Tyr			Gln			indel %			
APOE4C158R	G	A	A	G	C	G	G	T	G	G	C	A	G	T	G	T	A	A	C	C	A		G	A	G
A	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
C	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
G	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FIGURE 30A

untreated	Arg			Ala			Met			Ala			Ile			Cys			Lys			indel %	
	C	C	G	C	C	G	A	T	G	C	C	G	A	T	C	T	G	C	A	A	G		C
TP53Y163C	C	C	G	C	C	G	A	T	G	C	C	G	A	T	C	T	G <td>C</td> <td>A</td> <td>A</td> <td>G</td> <td>C</td> <td>0.0</td>	C	A	A	G	C	0.0
A	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.1	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	100.0	99.9	0.0	0.0	0.0
C	100.0	100.0	0.0	100.0	100.0	0.0	0.0	0.0	0.0	100.0	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	100.0
G	0.0	0.0	100.0	0.0	0.0	99.9	0.0	0.0	100.0	99.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.9	0.0	0.0	0.0	100.0
T	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3 treated	Arg			Ala			Met			Ala			Ile			Cys → Tyr			Lys			indel %	
TP53Y163C	C	C	G	C	C	G	A	T	G	C	C	G	A	T	C	T	G <td>C</td> <td>A</td> <td>A</td> <td>G</td> <td>C</td> <td>0.7</td>	C	A	A	G		C
A	0.0	0.0	0.0	0.0	0.1	0.0	100.0	0.0	0.0	0.1	0.0	0.0	100.0	0.0	0.0	0.0	0.0	7.6	0.0	100.0	100.0	0.0	0.0
C	100.0	100.0	0.0	100.0	0.0	100.0	0.0	0.0	0.0	99.9	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.4	100.0	0.0	0.0	0.0	100.0
G	0.0	0.0	100.0	0.0	99.9	0.0	0.0	0.0	99.9	99.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	97.8	0.0	0.0	0.0	0.0	100.0
T	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.1	0.0	0.1	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0
Cas9 + HDR	Arg			Ala			Met			Ala			Ile			Cys → Tyr			Lys			indel %	
TP53Y163C	C	C	G	C	C	G	A	T	G	C	C	G	A	T	C	T	G <td>C</td> <td>A</td> <td>A</td> <td>G</td> <td>C</td> <td>6.1</td>	C	A	A	G		C
A	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	99.9	100.0	0.0	0.0
C	100.0	100.0	0.0	100.0	0.0	100.0	0.0	0.0	0.0	100.0	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	100.0
G	0.0	0.0	100.0	0.0	100.0	0.0	0.0	0.0	100.0	99.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	100.0
T	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FIGURE 30B

GUIDE-seq
counts/ ChIP-seq
intensity
1670/ 117

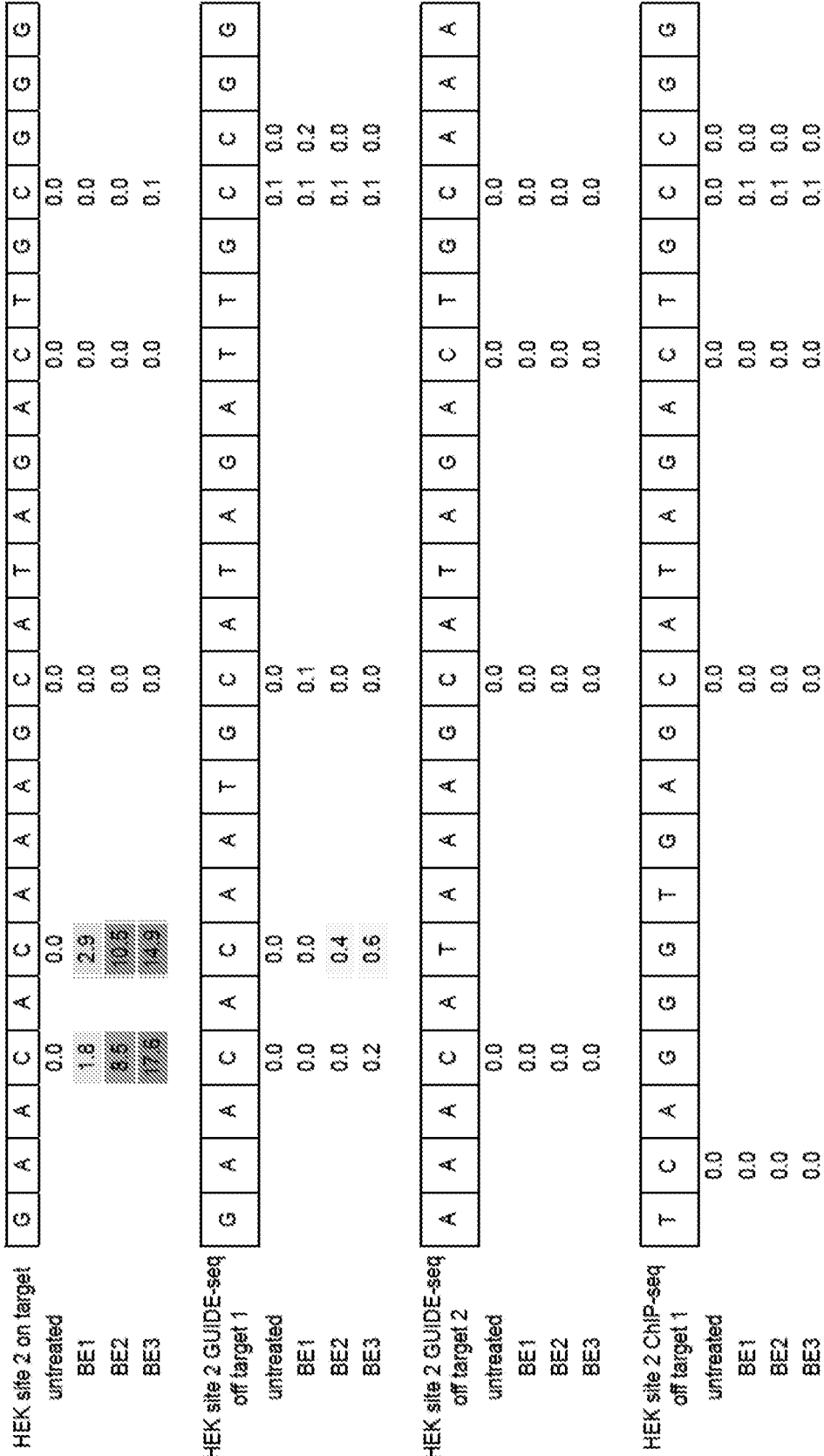


FIGURE 31

GUIDE-seq
counts/ ChIP-seq
intensity
1670/ 117
28

HEK site 2 ChIP-seq
off target 2

G	A	A	T	C	C	G	G	G	A	A	G	G	G	A	A	G	A	C	T	G	C	G	G	G
untreated	0.0																	0.0	0.0					
BE1	0.0																	0.0	0.1					
BE2	0.0																	0.0	0.1					
BE3	0.0																	0.0	0.1					

HEK site 2 ChIP-seq
off target 3

T	G	A	A	A	G	T	G	T	T	G	C	A	T	A	G	A	C	T	G	C	A	G	G
untreated	0.0																	0.0	0.0				
BE1	0.0																	0.0	0.1				
BE2	0.0																	0.0	0.0				
BE3	0.0																	0.0	0.0				

HEK site 2 ChIP-seq
off target 4

G	G	A	G	A	G	A	G	A	G	A	G	C	A	T	A	G	A	C	T	G	C	T	G	G
untreated	0.0																	0.0	0.0					
BE1	0.0																	0.0	0.0					
BE2	0.0																	0.0	0.0					
BE3	0.0																	0.0	0.0					

HEK site 2 ChIP-seq
off target 5

C	C	A	A	A	C	A	A	A	A	A	C	A	C	A	T	A	G	A	C	T	G	C	T	G	G
untreated	0.0	0.0																	0.0	0.0					
BE1	0.0	0.0																0.0	0.0						
BE2	0.0	0.0																0.0	0.0						
BE3	0.0	0.0																0.0	0.0					0.2	

FIGURE 31 (CONTINUED)

GUIDE-seq
counts/ ChIP-seq
intensity
2074/ 163

HEK site 3 on target

untreated	G	G	C	C	C	C	A	A	G	A	C	T	G	A	G	C	A	C	G	T	G	A	T	G	G
BE1	0.1	0.1	0.1	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.1	4.8	4.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3	3.6	27.3	20.4	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
	0.5	3.4	3.7	0.0	0.0	0.5	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

HEK site 3 GUIDE-seq off target 1

untreated	C	A	C	C	C	C	A	A	G	A	C	T	G	A	G	C	A	C	G	T	G	C	T	G	G
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

327

HEK site 3 GUIDE-seq off target 2

untreated	G	A	C	A	C	A	C	A	G	A	C	T	G	G	G	C	A	C	G	T	G	A	G	G	G
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

306

HEK site 3 GUIDE-seq off target 3

untreated	A	G	C	T	C	A	C	A	G	A	C	T	G	A	G	C	A	A	G	T	G	A	G	G	G
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3	0.0	0.0	0.0	0.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

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HEK site 3 GUIDE-seq off target 4

untreated	A	G	A	C	C	C	A	A	G	A	C	T	G	A	G	C	A	A	G	A	G	A	G	G	G
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

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FIGURE 32

GUIDE-seq
counts/ChIP-seq
intensity
3874/153
2

G	A	G	C	C	C	A	G	A	A	A	T	G	A	G	C	A	C	G	T	G	A	G	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 3 GUIDE-seq
off target 5
untreated
BE1
BE2
BE3

0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----

C	A	G	G	A	A	G	C	T	G	G	A	G	A	G	C	A	C	G	T	G	A	G	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 3 ChIP-seq
off target 1
untreated
BE1
BE2
BE3

0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----

A	A	G	G	C	T	G	A	G	G	G	A	G	A	G	C	A	C	G	T	G	A	A	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 3 ChIP-seq
off target 2
untreated
BE1
BE2
BE3

0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----

G	T	C	A	G	G	G	A	A	G	A	A	G	A	G	C	A	C	G	T	G	A	C	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 3 ChIP-seq
off target 3
untreated
BE1
BE2
BE3

0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----

G	T	T	G	T	G	A	A	C	T	G	A	G	A	G	C	A	C	G	T	G	A	G	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 3 ChIP-seq
off target 4
untreated
BE1
BE2
BE3

0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----

A	T	A	T	T	T	G	C	T	G	G	A	G	A	G	C	A	C	G	T	G	A	A	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 3 ChIP-seq
off target 5
untreated
BE1
BE2
BE3

0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----	-----

FIGURE 32 (CONTINUED)

GUIDE-seq
counts/ ChIP-seq
intensity
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HEK site 4 on target

untreated	G	G	C	A	C	T	G	C	G	G	C	T	G	G	A	G	G	T	G	G	G	G	G
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.0	0.0	0.4	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3	0.1	1.5	0.1	0.1	1.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

HEK site 4 GUIDE-seq off target 1

untreated	T	G	C	A	C	T	G	C	G	G	C	C	T	G	G	A	G	G	T	G	G	T	G	G
BE1	0.0	0.0	0.0	0.6	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE2	0.3	0.2	0.3	0.3	0.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE3	0.2	0.2	0.2	0.2	0.2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

2475

HEK site 4 GUIDE-seq off target 2

untreated	G	G	C	T	C	T	G	C	G	G	C	T	G	G	A	G	G	G	T	G	G	T	G	G
BE1	0.0	0.0	0.0	0.6	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE2	0.0	0.0	0.4	0.1	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE3	0.0	0.0	0.3	0.3	0.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

2020

HEK site 4 GUIDE-seq off target 3

untreated	G	G	C	A	C	G	A	C	G	G	C	T	G	G	A	G	G	T	G	G	G	T	G	G
BE1	0.0	0.1	0.0	0.3	4.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE2	0.0	0.0	0.0	0.6	3.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE3	0.1	0.1	0.1	0.9	1.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

1375

HEK site 4 GUIDE-seq off target 4

untreated	G	G	C	A	T	C	A	C	G	G	C	T	G	G	A	G	G	T	G	G	A	G	G	G
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE2	0.0	0.0	0.5	0.2	5.3	0.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE3	0.2	0.2	0.2	0.2	3.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

1097

FIGURE 33

GUIDE-seq
counts/ ChIP-seq
intensity
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G	G	C	G	C	T	G	C	G	C	G	G	A	G	G	T	G	G	A	G	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 4 GUIDE-seq
off target 5
untreated
BE1
BE2
BE3

1002

G	G	C	A	C	T	G	A	G	A	C	T	G	G	G	G	T	G	G	G	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 4 GUIDE-seq
off target 6
untreated
BE1
BE2
BE3

562

A	G	C	A	G	T	G	C	G	G	C	T	A	G	A	G	G	T	G	G	T	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 4 GUIDE-seq
off target 7
untreated
BE1
BE2
BE3

961

G	G	C	A	C	T	G	C	T	A	C	T	G	G	G	G	T	G	G	T	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 4 GUIDE-seq
off target 8
untreated
BE1
BE2
BE3

769

G	G	C	A	C	T	G	G	G	G	C	T	G	G	G	G	A	G	G	G	G	G
---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---	---

HEK site 4 GUIDE-seq
off target 9
untreated
BE1
BE2
BE3

712

FIGURE 33 (CONTINUED)

GUIDE-seq
counts/ ChIP-seq
intensity
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HEK site 4 GUIDE-seq off target 10	G	G	C	A	C	C	T	T	G	G	G	G	T	T	G	G	A	G	G	T	G	G	G	G	G	664
untreated	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.6	0.6	0.6	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5	3.5
BE3	0.3	0.3	0.3	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0

HEK site 4 ChIP-seq off target 1	G	T	G	G	C	C	T	T	G	G	A	G	G	T	G	G	A	G	G	T	G	G	G	G	G	110
untreated	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

HEK site 4 ChIP-seq off target 3	G	A	G	G	G	G	A	A	A	G	G	G	C	T	G	G	A	G	G	T	G	G	A	G	G	89
untreated	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
BE3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FIGURE 33 (CONTINUED)

APOE4	C30	A49	C48	C47	T46	G45	C44	G43	C42	A41	A40	G39	C38	T37	G36	C35	G34	T33	A32	A31	G30	C29	G28	G27	C26	
Untreated																										
A	0.140.1	100±0.1	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1
C	100±0.1	0.140.1	100±0.1	0.140.1	0.140.1	100±0.1	0.140.1	100±0.1	0.140.1	0±0	0.140.1	0±0	100±0.1	0.140.1	0.140.1	0±0	0.140.1	0±0	100±0.1	0.140.1	0±0	100±0.1	0.140.1	0.140.1	0.140.1	100±0.1
G	0.140.1	0.140.1	0±0	0±0	0.140.1	100±0.1	0±0	99.9±0.1	0.140.1	0.140.1	0.140.1	100±0.1	0±0	0±0	100±0.1	0.140.1	100±0.1	0±0	0.140.1	0.140.1	100±0.1	0.140.1	100±0.1	100±0.1	0±0	0±0
T	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1
BEC + on-target sgRNA																										
A	0±0	100±0.1	0±0	0.140.1	0.140.1	0.140.1	0±0	0.140.1	0.140.1	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0±0	0.140.1	0±0	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1
C	100±0.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0±0	100±0.1	0.140.1	0.140.1	0±0	0.140.1	0±0	100±0.1	0.140.1	0±0	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1
G	0.140.1	0.140.1	0±0	0.140.1	0.140.1	100±0.1	0±0	0.140.1	0.140.1	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	100±0.1	100±0.1	0.140.1	0.140.1
T	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1
BEC + off-target sgRNA																										
A	0±0	100±0.1	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0±0	0.140.1	0.140.1	0.140.1	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1
C	100±0.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1
G	0.140.1	0.140.1	0±0	0±0	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	100±0.1	100±0.1	0.140.1	0.140.1
T	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	100±0.1	0.140.1	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1
BPOE4	T26	C24	C23	T22	C21	C20	G19	G18	G17	A16	T15	G14	C13	C12	G11	A10	T9	G8	A7	C6	C5	T4	G3	C2	A1	
Untreated																										
A	0.140.1	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
C	0.140.1	100±0	100±0.1	0.140.1	100±0.1	100±0.1	0±0	100±0.1	0.140.1	0±0	0.140.1	0±0	100±0.1	100±0.1	0±0	0±0	100±0.1	100±0.1	0.140.1	0.140.1	100±0.1	100±0.1	0.140.1	0.140.1	0.140.1	
G	0±0	0±0	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
T	100±0.1	0±0	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
BEC + on-target sgRNA																										
A	0±0	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
C	0.140.1	99.9±0.1	100±0.1	0.140.1	99.9±0.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
G	0±0	0.140.1	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
T	100±0.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
BEC + off-target sgRNA																										
A	0.140.1	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
C	0.140.1	100±0.1	100±0.1	0.140.1	100±0.1	100±0.1	0±0	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
G	0±0	0±0	0±0	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	
T	100±0.1	0.140.1	0.140.1	100±0.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	0.140.1	

FIGURE 34

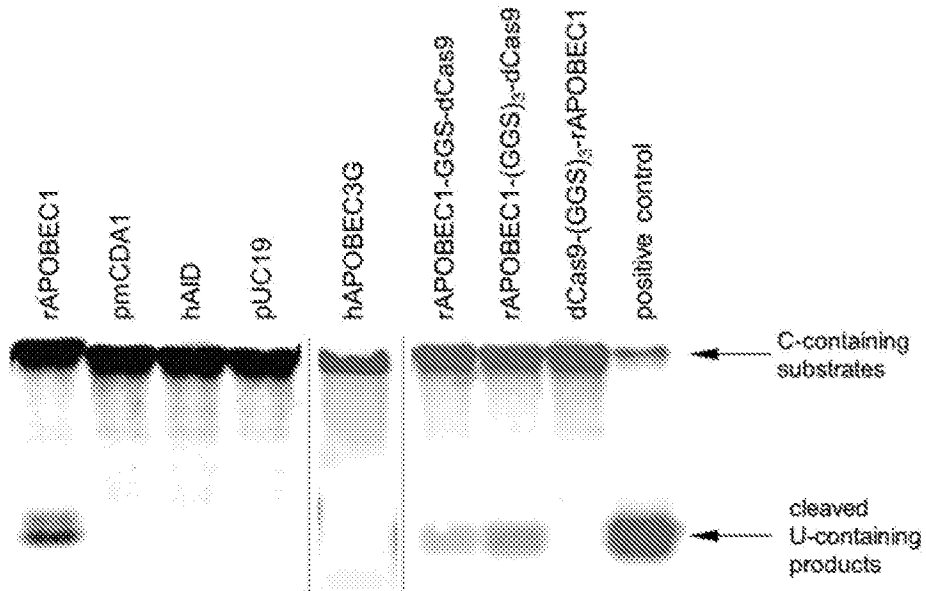


FIGURE 36A

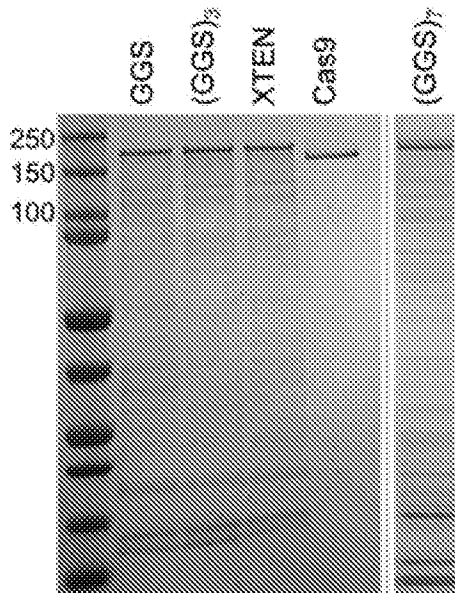


FIGURE 36B

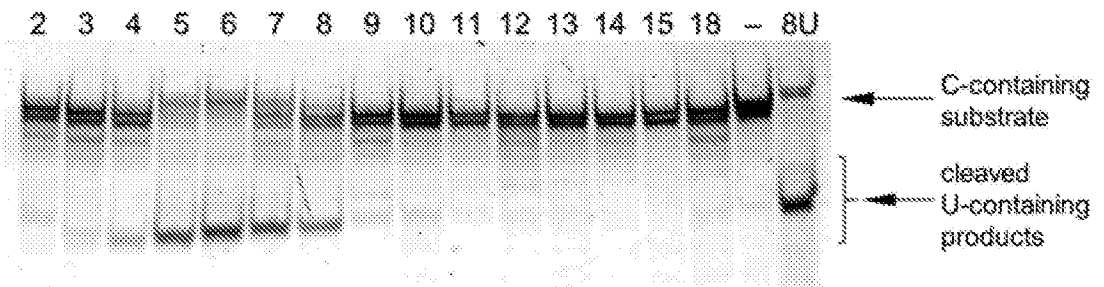


FIGURE 36C

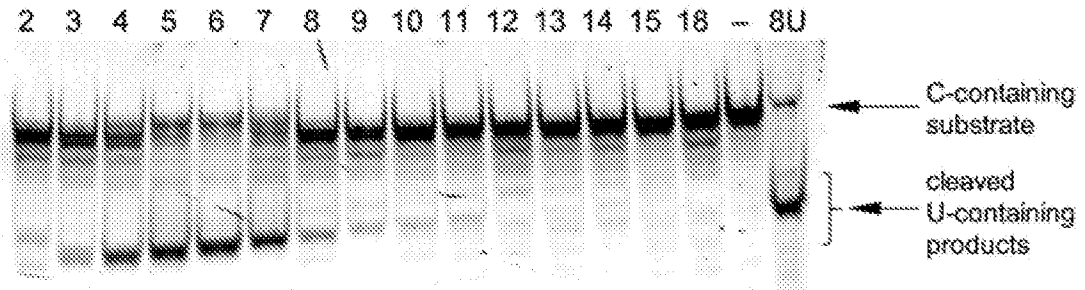


FIGURE 36D

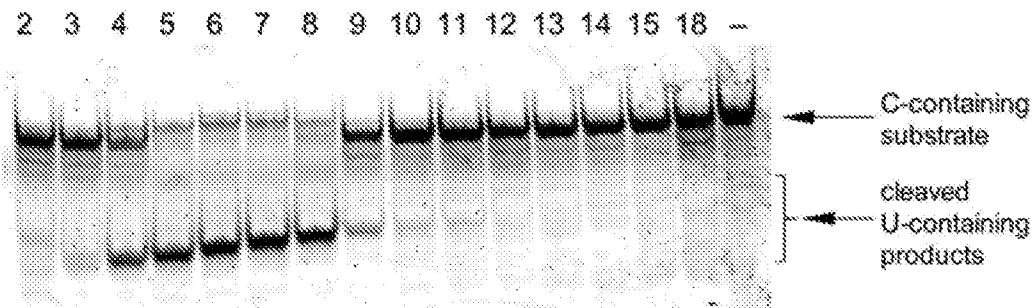


FIGURE 36E

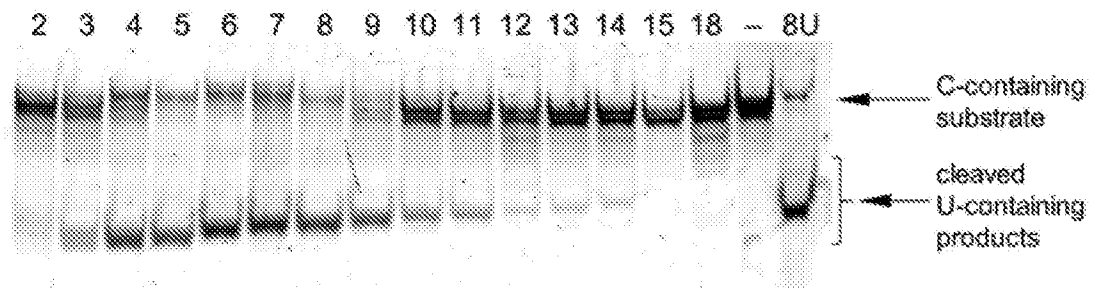


FIGURE 36F

EMX1: GAGTC₅C₆GAGCAGAAGAAGAAGGG
FANCF: GGAATC₆C₇C₈TTC₁₁TGCAGCACCTGG
HEK293 site 2: GAAC₄AC₆AAAGCATAGACTGCGGG
HEK293 site 3: GGCC₄C₅AGACTGAGCACGTGATGG
HEK293 site 4: GGCAC₅TGCGGCTGGAGGTCCGGG
RNF2: GTC₃ATC₆TTAGTC₁₂ATTACCTGAGG

FIGURE 37A

EMX1		G	A	G	T	C ₁	C ₂	G	A	G	C	A	G	A	G	A	A	G	A	G	A	A	G	G
1.85 uM BE1	A	0.2	99.9	0.1	0.1	0.1	0.0	0.3	100.0	0.1	0.1	99.9	0.2	99.9	99.7	0.3	100.0	99.9	0.2	99.9	97.9	0.1	0.2	0.2
	C	0.0	0.0	0.0	0.0	60.4	61.0	0.0	0.0	0.0	99.1	0.0	0.0	0.2	0.0	0.0	0.0	0.0	0.0	0.0	2.0	0.0	0.0	0.0
	G	99.7	0.0	99.9	0.0	0.0	0.0	99.7	0.0	99.9	0.1	0.0	99.7	0.0	0.0	99.8	0.0	0.0	99.8	0.0	0.0	99.5	99.8	99.8
	T	0.1	0.0	0.0	99.9	99.5	99.0	0.0	0.0	0.0	0.7	0.0	0.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FANCF		G	G	A	A	T	C ₁	C ₂	T	G	C	A	G	A	G	A	C	A	G	C	A	C	T	G	G
1.85 uM BE1	A	0.1	0.2	99.8	99.9	0.0	0.1	0.1	0.1	0.1	0.1	0.1	99.9	0.2	99.9	0.2	95.0	0.1	0.1	0.1	0.1	0.1	0.1	0.3	0.4
	C	0.0	0.0	0.1	0.1	0.0	63.9	64.7	65.0	0.0	0.0	72.6	0.0	99.4	0.1	0.0	99.7	4.9	99.8	99.9	0.0	0.0	0.0	0.0	0.0
	G	99.8	99.7	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.8	0.2	0.0	99.7	0.2	0.0	0.1	0.0	0.0	0.0	0.0	99.5	99.5
	T	0.1	0.1	0.0	0.0	99.9	36.0	35.1	34.9	99.9	99.9	27.3	99.8	0.1	0.3	0.0	0.1	0.0	0.0	0.0	0.0	0.0	99.9	0.1	0.1

HEK293 site 2		G	A	A	C ₁	A	C ₂	A	A	A	G	C	A	T	A	G	A	C	T	G	C	A	G	G		
1.85 uM BE1	A	0.1	100.0	99.8	0.1	100.0	0.1	99.9	100.0	99.9	0.1	0.1	100.0	0.0	99.9	0.2	100.0	0.1	0.0	0.1	0.1	0.1	0.1	0.1	0.2	0.2
	C	0.0	0.0	0.1	80.6	0.0	76.9	0.0	0.0	0.0	0.0	99.6	0.0	0.0	0.1	0.0	0.0	99.9	0.0	0.0	99.9	0.0	0.0	0.0	0.0	
	G	99.8	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.8	0.0	0.1	0.0	99.8	0.0	0.0	0.0	0.0	0.0	99.8	0.0	99.9	99.8	99.8
	T	0.1	0.0	0.0	19.3	0.0	22.9	0.0	0.0	0.0	0.1	0.3	0.0	99.9	0.0	0.1	0.0	0.0	99.9	0.0	0.1	0.0	0.1	0.0	0.1	0.0

HEK293 site 3		G	G	C	C ₁	C ₂	A	G	A	C	T	G	A	G	A	C	G	T	G	A	T	G	A	T	G	G
1.85 uM BE1	A	0.2	0.1	0.1	0.1	0.0	99.9	0.2	99.9	0.1	0.0	0.2	99.9	0.1	0.1	100.0	0.1	0.1	0.0	0.2	99.8	0.0	0.1	0.1	0.2	0.1
	C	0.0	0.0	92.2	74.8	71.5	0.0	0.0	0.0	96.6	0.0	0.0	99.8	0.0	0.0	99.7	0.0	0.0	0.0	0.0	1.3	0.0	0.0	0.0	0.0	0.0
	G	99.7	99.8	0.0	0.0	0.0	0.0	99.8	0.0	0.0	0.0	99.8	0.0	99.8	0.0	0.0	99.8	0.0	99.8	0.0	0.0	0.0	99.9	99.8	99.8	99.8
	T	0.1	0.1	7.7	25.1	28.5	0.0	0.0	0.0	3.3	99.9	0.1	0.0	0.0	0.1	0.0	0.2	0.0	99.9	0.0	0.0	100.0	0.0	0.0	0.0	0.0

HEK293 site 4		G	G	C	A	C ₁	A	T	G	C	G	G	C	T	G	A	G	G	T	G	A	T	G	G	G	G
1.85 uM BE1	A	0.3	0.2	0.1	100.0	0.1	0.0	0.2	0.1	0.2	0.1	0.0	0.3	0.5	100.0	0.1	0.1	0.0	0.1	0.1	0.1	0.1	0.1	0.1	0.2	0.2
	C	0.0	0.0	96.6	0.0	60.1	0.0	0.0	97.0	0.0	0.0	99.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
	G	99.7	99.7	0.0	0.0	0.0	0.0	99.8	0.0	99.8	99.8	0.0	0.0	99.7	99.4	0.0	99.8	99.8	0.1	99.8	99.8	99.8	99.8	99.8	99.8	99.7
	T	0.0	0.1	1.1	0.0	39.8	100.0	0.1	2.9	0.0	0.1	0.5	100.0	0.0	0.1	0.0	0.0	0.0	99.9	0.0	0.1	0.1	0.1	0.1	0.0	0.0

RNF2		G	T	C ₁	A	T	C ₂	T	A	G	T	C ₂	A	T	A	C	T	G	A	C	T	G	A	G	G	G	
1.85 uM BE1	A	0.1	0.0	0.1	99.9	0.0	0.0	0.0	99.9	0.1	0.0	0.1	99.9	0.0	0.0	99.9	0.1	0.0	0.1	99.9	0.0	0.0	0.1	99.9	0.0	0.0	0.0
	C	0.0	0.0	59.1	0.0	0.0	57.8	0.0	0.0	0.1	0.0	86.4	0.0	0.0	0.0	0.1	99.9	99.9	0.0	0.0	0.0	0.0	1.4	0.0	0.0	0.0	
	G	99.9	0.1	0.0	0.0	0.0	0.0	0.0	99.8	0.0	0.0	99.8	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.8	0.0	99.9	99.9	99.9	
	T	0.0	99.9	40.8	0.0	100.0	42.1	100.0	100.0	0.0	0.1	99.9	13.4	0.0	100.0	99.9	0.0	0.0	99.9	0.1	0.0	99.9	0.1	0.0	0.1	0.0	

FIGURE 37B

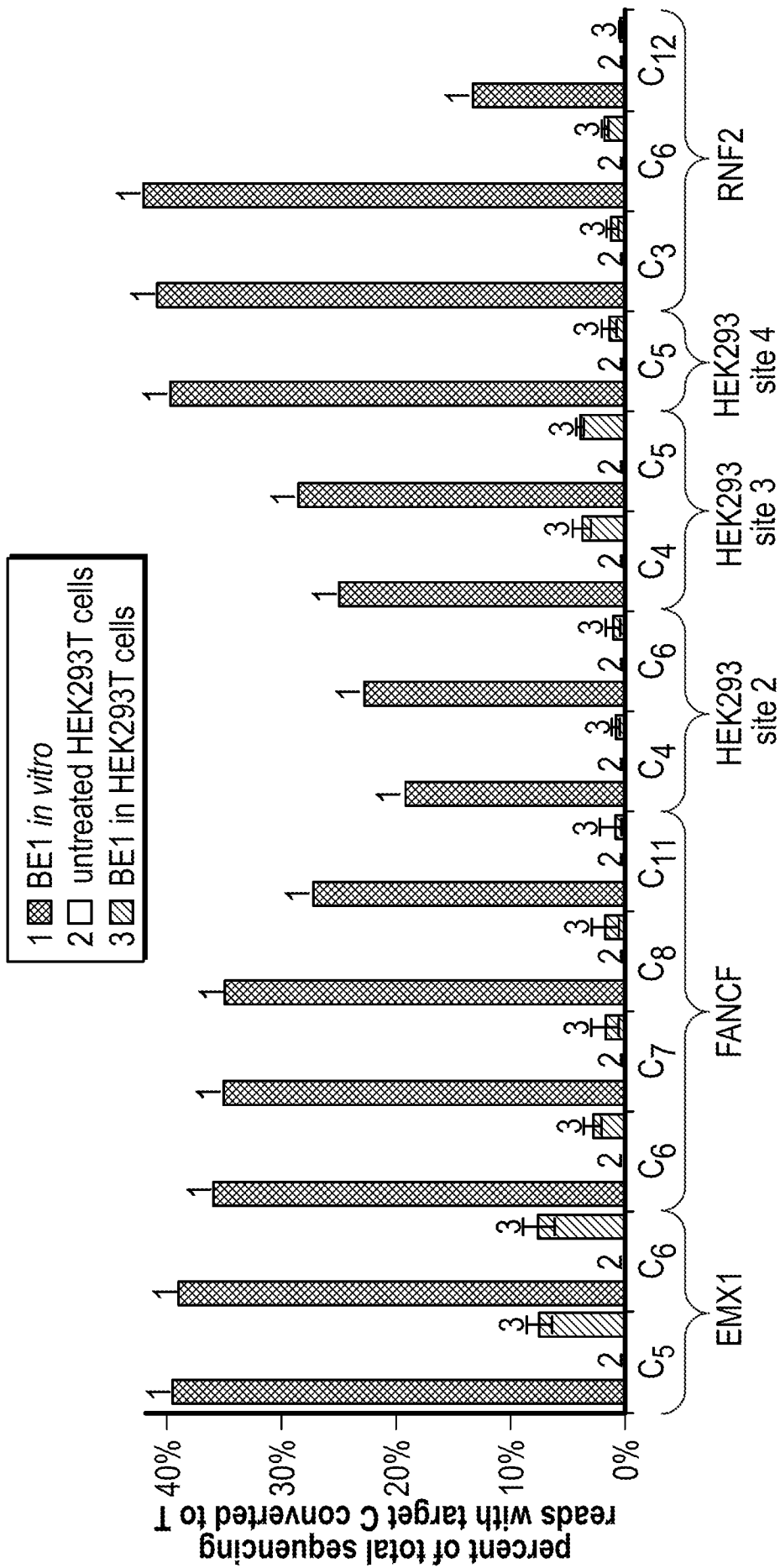


FIGURE 37C

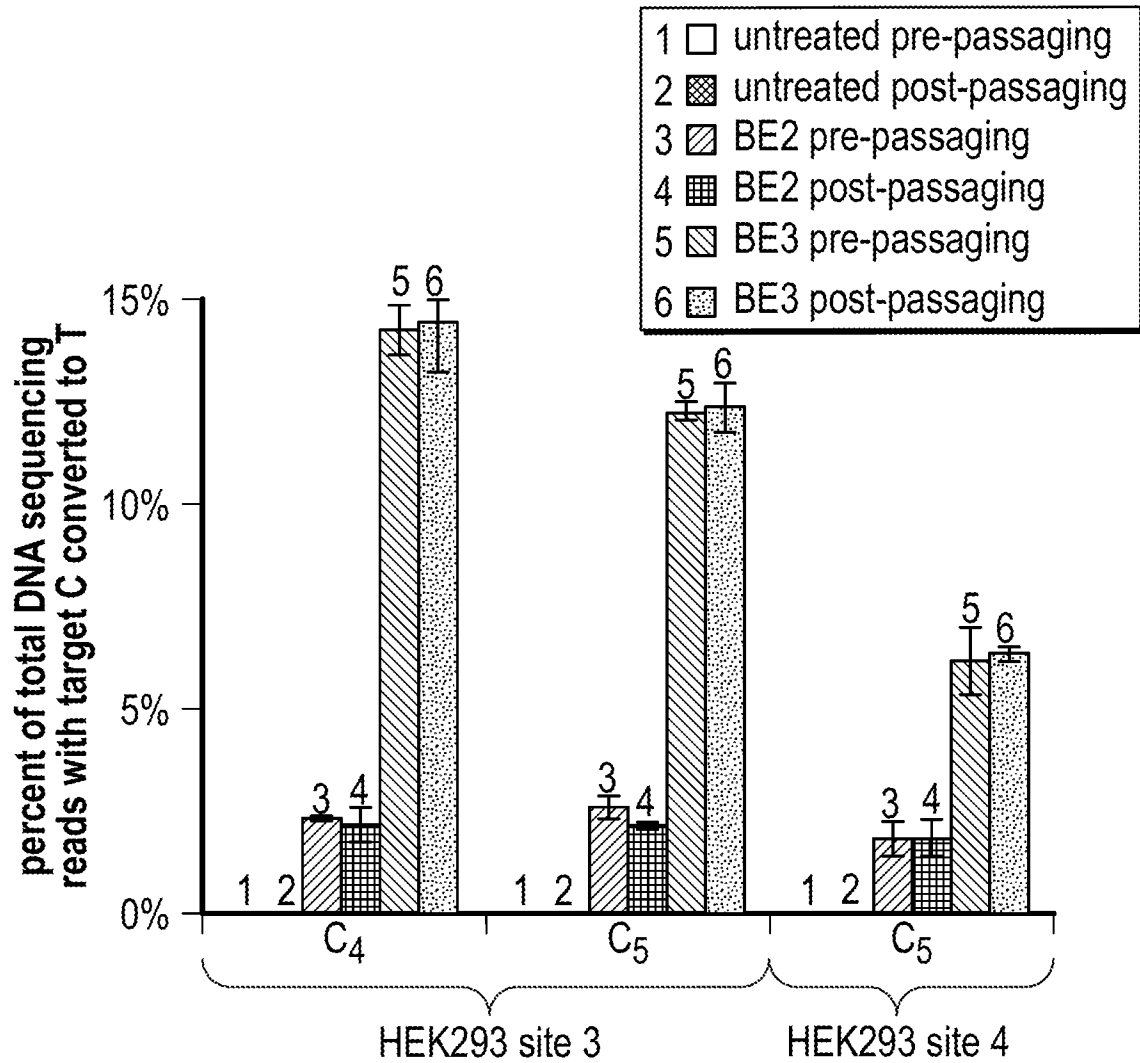


FIGURE 38

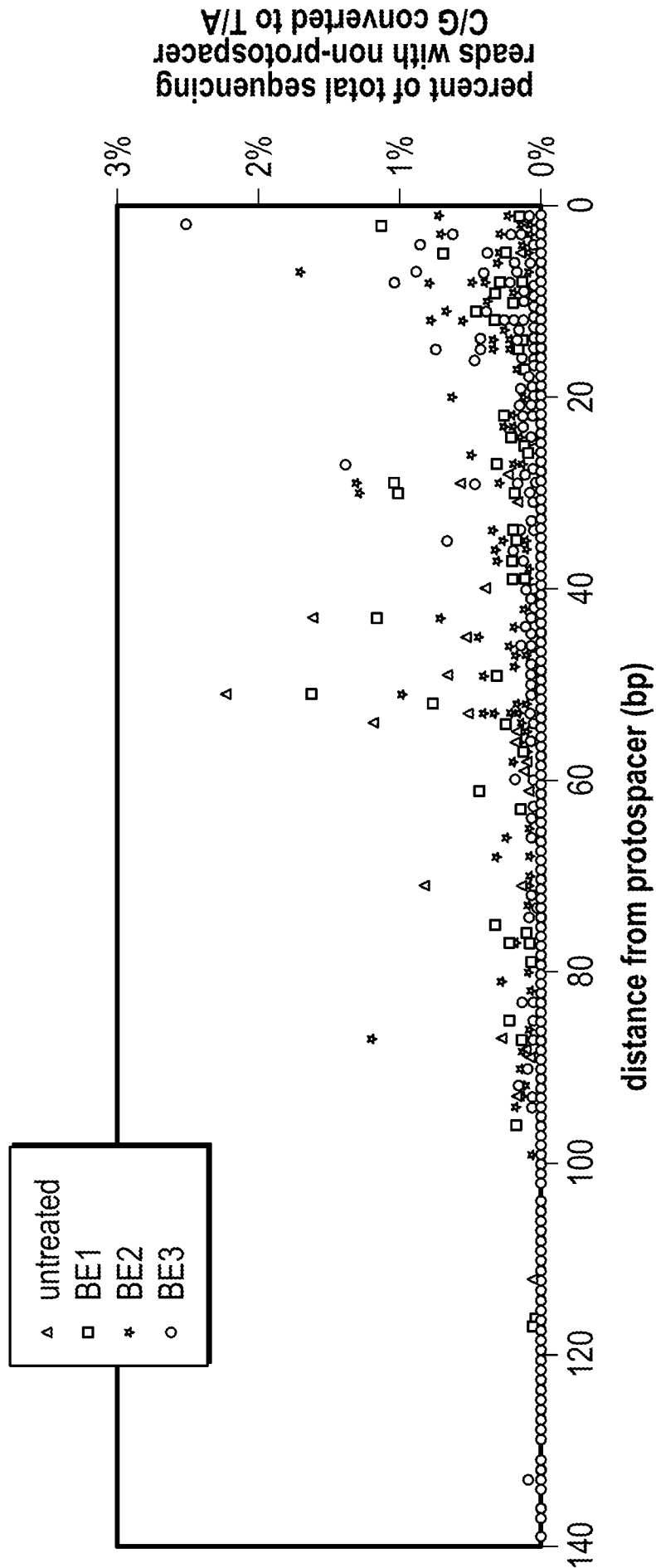


FIGURE 39A

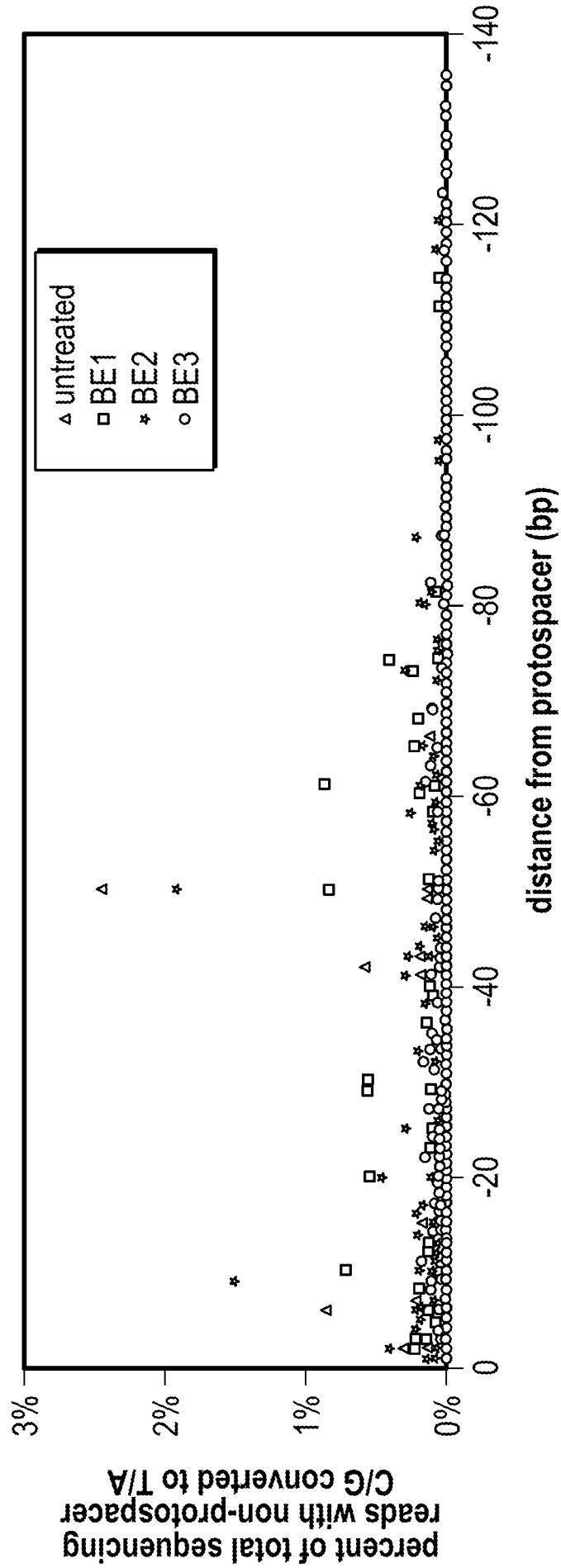


FIGURE 39B

non-protospacer C/Gs	average C/G (%)	average T/A (%)	lowest T/A (%)	highest T/A (%)
untreated	99.95 ± 0.14	0.02 ± 0.02	0.00	2.44
BE1	99.95 ± 0.24	0.03 ± 0.03	0.00	1.64
BE2	99.95 ± 0.13	0.03 ± 0.03	0.00	1.92
BE3	99.97 ± 0.09	0.02 ± 0.02	0.00	2.52

FIGURE 39C

untreated APOE4C158R	Lys			Arg			Leu			Ala			Val			Tyr			Gln			indel %	
	A	A	G	C ₄	G	C	C	T	G	C	C	A	G	T	G	A	T	A	C	C	A		
	G	A	G	C ₄	G	C	C	T	G	C	C	A	G	T	G	A	T	A	C	C	A		
A	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	100.0	0.0	0.0
C	0.0	0.0	0.0	100.0	0.0	100.0	100.0	0.0	0.1	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	100.0	0.0	0.0	0.0
G	100.0	0.0	0.0	99.9	0.0	100.0	0.0	0.0	99.9	0.0	0.0	0.0	100.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	99.9	100.0
T	0.0	0.0	0.1	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1

BE3 + on-target sgRNA APOE4C158R	Lys			Arg → Cys			Leu → Leu			Ala			Val			Tyr			Gln			indel %	
	A	A	G	C ₄	G	C	C	T	G	C	C	A	G	T	G	A	T	A	C	C	A		
	G	A	G	C ₄	G	C	C	T	G	C	C	A	G	T	G	A	T	A	C	C	A		
A	0.1	100.0	0.0	1.0	0.0	1.6	0.6	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	100.0	0.0	0.1
C	0.0	0.0	0.0	39.1	0.0	62.0	61.3	0.0	0.0	0.0	0.0	99.9	0.0	0.0	0.0	0.0	0.0	0.0	100.0	100.0	0.0	0.0	0.0
G	99.9	0.0	0.0	100.0	1.5	99.9	0.7	0.5	0.0	100.0	100.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.9	
T	0.0	0.0	0.0	59.1	0.1	35.7	37.5	100.0	0.0	0.0	0.1	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

BE3 + off-target sgRNA APOE4C158R	Lys			Arg			Leu			Ala			Val			Tyr			Gln			indel %	
	A	A	G	C ₄	G	C	C	T	G	C	C	A	G	T	G	A	T	A	C	C	A		
	G	A	G	C ₄	G	C	C	T	G	C	C	A	G	T	G	A	T	A	C	C	A		
A	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	100.0	0.0	0.0
C	0.0	0.0	0.0	99.9	0.0	100.0	100.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	100.0	0.0	0.0	0.0
G	100.0	0.0	0.0	100.0	0.0	100.0	0.0	0.0	100.0	100.0	0.0	0.0	100.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	100.0
T	0.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

Cas9 + HDR APOE4C158R	Lys			Arg → Cys			Leu			Ala			Val			Tyr			Gln			indel %	
	A	A	G	C ₄	G	C	C	T	G	C	C	A	G	T	G	A	T	A	C	C	A		
	G	A	G	C ₄	G	C	C	T	G	C	C	A	G	T	G	A	T	A	C	C	A		
A	0.0	99.7	99.7	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0	98.0	0.0	0.0	0.0	0.0	0.0	99.1	0.1	0.5	98.8	0.0	0.0
C	0.0	0.3	0.2	0.0	99.8	0.0	99.8	99.9	0.1	0.1	0.2	99.9	1.8	0.4	0.2	0.3	0.2	0.5	99.7	99.6	0.7	0.0	0.0
G	99.7	0.0	0.0	99.9	0.0	99.7	0.1	0.0	0.1	99.8	99.7	0.0	0.1	99.6	1.4	99.4	0.2	0.3	0.1	0.0	0.0	99.4	100.0
T	0.2	0.0	0.1	0.1	0.1	0.3	0.1	0.1	99.8	0.2	0.1	0.1	1.1	99.3	0.4	99.7	0.2	0.2	0.0	0.0	0.5	0.5	0.0

FIGURE 40A

untreated TP53Y163C	Arg			Ala			Met			Ala			Ile			Cys			Lys			indel %
	C	C	G	C	C	G	A	T	G	C	C	G	A	T	C	T	G	C	A	A	G	
	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
A	100.0	100.0	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
G	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
BE3 + on-target sgRNA TP53Y163C	Arg			Ala			Met			Ala			Ile			Cys → Tyr			Lys			indel %
C	C	G	C	C	G	A	T	G	C	C	G	A	T	C	T	G	C	A	A	G		
0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.5	0.0	0.0	0.0		
A	100.0	100.0	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
G	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.5	0.0	0.0	0.0	
BE3 + off-target sgRNA TP53Y163C	Arg			Ala			Met			Ala			Ile			Cys			Lys			indel %
C	C	G	C	C	G	A	T	G	C	C	G	A	T	C	T	G	C	A	A	G		
0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0		
A	100.0	100.0	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
G	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
Cas9 + HDR TP53Y163C	Arg			Ala			Met			Ala			Ile			Cys → Tyr			Lys			indel %
C	C	G	C	C	G	A	T	G	C	C	G	A	T	C	T	G	C	A	A	G		
0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0		
A	100.0	100.0	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
G	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	

FIGURE 40B

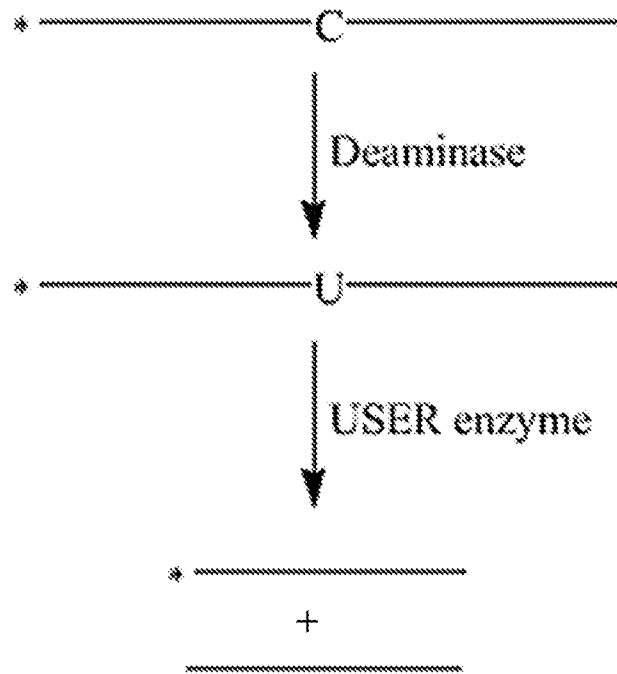
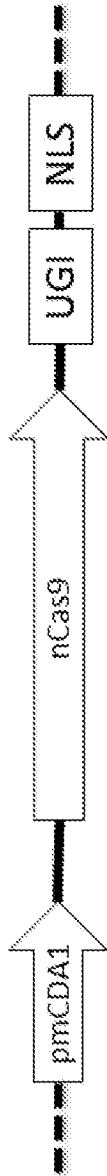


FIGURE 41



P A M 20 19 18 17 16 15 14 13 12 11 10 9 8 7 6 5 4 3 2 1 -1 -2 -3
 T G C C A T C A C G T G C T C A A T C G G G G C C C C A

CDA	A	C	G	T	A	M	20	19	18	17	16	15	14	13	12	11	10	9	8	7	6	5	4	3	2	1	-1	-2	-3			
A	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.5%	0.0%	0.0%	0.1%	100.0%	5.8%	0.0%	0.0%	0.0%	0.0%	33.9%	33.9%	33.1%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	
C	0.0%	0.0%	99.9%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	99.9%	0.0%	1.0%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	100.0%	100.0%	99.9%	99.9%	0.0%	0.0%		
G	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	99.5%	0.0%	0.0%	0.0%	0.0%	91.8%	0.0%	0.0%	0.0%	0.0%	65.7%	65.0%	65.2%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
T	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	1.5%	100.0%	0.0%	100.0%	0.3%	1.1%	1.7%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%		
untreated	A	C	G	T	A	M	20	19	18	17	16	15	14	13	12	11	10	9	8	7	6	5	4	3	2	1	-1	-2	-3			
A	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.1%	100.0%	0.8%	0.0%	0.0%	0.0%	0.0%	38.5%	41.9%	1.4%	0.0%	0.0%	0.1%	0.0%	0.0%	100.0%	
C	0.0%	0.0%	99.9%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	99.9%	0.0%	0.1%	0.0%	99.9%	0.0%	0.0%	8.1%	0.7%	0.0%	100.0%	99.9%	99.9%	100.0%	0.0%	0.0%	
G	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	99.0%	0.0%	0.0%	0.0%	0.0%	51.3%	56.7%	98.6%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
T	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	100.0%	2.2%	0.7%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
A	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.1%	100.0%	0.0%	0.0%	0.0%	0.1%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	
C	0.0%	0.0%	99.9%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	99.9%	0.0%	0.0%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	100.0%	99.9%	100.0%	0.0%	0.0%	
G	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	99.9%	99.9%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%
T	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%

FIGURE 42



Hek-2 site, shown: coding strand (G→A), target: template stand (C→T)

	P	A	M	20	C	C	G	C	A	G	T	C	T	A	T	G	C	T	T	T	G	T	G	T	3	2	1	-1	-2	-3		
C	G	C	C	C	G	C	C	A	G	T	C	T	C	T	A	T	G	C	T	T	T	G	T	G	T	T	C	C	C	A	G	
A	0.0%	0.0%	0.2%	0.0%	0.5%	0.2%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.9%	0.8%		
C	99.9%	0.0%	100.0%	99.7%	99.9%	99.5%	99.9%	0.0%	99.9%	0.0%	0.0%	99.9%	0.0%	0.0%	0.5%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.9%	99.6%	0.0%	0.3%	
G	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	80.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	98.8%	0.1%	
T	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	100.0%	0.3%	0.0%	0.0%	0.0%	100.0%	100.0%	100.0%	2.4%	100.0%	8.0%	100.0%	100.0%	0.2%	0.4%	0.0%	0.0%	0.0%	0.1%	
TAPOBEC1																																
A	0.0%	0.0%	0.0%	0.0%	0.0%	0.1%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.7%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.9%	0.0%	0.0%	
C	99.9%	0.0%	100.0%	100.0%	99.9%	99.9%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.9%	99.9%	0.0%	0.0%
G	0.0%	99.9%	0.0%	0.0%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.3%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.7%	0.3%	
T	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	100.0%	100.0%	100.0%	0.7%	100.0%	1.1%	100.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.3%	
untreated																																
A	0.0%	0.0%	0.0%	0.0%	0.0%	0.1%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.9%	0.0%	0.0%	
C	99.9%	0.0%	100.0%	100.0%	99.9%	99.9%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.9%	99.9%	0.0%	0.0%
G	0.0%	99.9%	0.0%	0.0%	0.0%	99.9%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	99.9%	0.0%	0.0%
T	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	100.0%	0.0%	100.0%	0.0%	0.0%	0.0%	0.0%	100.0%	100.0%	100.0%	0.0%	100.0%	0.0%	100.0%	100.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%	0.0%

FIGURE 43

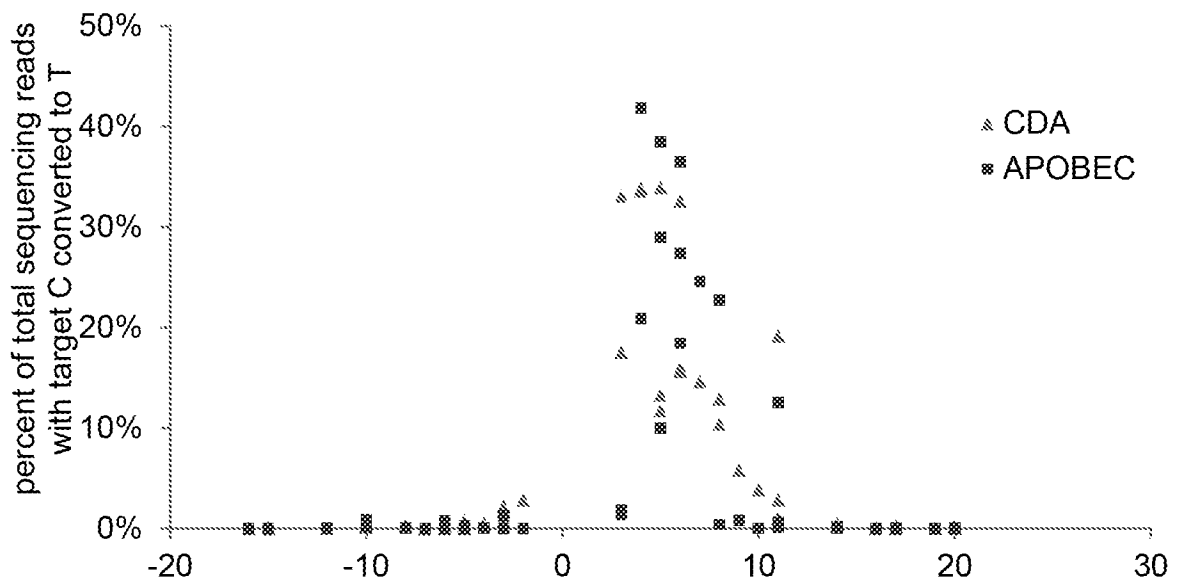


FIGURE 44

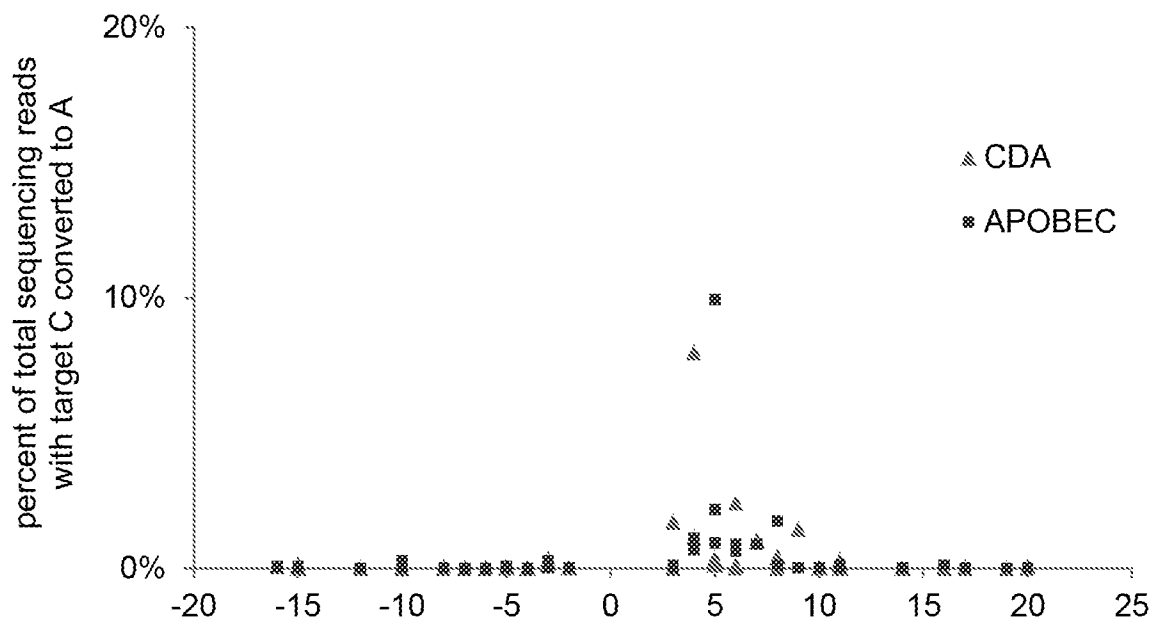


FIGURE 45

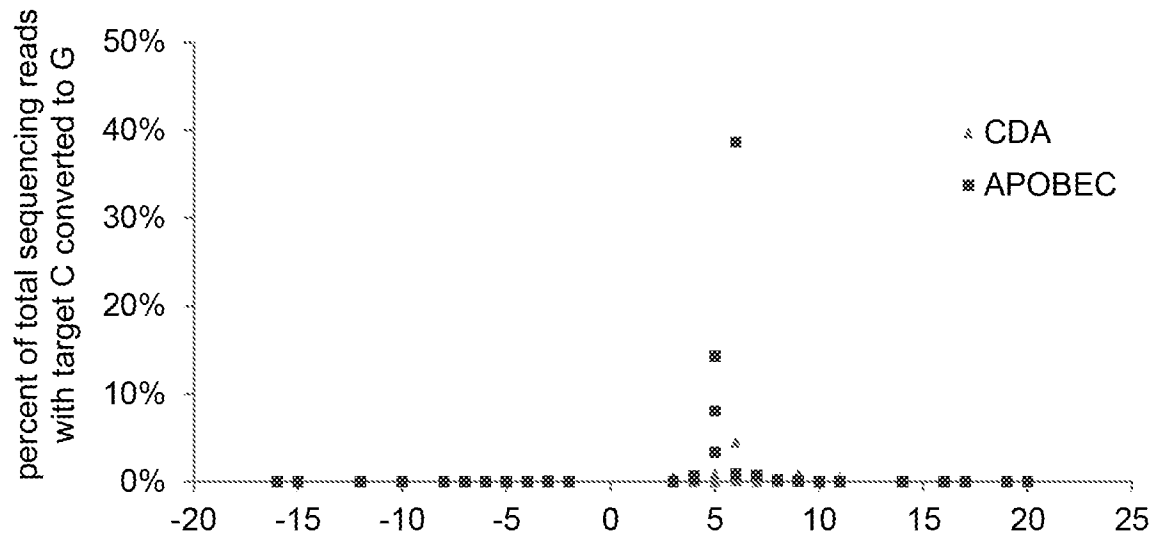
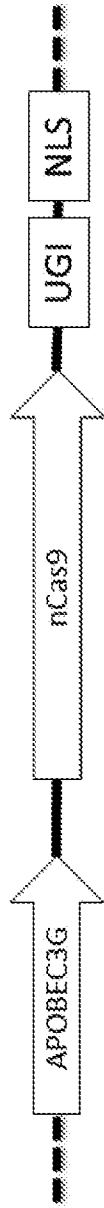
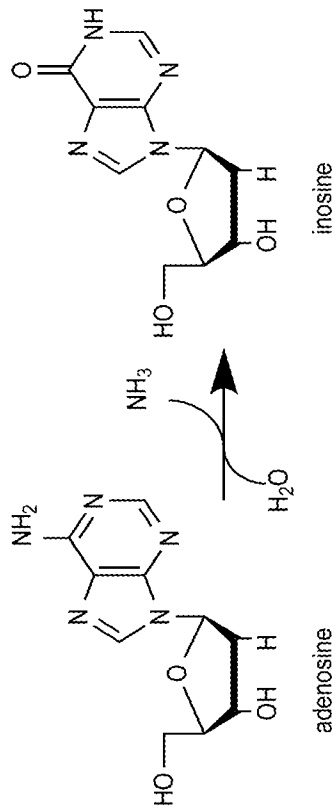


FIGURE 46

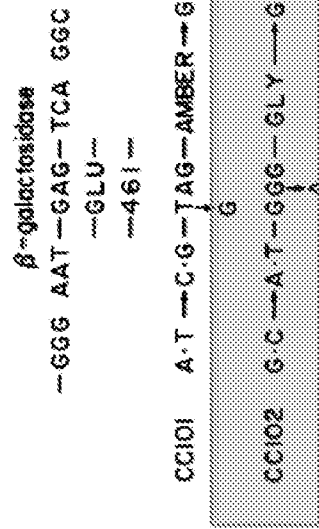


	P	A	M	20	19	18	17	16	15	14	13	12	11	10	9	8	7	6	5	4	3	2	1	
	C	C	C	G	C	A	G	T	C	T	A	T	G	C	T	T	T	G	T	G	T	T	C	
TAPOBEC1																								
A	0.02%	0.02%	0.01%	0.02%	0.02%	0.02%	0.05%	0.01%	0.01%	99.95%	0.01%	0.00%	0.01%	0.02%	0.01%	0.00%	0.00%	0.00%	0.01%	9.30%	0.01%	10.87%	0.00%	0.01%
C	99.96%	0.00%	99.98%	99.97%	99.97%	99.97%	0.00%	99.98%	0.01%	0.01%	0.01%	0.01%	99.98%	0.01%	0.01%	0.01%	0.01%	18.14%	0.01%	18.14%	0.01%	0.54%	0.01%	0.01%
G	0.00%	99.96%	0.00%	0.00%	0.00%	0.01%	99.86%	0.00%	0.00%	0.01%	0.00%	99.96%	0.00%	0.00%	0.00%	0.00%	0.00%	72.40%	0.00%	72.40%	0.00%	88.10%	0.00%	0.01%
T	0.02%	0.02%	0.01%	0.01%	0.01%	0.01%	0.03%	99.99%	0.01%	99.99%	0.02%	99.98%	0.01%	0.01%	99.99%	99.98%	99.99%	0.17%	99.98%	0.49%	99.98%	99.98%	99.98%	0.03%
APOBEC3G																								
A	0.01%	0.01%	0.01%	0.02%	0.01%	0.01%	0.05%	0.01%	0.01%	99.97%	0.01%	0.00%	0.01%	0.01%	0.00%	0.01%	0.00%	2.18%	0.01%	0.34%	0.00%	0.00%	0.01%	0.01%
C	99.97%	0.01%	99.98%	99.98%	99.96%	99.98%	0.00%	99.99%	0.01%	99.99%	0.01%	0.00%	99.98%	0.01%	0.01%	0.01%	0.01%	5.34%	0.01%	0.15%	0.01%	0.02%	99.98%	99.98%
G	0.00%	99.97%	0.00%	0.00%	0.00%	0.01%	99.98%	0.00%	0.00%	0.00%	0.00%	99.98%	0.00%	0.00%	0.00%	0.00%	0.00%	92.44%	0.00%	99.46%	0.00%	0.00%	0.00%	0.00%
T	0.02%	0.01%	0.01%	0.02%	0.01%	0.00%	0.01%	99.99%	0.01%	99.99%	0.02%	99.98%	0.01%	0.01%	99.99%	99.98%	99.99%	0.04%	99.98%	0.05%	99.98%	99.98%	99.98%	0.01%
APOBEC3G_RR																								
A	0.01%	0.02%	0.01%	0.01%	0.02%	0.01%	0.05%	0.01%	0.01%	99.97%	0.01%	0.04%	0.01%	0.01%	0.00%	0.01%	0.00%	2.18%	0.01%	6.12%	0.00%	0.00%	0.01%	0.01%
C	99.97%	0.00%	99.98%	99.99%	99.97%	99.97%	0.00%	99.98%	0.01%	99.99%	0.01%	0.01%	99.98%	0.01%	0.01%	0.01%	0.01%	4.98%	0.01%	2.49%	0.01%	0.02%	99.99%	99.99%
G	0.00%	99.96%	0.00%	0.00%	0.00%	0.01%	99.92%	0.00%	0.00%	0.00%	99.94%	0.00%	99.94%	0.00%	0.00%	0.00%	0.00%	92.80%	0.00%	91.04%	0.00%	0.00%	0.00%	0.00%
T	0.02%	0.02%	0.01%	0.01%	0.02%	0.01%	0.02%	99.99%	0.01%	99.99%	0.01%	99.98%	0.01%	0.01%	99.99%	99.99%	99.98%	0.05%	99.98%	0.35%	99.98%	99.98%	99.98%	0.01%
untreated																								
A	0.02%	0.02%	0.00%	0.01%	0.02%	0.01%	0.05%	0.01%	0.01%	99.98%	0.01%	0.00%	0.01%	0.00%	0.00%	0.00%	0.00%	0.01%	0.01%	0.01%	0.00%	0.00%	0.01%	0.01%
C	99.96%	0.00%	99.99%	99.96%	99.97%	99.97%	0.00%	99.98%	0.01%	99.99%	0.01%	0.00%	99.98%	0.01%	0.01%	0.01%	0.01%	0.01%	0.01%	0.01%	0.00%	0.01%	0.01%	99.98%
G	0.00%	99.96%	0.00%	0.00%	0.00%	0.01%	99.91%	0.00%	0.00%	0.00%	99.98%	0.00%	99.98%	0.00%	0.00%	0.00%	0.00%	99.96%	0.00%	99.95%	0.00%	0.00%	0.00%	0.00%
T	0.02%	0.02%	0.01%	0.01%	0.02%	0.01%	0.03%	0.01%	0.01%	99.99%	0.02%	99.98%	0.01%	0.01%	99.99%	99.99%	99.99%	0.02%	99.98%	0.02%	99.98%	99.98%	99.98%	0.01%

FIGURE 47



Utilizing a bacterial strain with lacZ encoded on the F plasmid
 Critical glutamic acid residue is mutated. Reversion restores lacZ activity.



Strain used to develop selection assay
 Constructs used: APOBEC1 and CDA

FIGURE 48

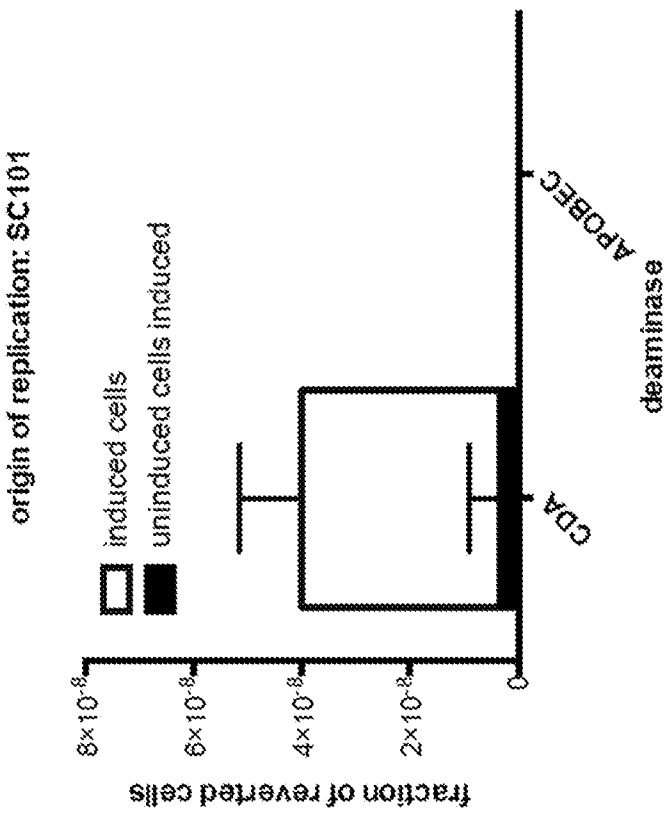
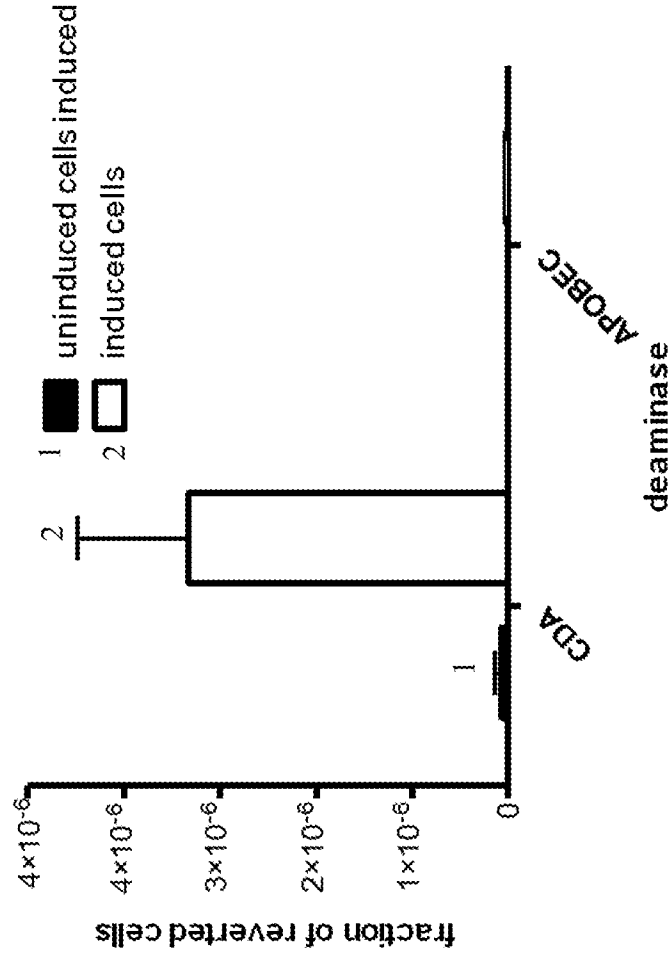


FIGURE 49

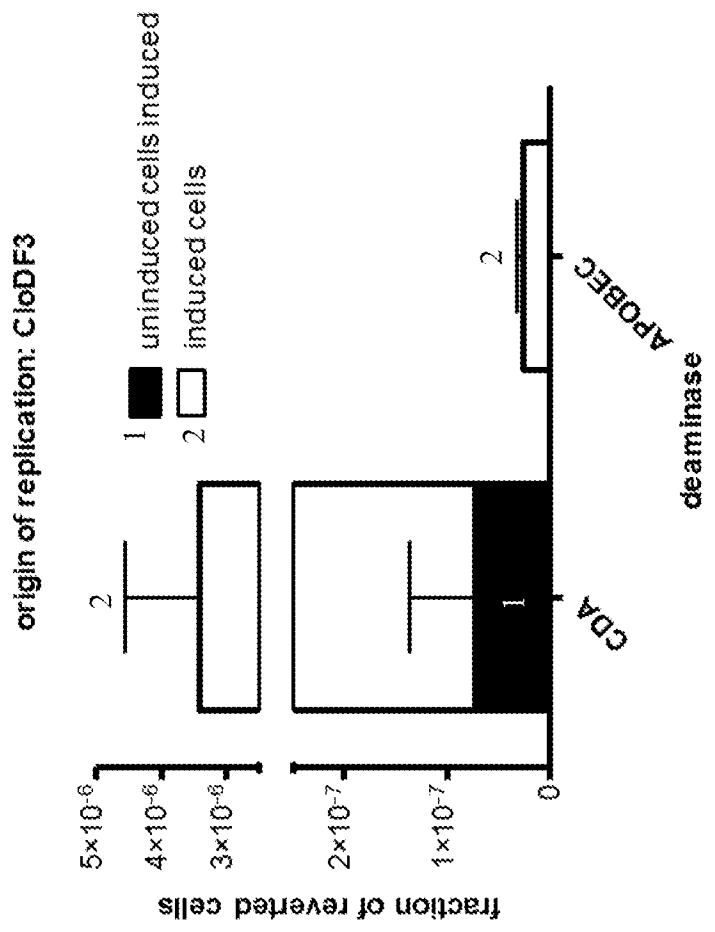
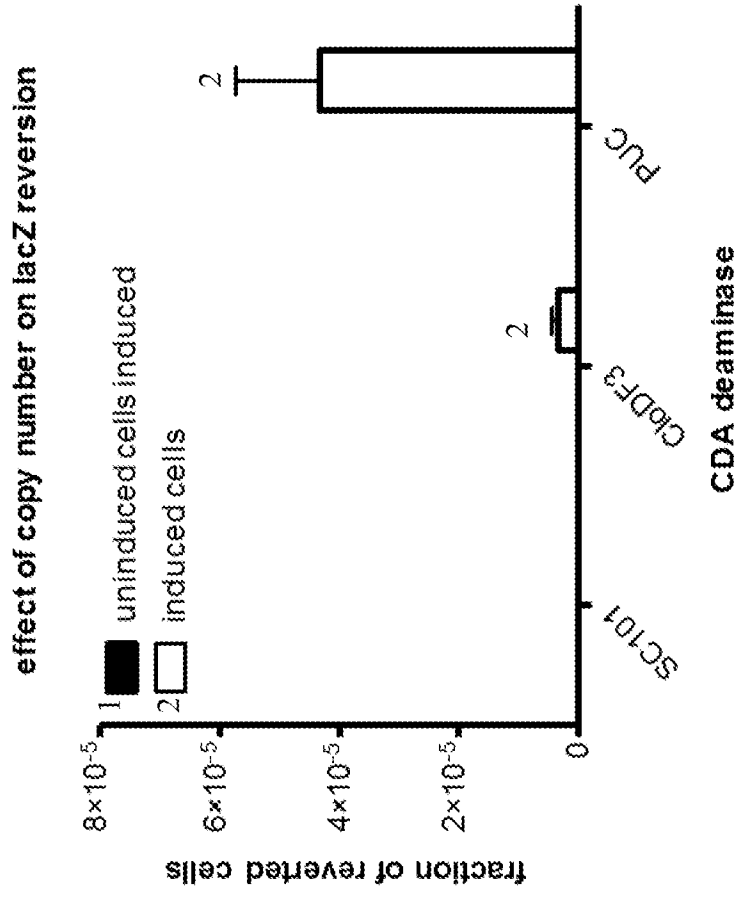


FIGURE 49 (CONTINUED)

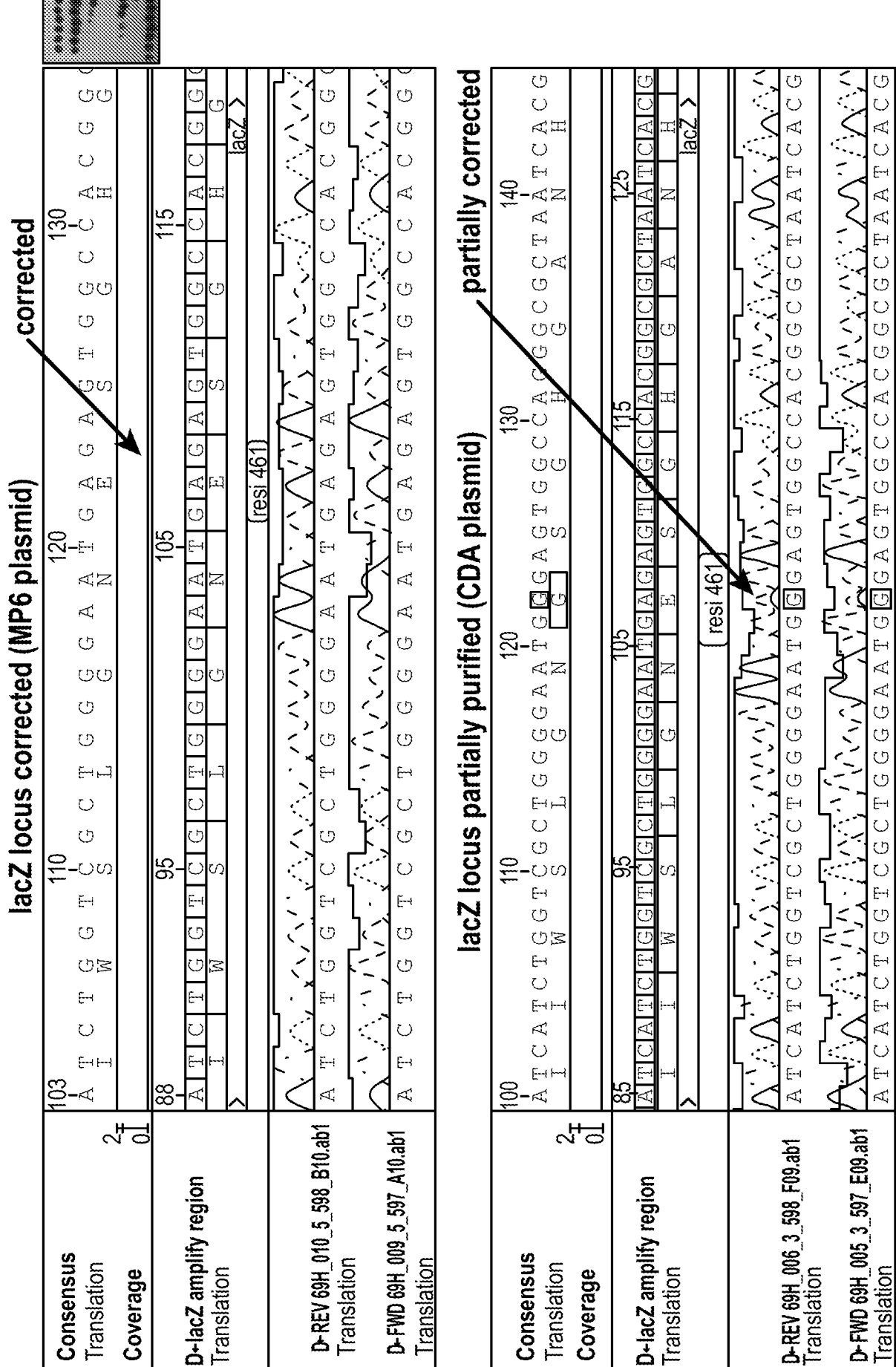


FIGURE 50

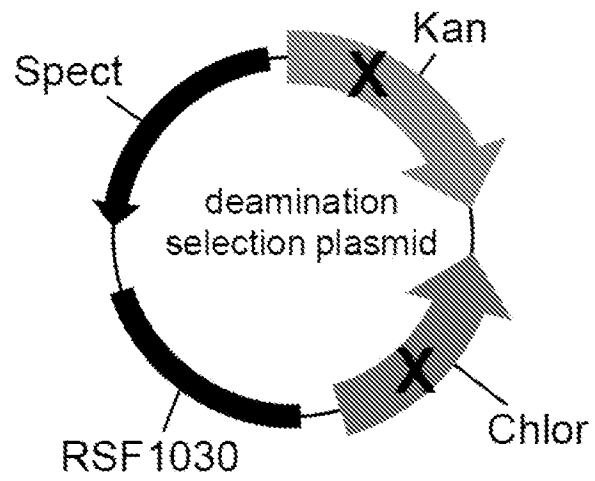
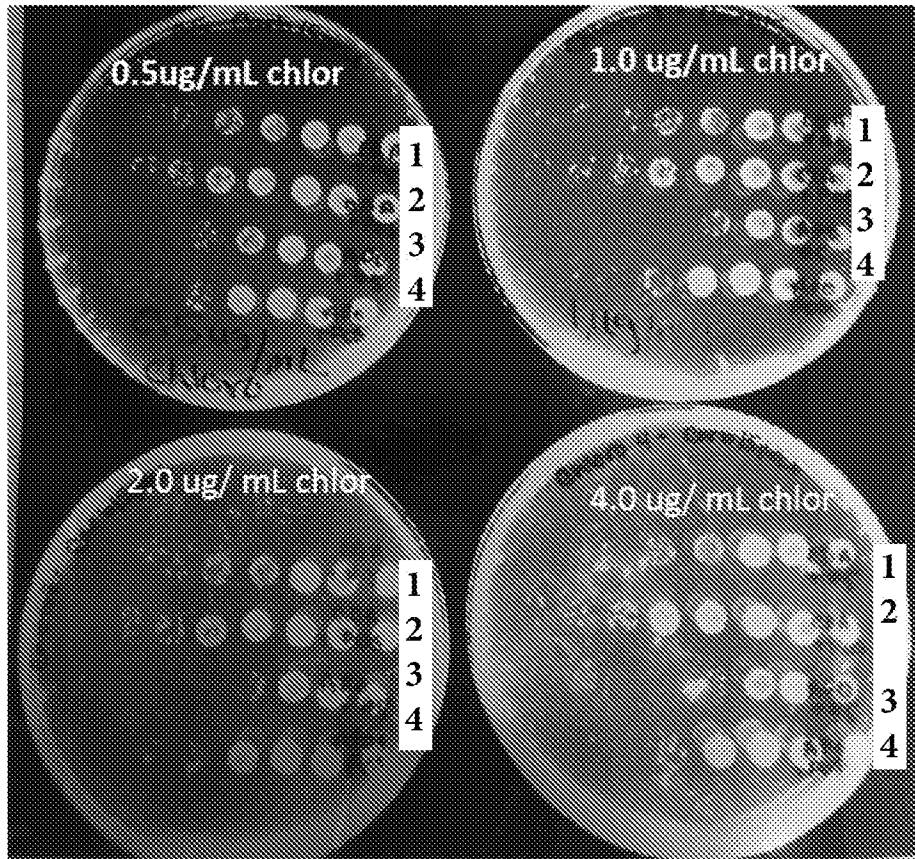
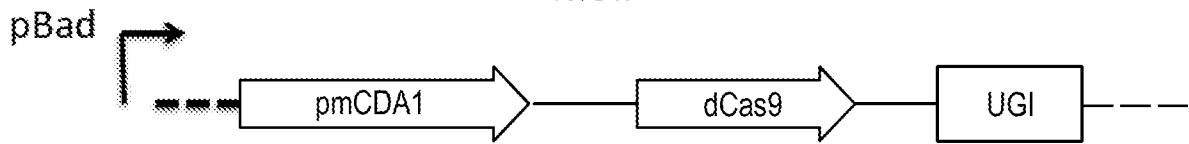


FIGURE 51



Row 1: CDA-dCas9 + selection plasmid (chlor^S)

Row 2: CDA-dCas9 + pos. control selection (chlor^R)

Row 3: rAPOBEC-dCas9 + selection plasmid (chlor^S)

Row 4: rAPOBEC-dCas9 + pos. control selection (chlor^R)

FIGURE 52

rAPOBEC-XTEN-dCas9-UGI on 8 ug/mL chlor

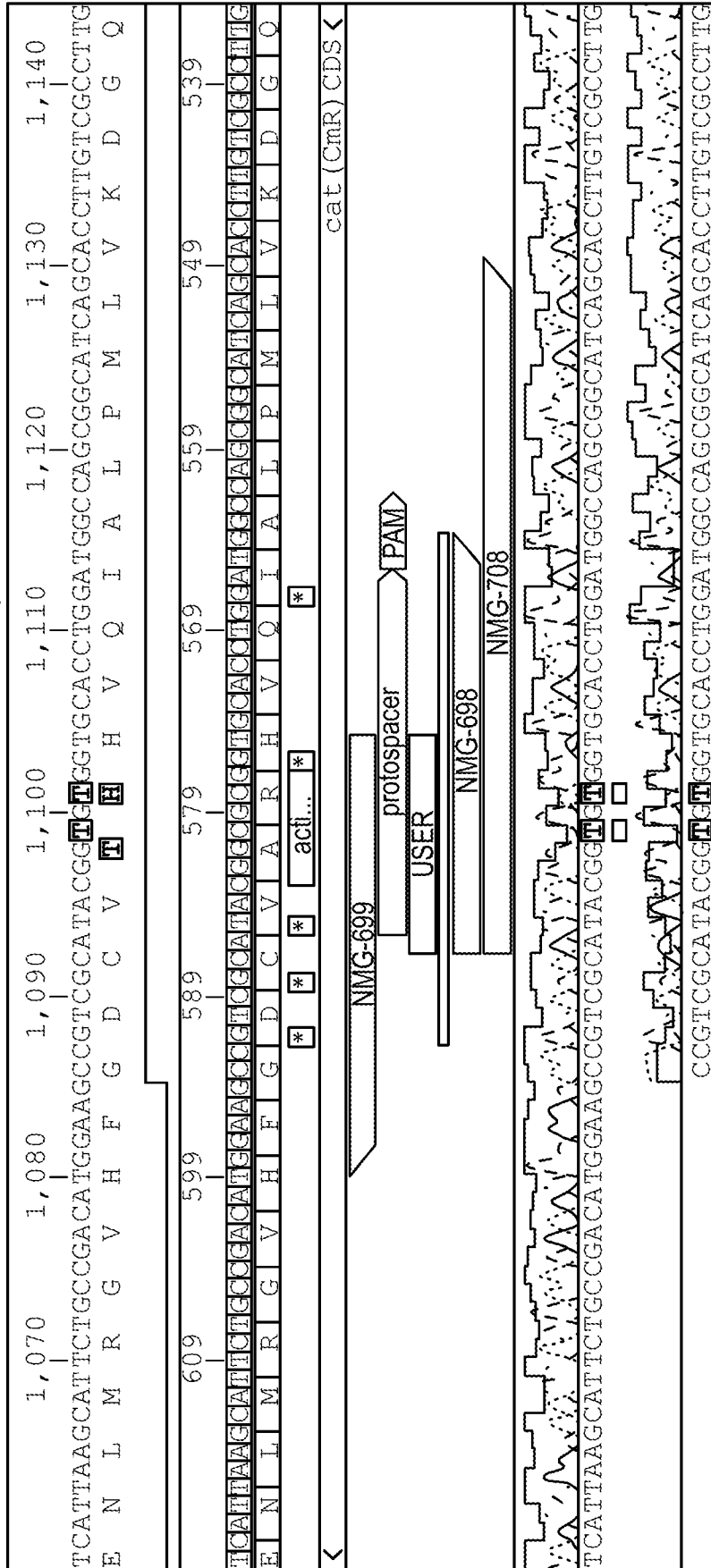


FIGURE 53A

CDA-XTEN-dCas9-UGI: survival on 8 ug/mL chloramphenicol

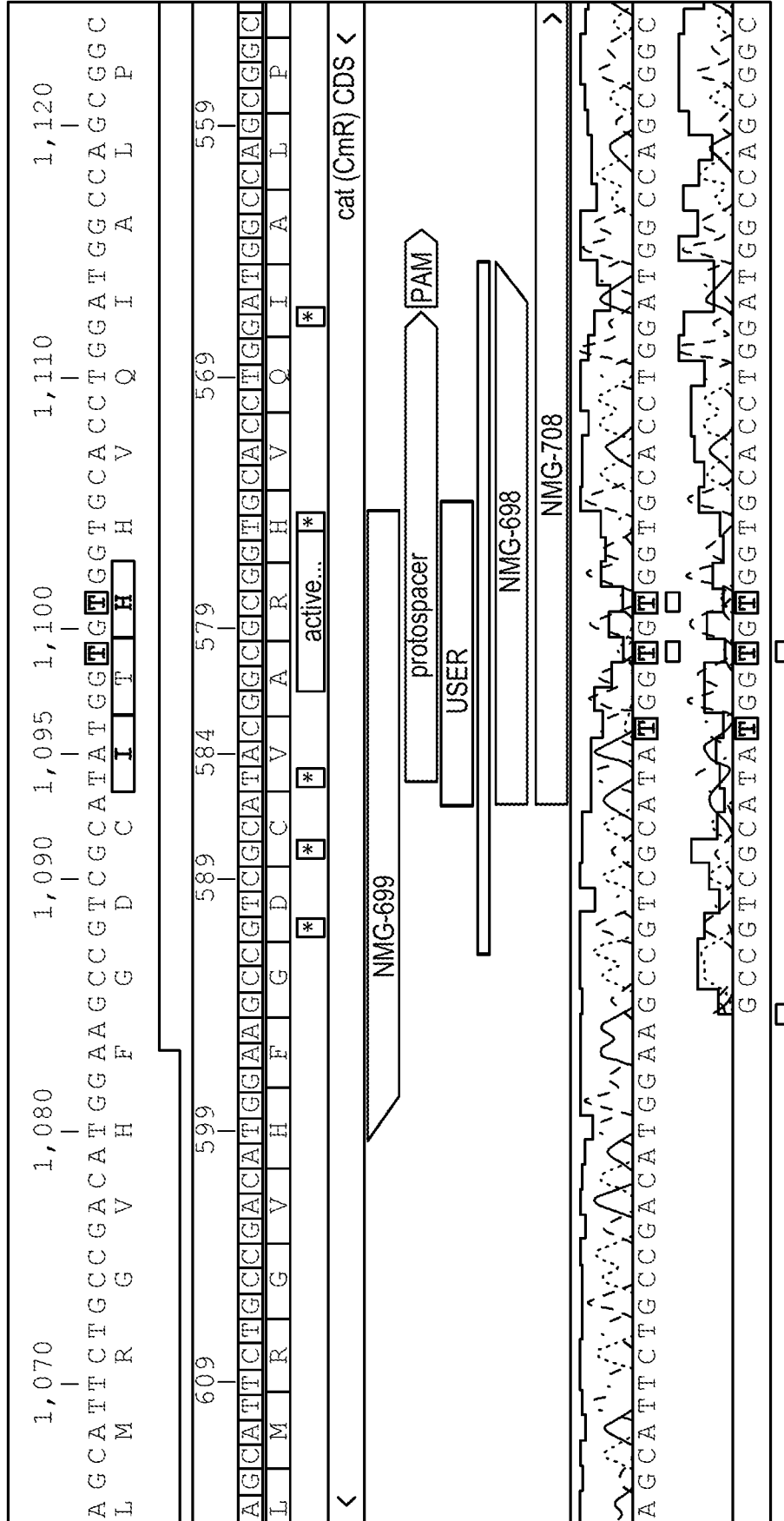


FIGURE 53B

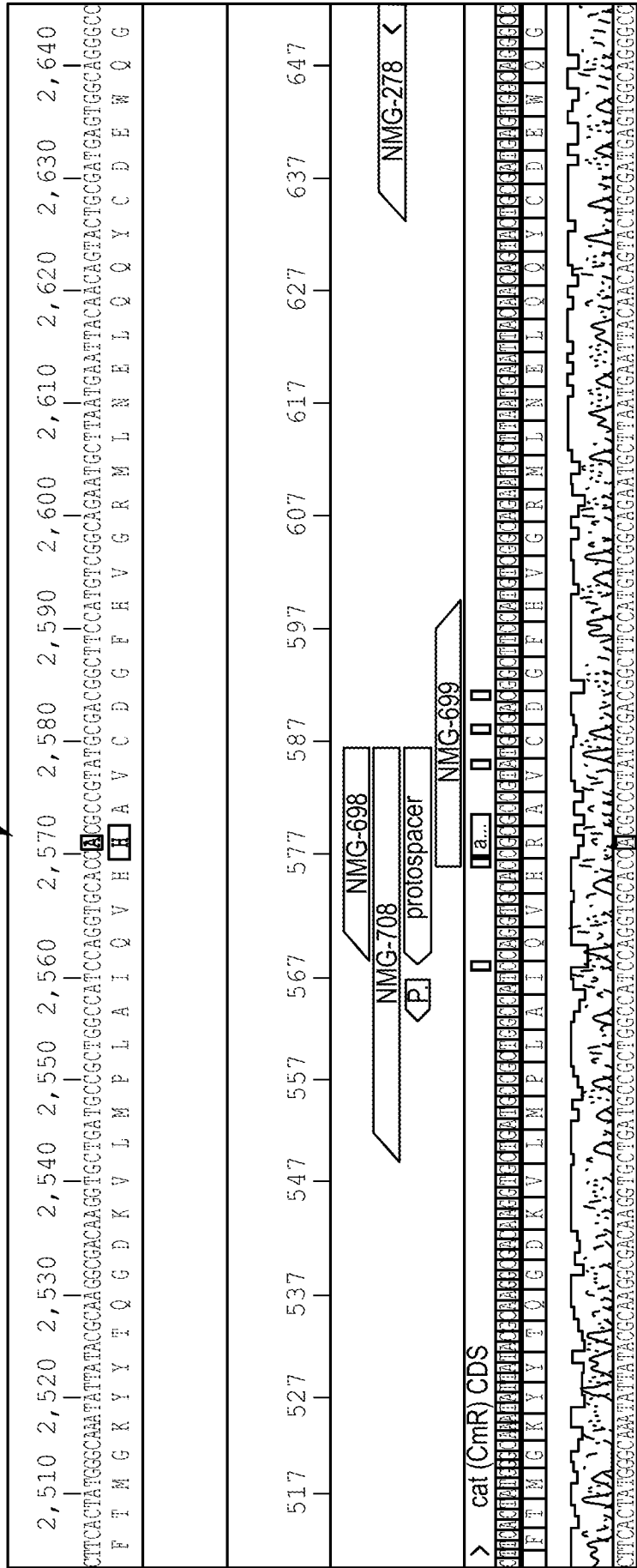
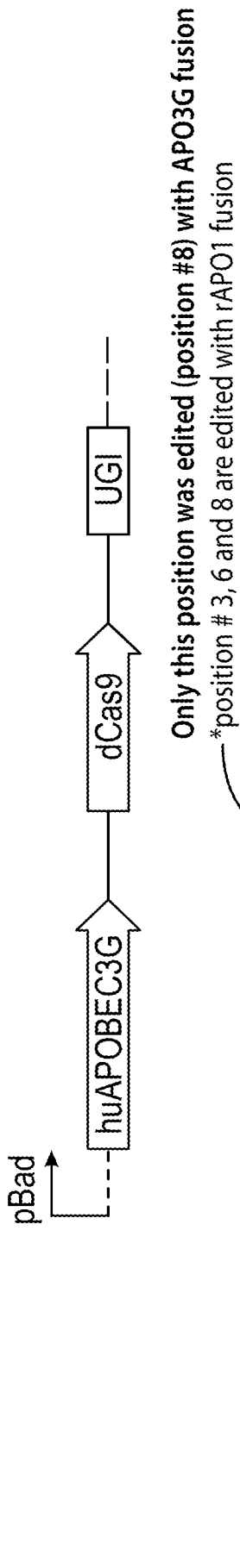


FIGURE 54

EMX1 on target	G	A	G	T	C	C	G	A	G	C	A	G	A	A	G	A	A	G	A	A	G	G	G	G	G	
untreated					0±0	0±0																				
BE3					42.8±3.1	41.6±5.5																				
HF-BE3					40.3±5.5	40.7±5.0																				
EMX1 off target 1	G	A	G	T	T	A	G	A	G	C	A	G	A	A	G	A	A	G	A	A	G	A	A	G	G	
untreated																										
BE3																										
HF-BE3																										
EMX1 off target 2	G	A	G	T	C	T	A	A	G	C	A	G	A	A	G	A	A	G	A	A	G	A	A	G	A	G
untreated					0±0																					
BE3					17.1±0.9																					
HF-BE3					1.7±0.3																					
EMX1 off target 3	G	A	G	G	C	C	G	A	G	C	A	G	A	A	G	A	A	G	A	A	G	A	C	G	G	
untreated					0±0	0±0																				
BE3					6.0±2	8.1±2.5																				
HF-BE3					0±0	0±0																				
EMX1 off target 4	G	A	G	T	C	C	T	A	G	C	A	G	A	A	G	A	A	G	A	A	G	A	A	G	A	G
untreated					0±0	0±0																				
BE3					1.3±0.1	1.3±0.1																				
HF-BE3					0.1±0	0.2±0																				

FIGURE 55

EMX1 off target 6	G	A	G	T	C	C	C	G	G	G	A	A	G	A	G	A	A	A	A	G	G
untreated					0±0	0±0	0±0														
BE3					0.4±0	0.4±0															
HF-BE3					0±0	0±0															
EMX1 off target 7	G	A	G	C	C	G	G	A	G	C	A	G	A	A	G	A	A	G	A	G	G
untreated				0±0	0±0	0±0															
BE3				0±0	0±0	0±0															
HF-BE3				0±0	0±0	0±0															
EMX1 off target 8	A	A	G	T	C	C	C	G	A	G	A	G	A	A	G	A	A	A	A	G	G
untreated					0±0	0±0	0±0														
BE3					0±0	0±0	0±0														
HF-BE3					0±0	0±0	0±0														
EMX1 off target 9	G	A	A	T	C	C	C	A	A	G	C	A	G	A	G	A	A	A	G	G	A
untreated					0±0	0±0	0±0														
BE3					0.1±0	0.1±0	0.1±0														
HF-BE3					0±0	0±0	0±0														
EMX1 off target 10	A	C	G	T	C	T	T	G	A	G	C	A	G	A	A	G	A	A	T	G	G
untreated		0±0			0±0																
BE3		0±0			15±0.1																
HF-BE3		0±0			11±0.2																

FIGURE 55 (CONTINUED)

EMX1		C ₃	C ₆	C ₁₂	indel %
untreated	A	0±0	0±0	0±0	0±0
	C	99.9±0	99.9±0	99.9±0	
	G	0±0	0±0	0±0	
	T	0±0	0±0	0±0	

BE3	A	1.3±0.1	0.7±0.2	0±0	2.6±0.3
	C	51.7±1	55.2±0.1	99.9±0	
	G	4±0.7	1.9±0.4	0±0	
	T	42.8±0.2	41.9±0.4	0±0	

HF BE3	A	1.5±0.7	0.5±0.1	0±0	1±0.1
	C	44.8±8.5	49.9±6.6	99.9±0	
	G	4.2±1.2	0.6±0.2	0±0	
	T	49.3±6.5	48.7±6.3	0±0	

RNF2		C ₃	C ₆	C ₁₂	indel %
untreated	A	0±0	0±0	0±0	0±0
	C	99.9±0	99.9±0	99.9±0	
	G	0±0	0±0	0±0	
	T	0±0	0±0	0±0	

BE3	A	0.4±0	1.5±0.2	0.1±0	2.3±0.3
	C	70.9±2.7	45±0	95±0.2	
	G	0±0	12.6±2.4	0.1±0	
	T	28.5±2.7	40.7±2.7	4.6±0.2	

HF BE3	A	0±0	0.2±0	0±0	0.5±0.1
	C	76.8±1.8	64.5±3.3	95.4±0.1	
	G	0±0	2.4±0.5	0±0	
	T	23±1.8	32.7±2.6	4.5±0.1	

FANCF		C ₅	C ₇	C ₈	C ₁₁	indel %
untreated	A	0±0	0±0	0±0	0±0	0±0
	C	99.9±0	99.9±0	99.9±0	99.9±0	
	G	0±0	0±0	0±0	0±0	
	T	0±0	0±0	0±0	0±0	

BE3	A	0.8±0.1	1.2±0.3	1.2±0.1	0.2±0.1	5.8±0.9
	C	64.7±5.9	67.9±5.1	68.9±5.2	85.4±2.7	
	G	0.6±0	0.7±0	0.7±0.1	0.1±0	
	T	33.8±5.7	29.9±4.8	29±5	14.1±2.6	

HF BE3	A	0.9±0.2	2±0.4	0.9±0.2	0.4±0	5.9±0.7
	C	52.5±8.4	57.7±7.6	61.9±6.8	84.7±2.4	
	G	1.4±0.2	0.6±0	0.5±0.1	0.3±0	
	T	45±8	39.5±7.2	36.5±6.5	14.4±2.2	

HEK3		C ₅	C ₇	C ₈	C ₁₁	indel %
untreated	A	0±0	0±0	0±0	0±0	0±0
	C	99.9±0	99.9±0	99.9±0	99.9±0	
	G	0±0	0±0	0±0	0±0	
	T	0±0	0±0	0±0	0±0	

BE3	A	0±0	1.6±0.2	1.5±0.3	0.2±0	2.7±0.4
	C	98.7±0.2	48.6±5.4	40.8±6.7	98.6±0.1	
	G	0±0	1.4±0.2	10.5±1.2	0.3±0	
	T	1.1±0.2	48.2±4.9	47±5.2	0.7±0.1	

HF BE3	A	0±0	0.8±0.2	1.1±0.3	0.4±0	4.1±0.7
	C	97.3±0.4	65±3.8	40.4±6.3	95.9±0.5	
	G	0±0	0.9±0.2	6.1±1.1	0.7±0.2	
	T	2±0.4	33±3.4	52.2±5.2	2.7±0.2	

HEK4		C ₅	C ₇	C ₈	C ₁₁	indel %
untreated	A	0±0	0±0	0±0	0±0	0±0
	C	99.9±0	99.9±0	99.9±0	99.9±0	
	G	0±0	0±0	0±0	0±0	
	T	0±0	0±0	0±0	0±0	

BE3	A	0±0	5.4±1.3	0±0	0±0	3±0.7
	C	98.7±0	44.5±8.2	98.9±0	99.8±0	
	G	0±0	18.1±2.3	0±0	0±0	
	T	1.1±0.1	31.9±4.8	0.9±0.1	0.1±0	

HF BE3	A	0±0	5.2±0.8	0±0	0±0	1.3±0
	C	99.6±0	38.6±9	98.5±0.1	99.9±0	
	G	0±0	16±2.8	0±0	0±0	
	T	0.2±0	42.1±5.3	1.3±0.1	0±0	

FIGURE 56

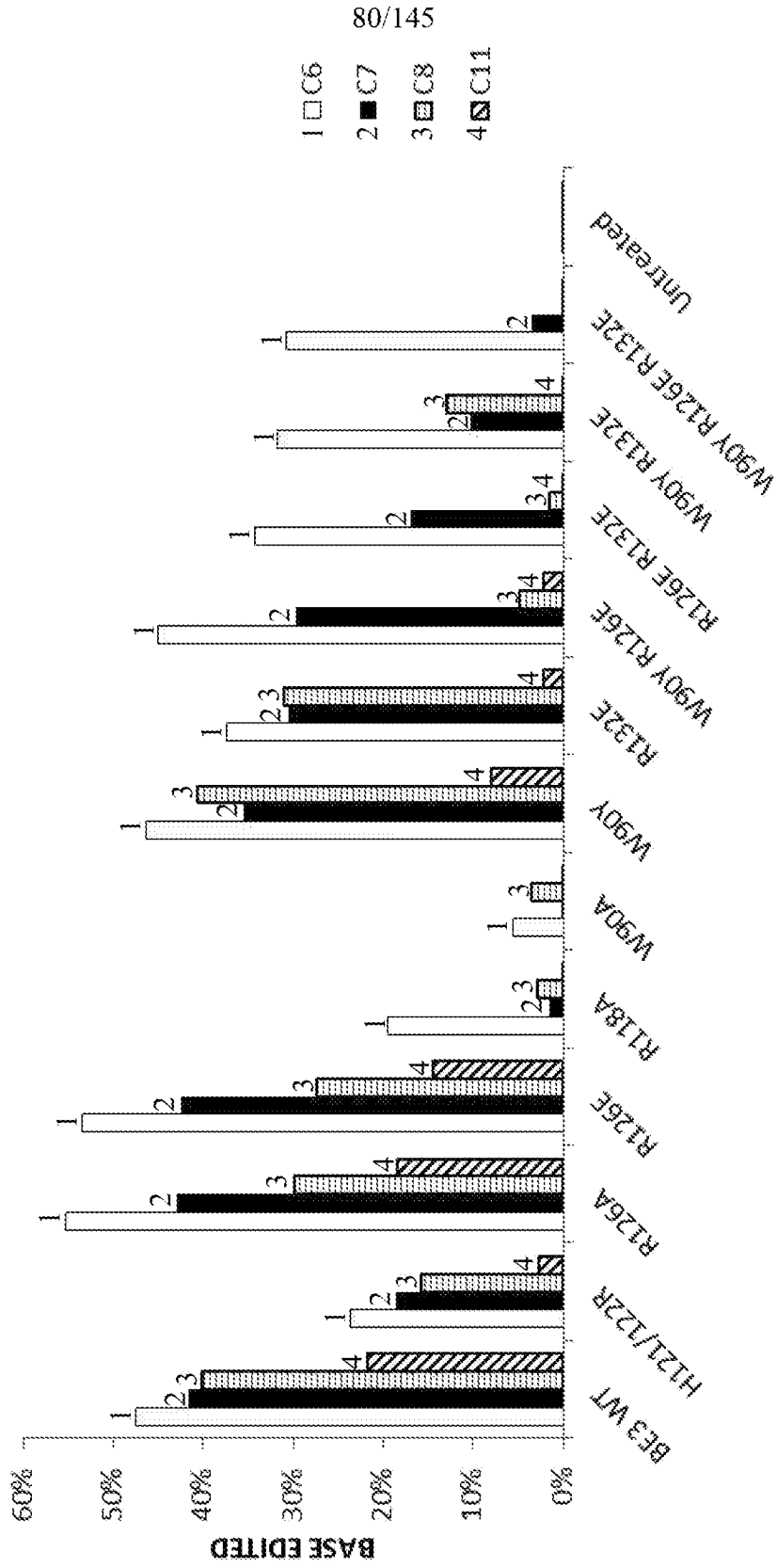


FIGURE 57

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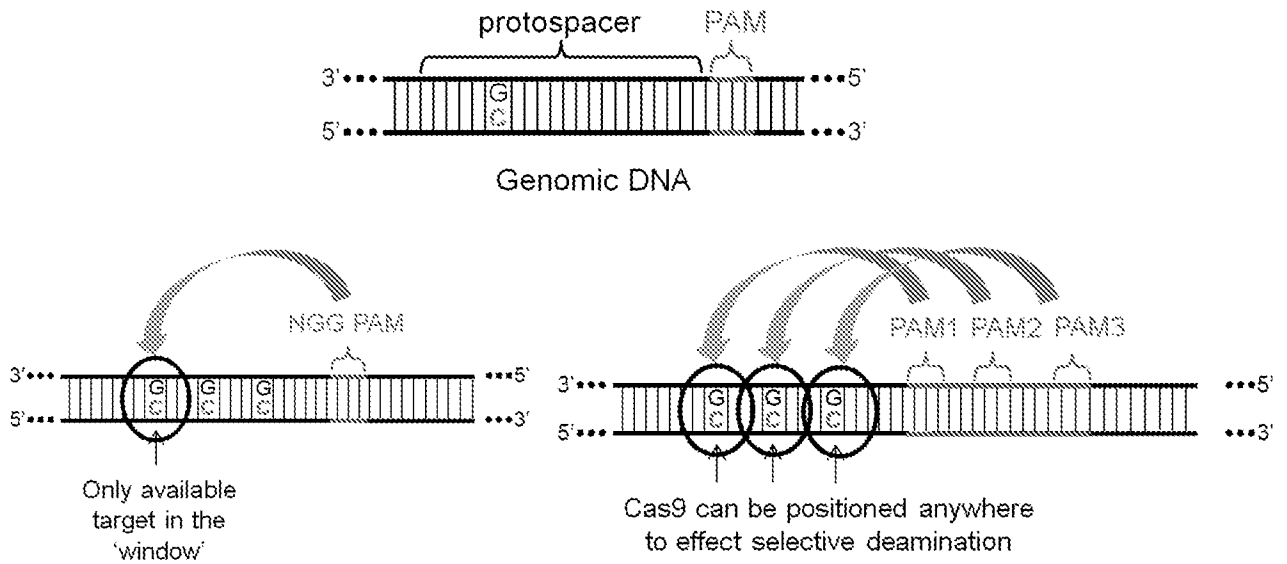


FIGURE 58

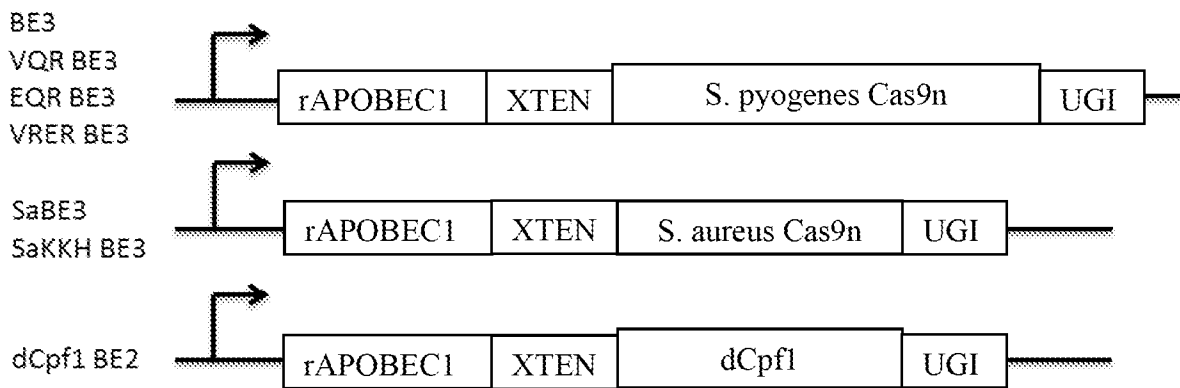


FIGURE 59

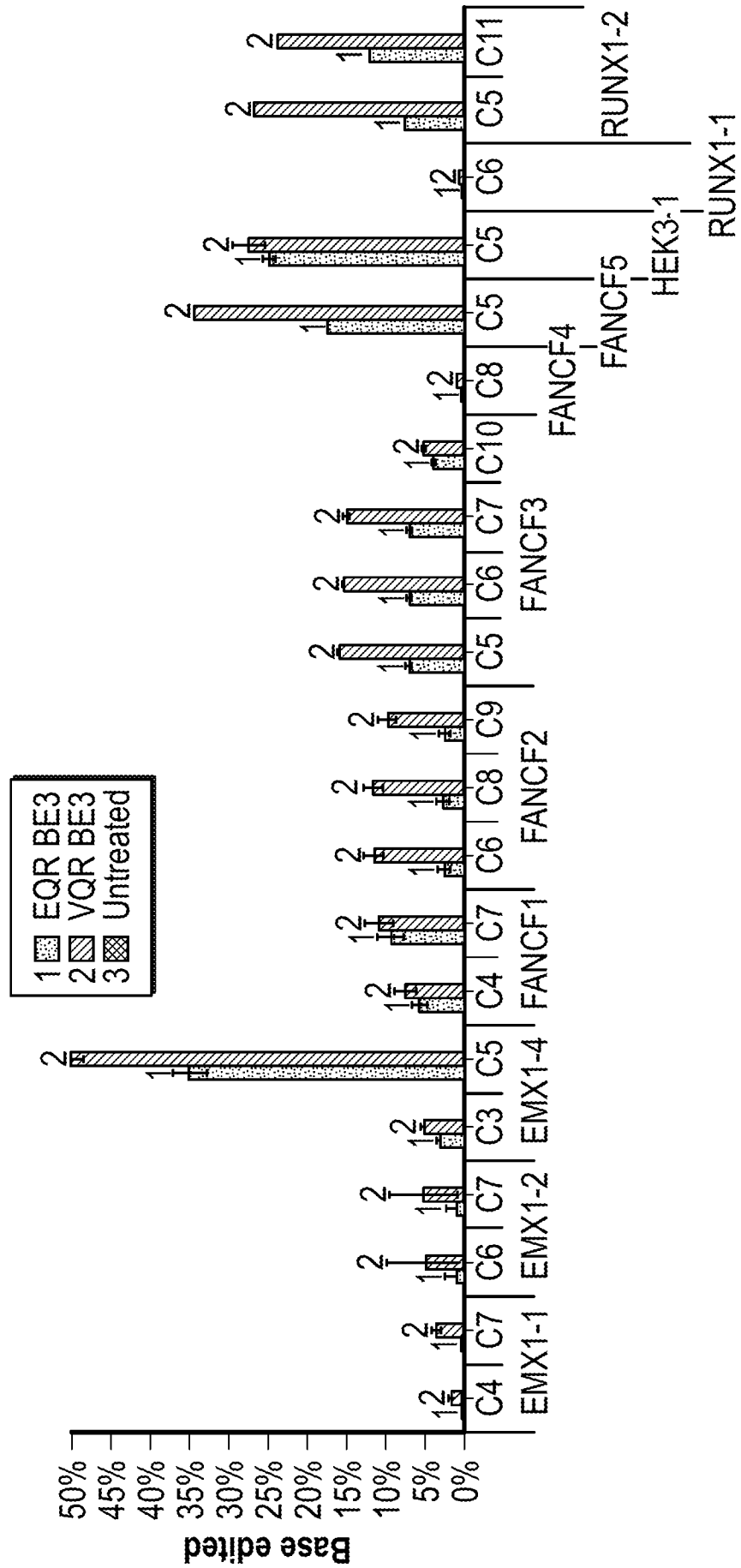


FIGURE 60

NGCG PAM EMX (VRER BE3)

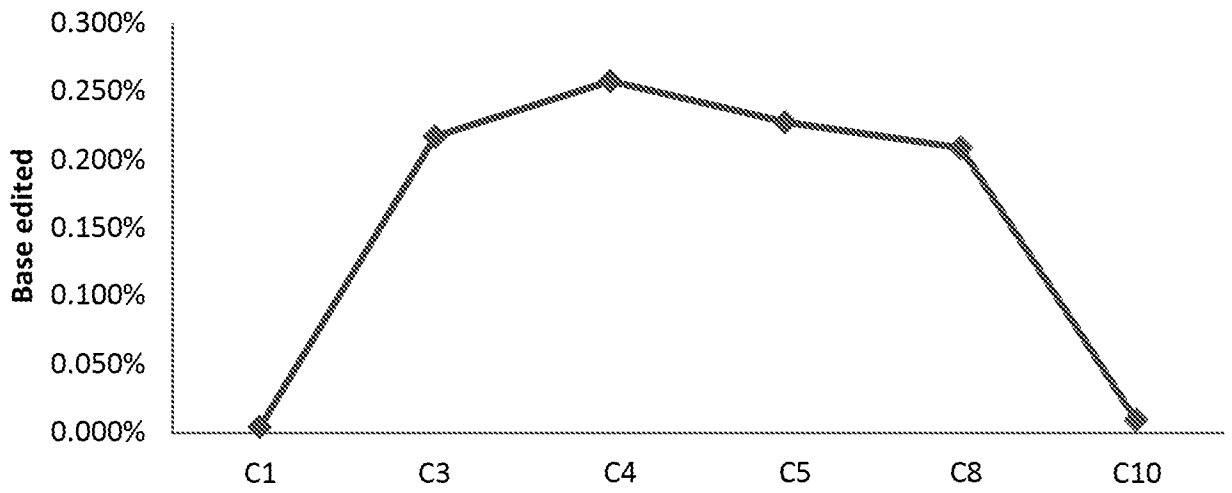


FIGURE 61

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□ SaKKH BE3
■ SaBE3
▨ Untreated

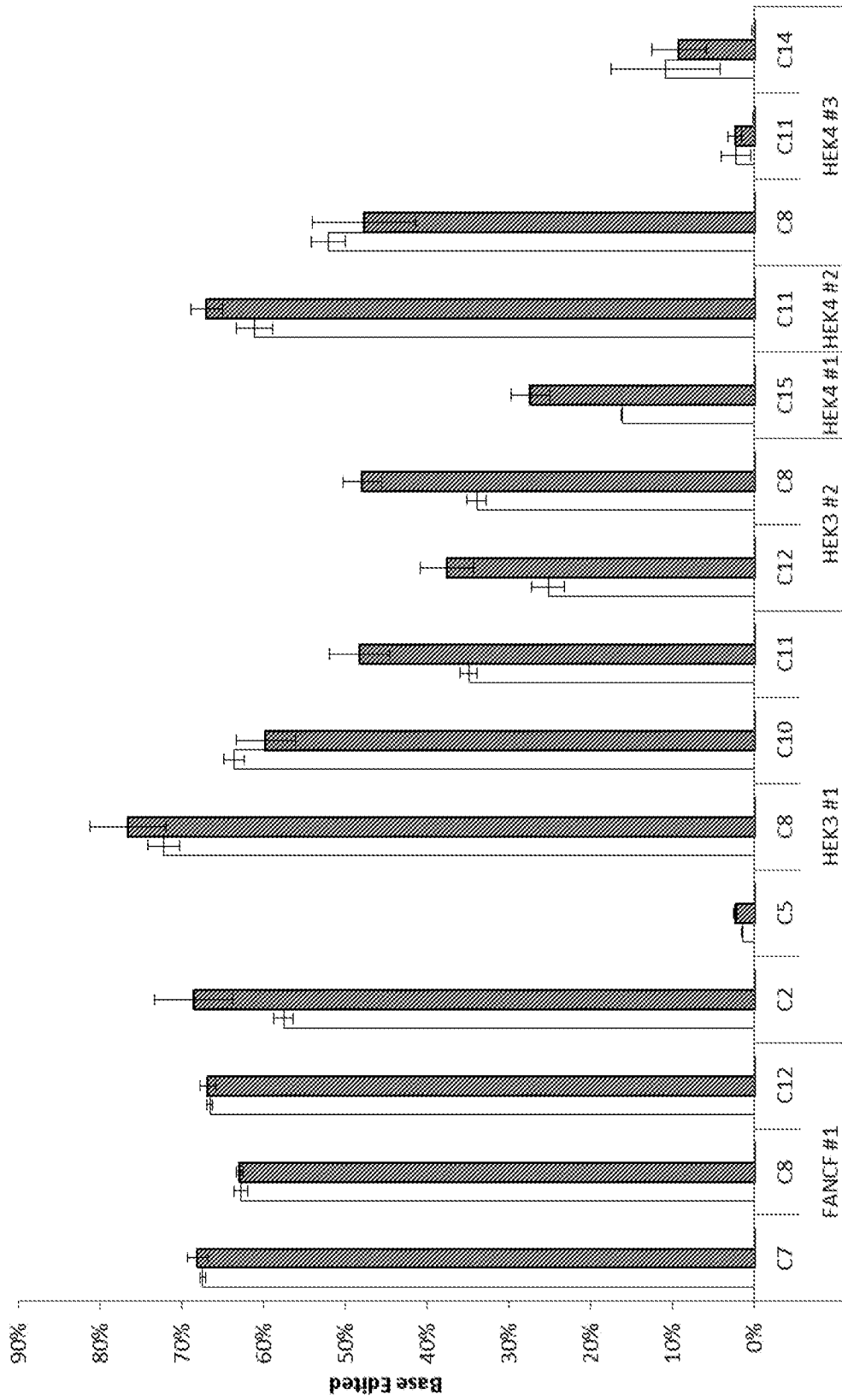


FIGURE 62

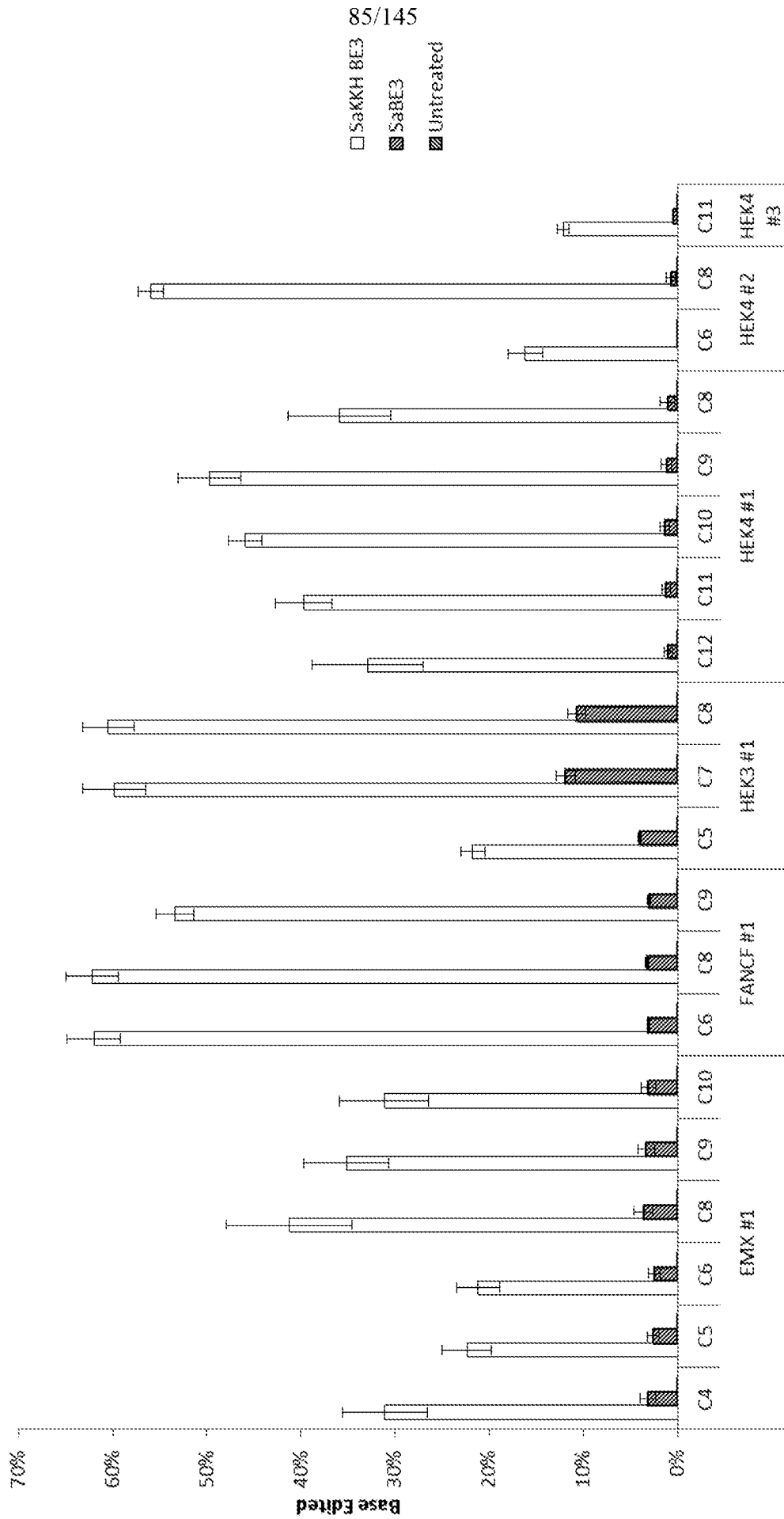


FIGURE 63

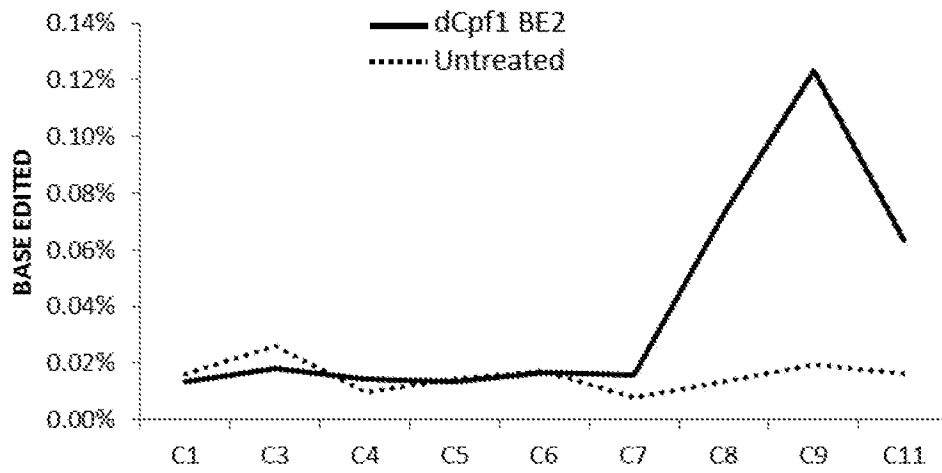


FIGURE 64A

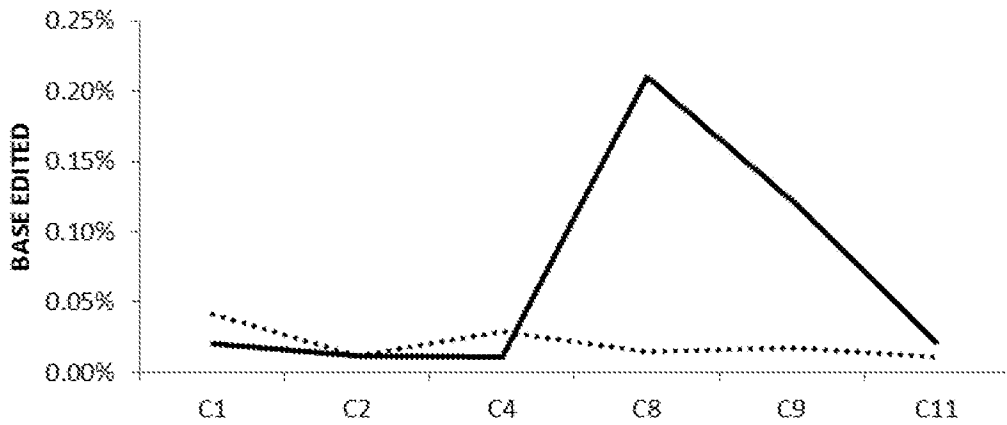


FIGURE 64B

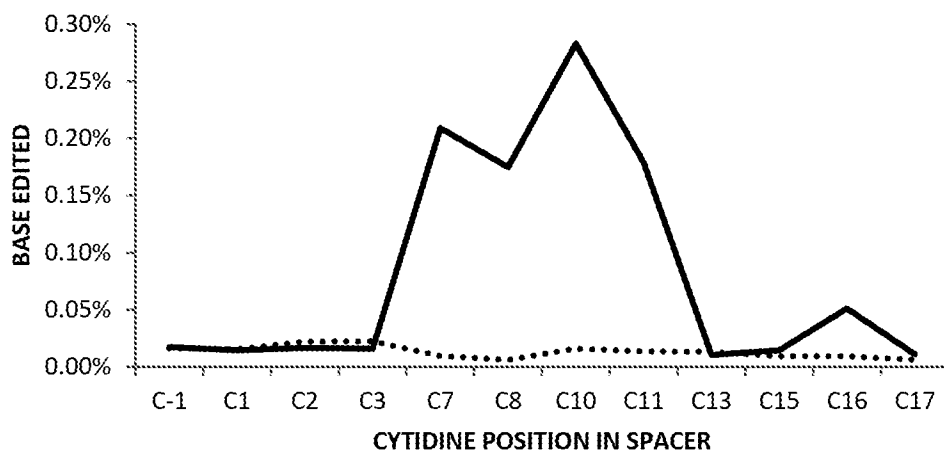


FIGURE 64C

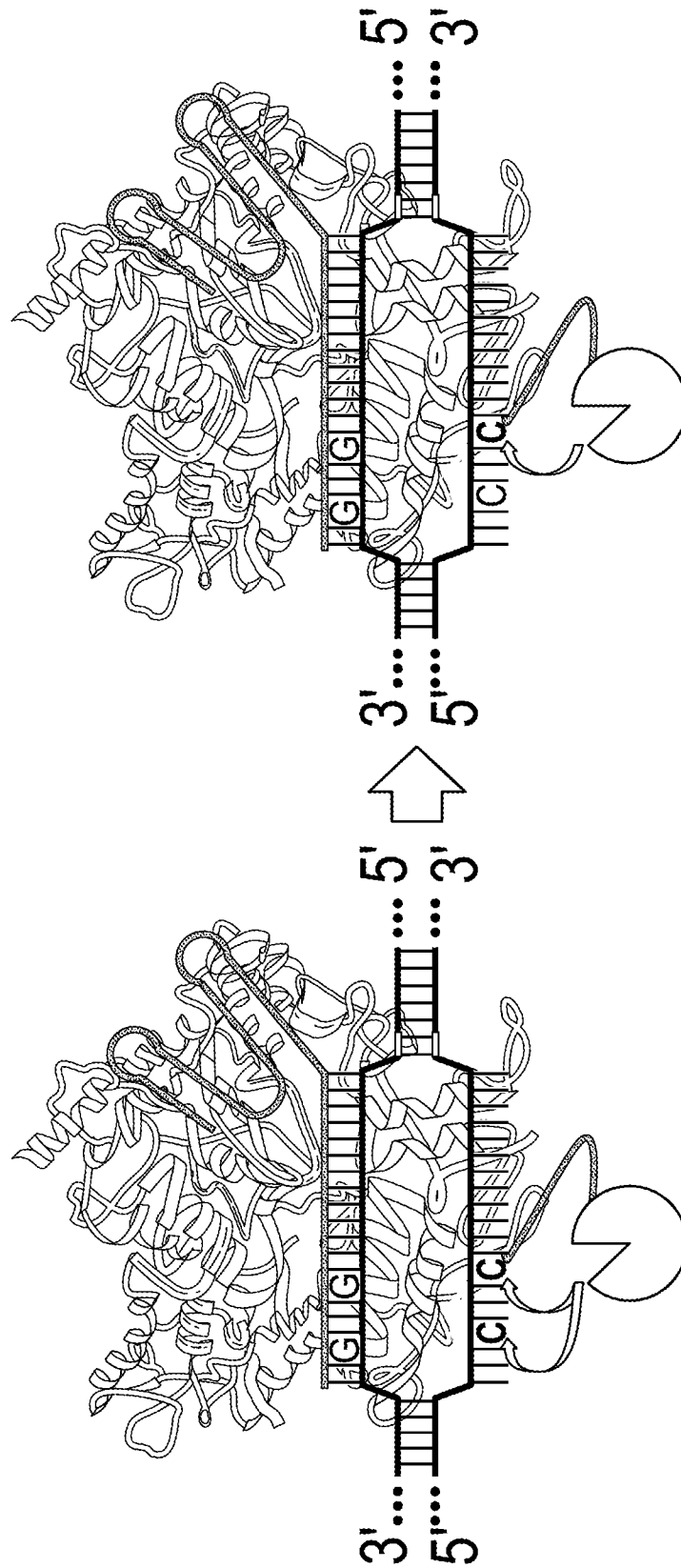


FIGURE 65

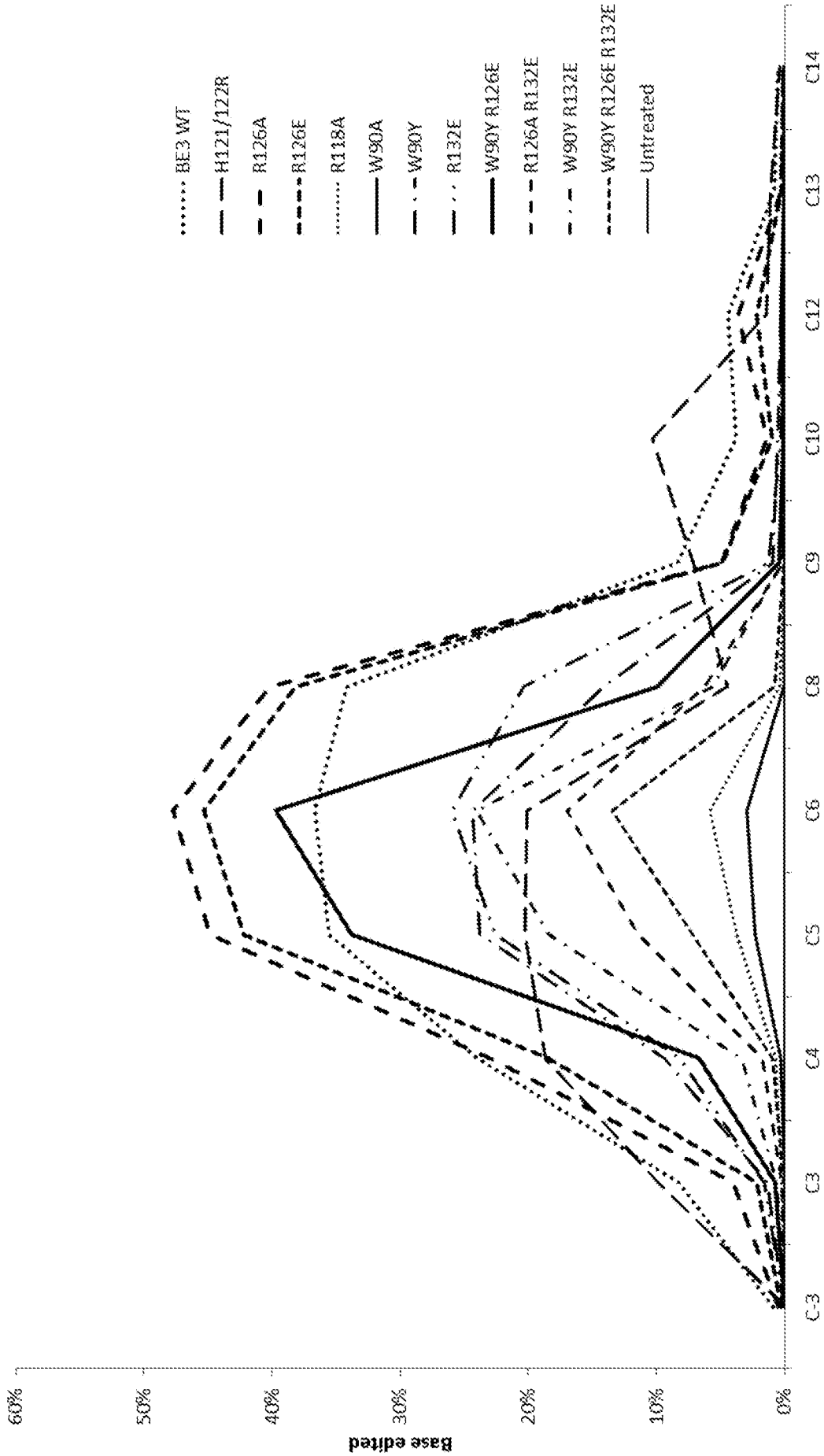


FIGURE 66

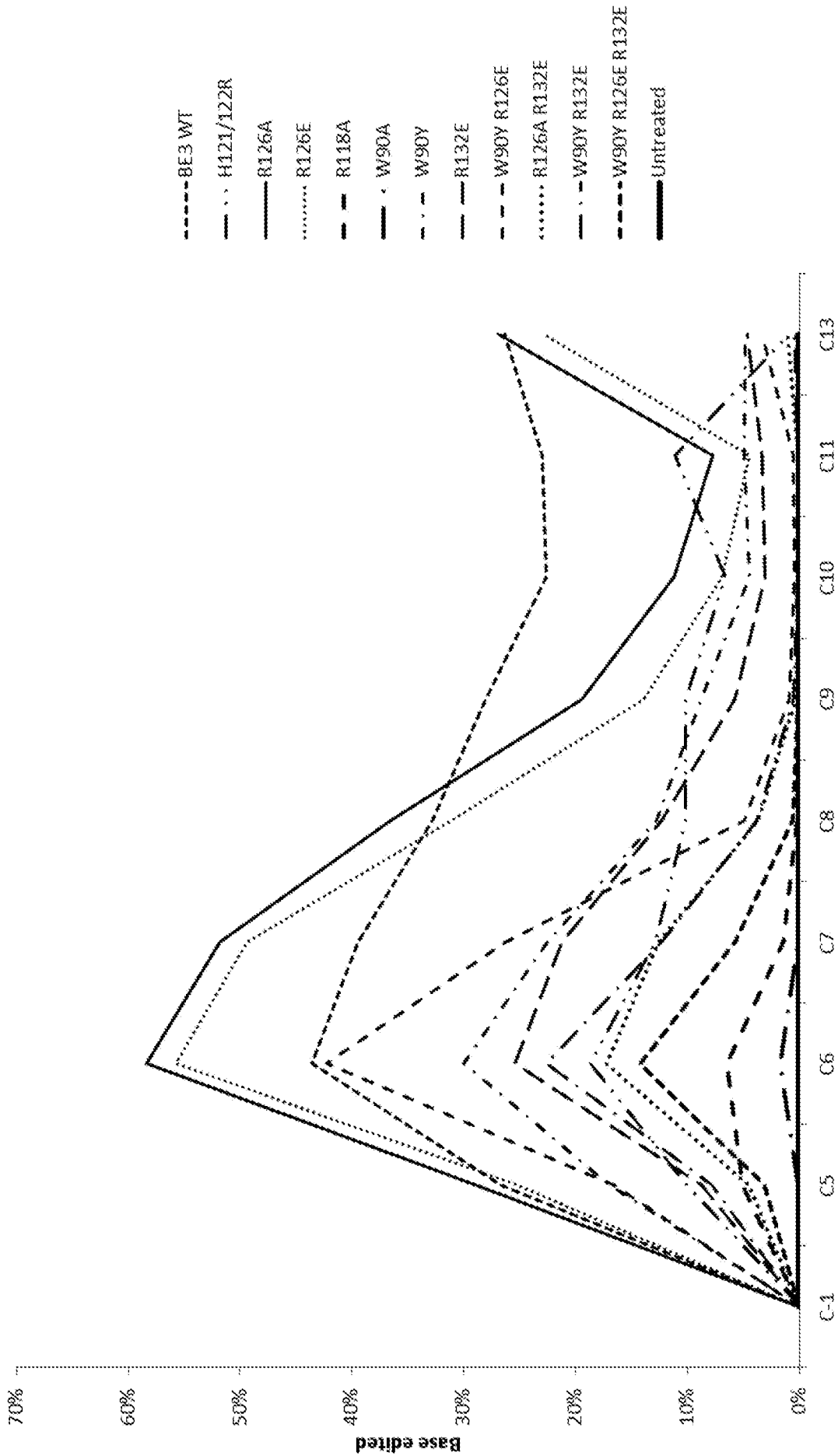


FIGURE 67

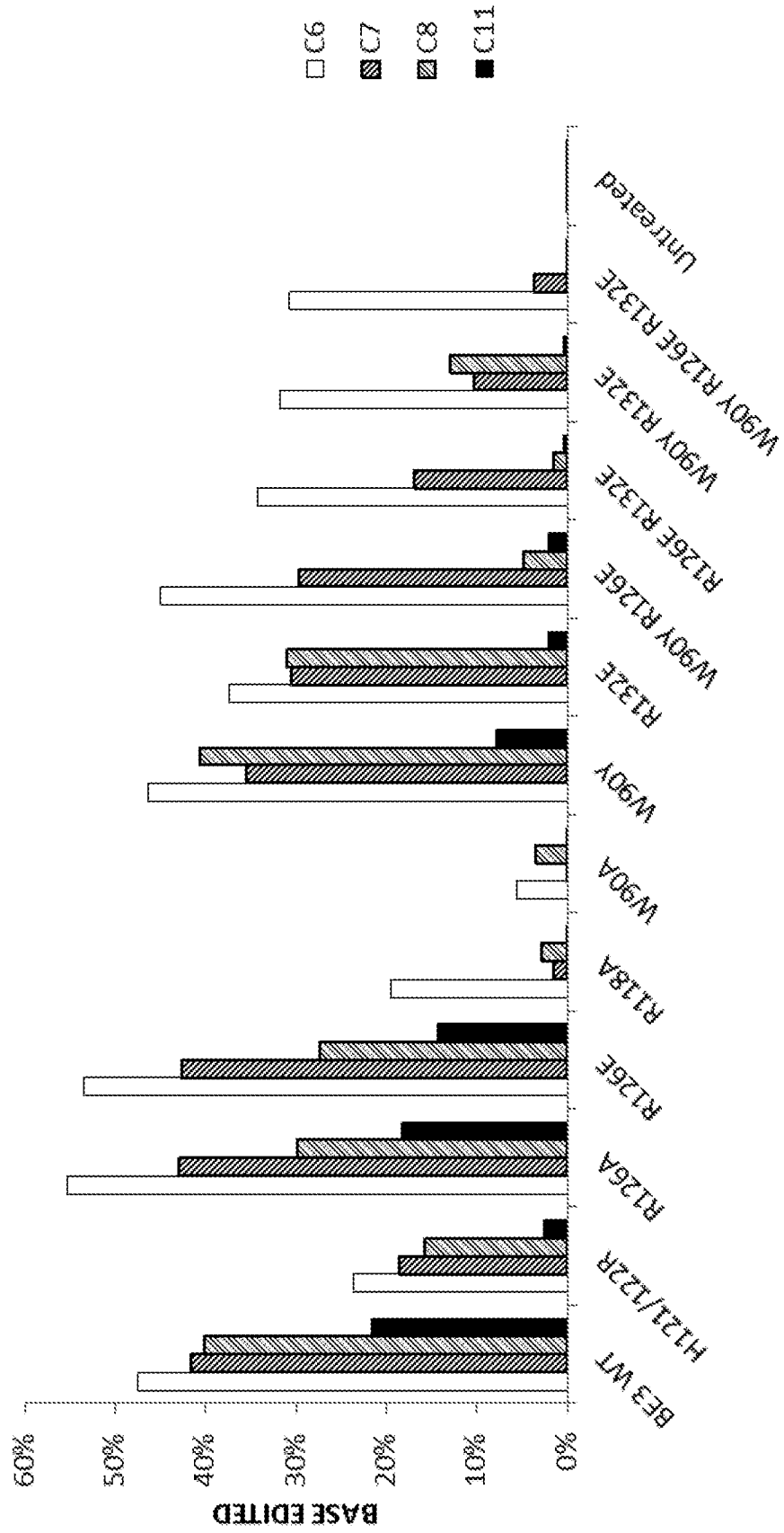


FIGURE 68

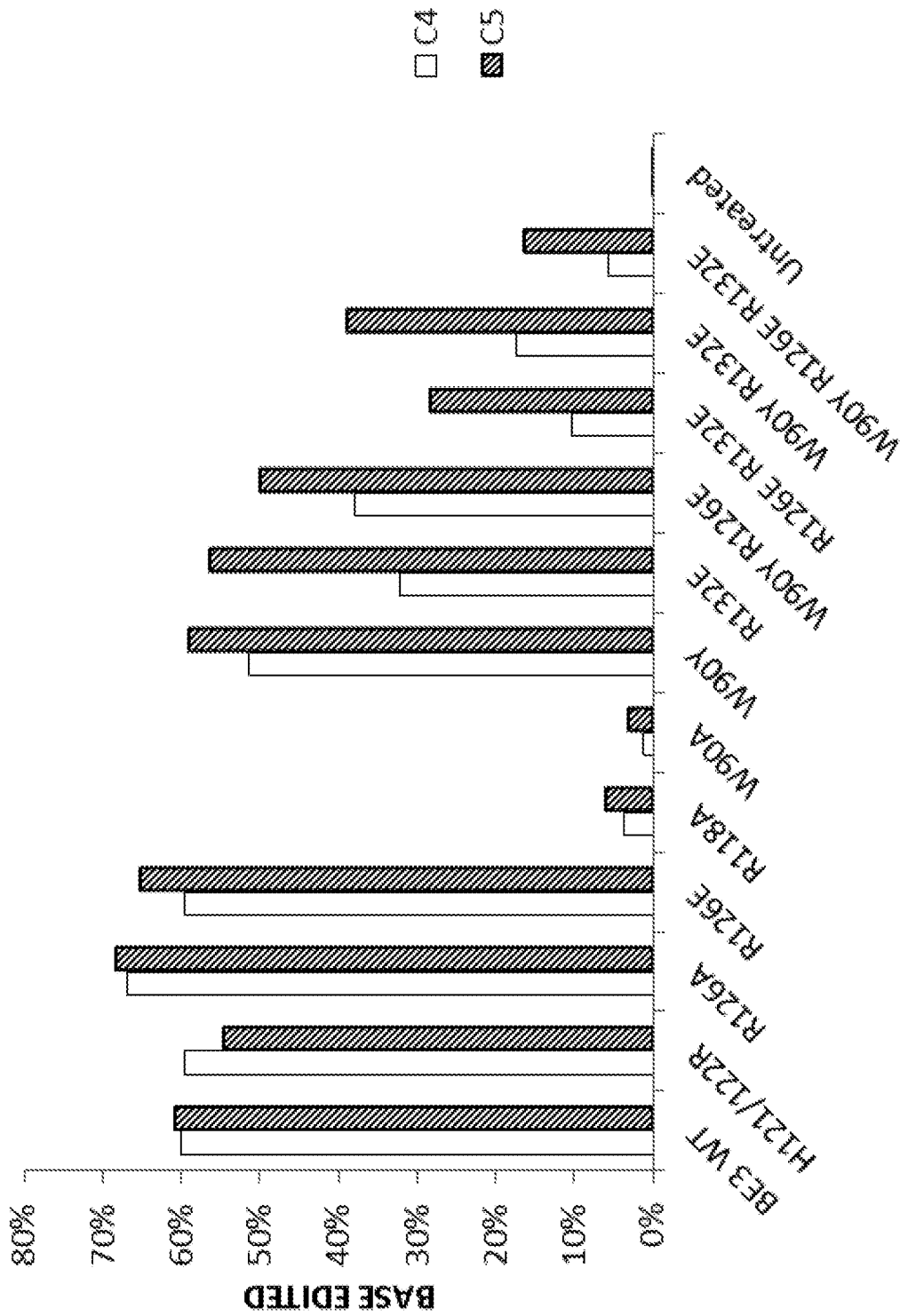


FIGURE 69

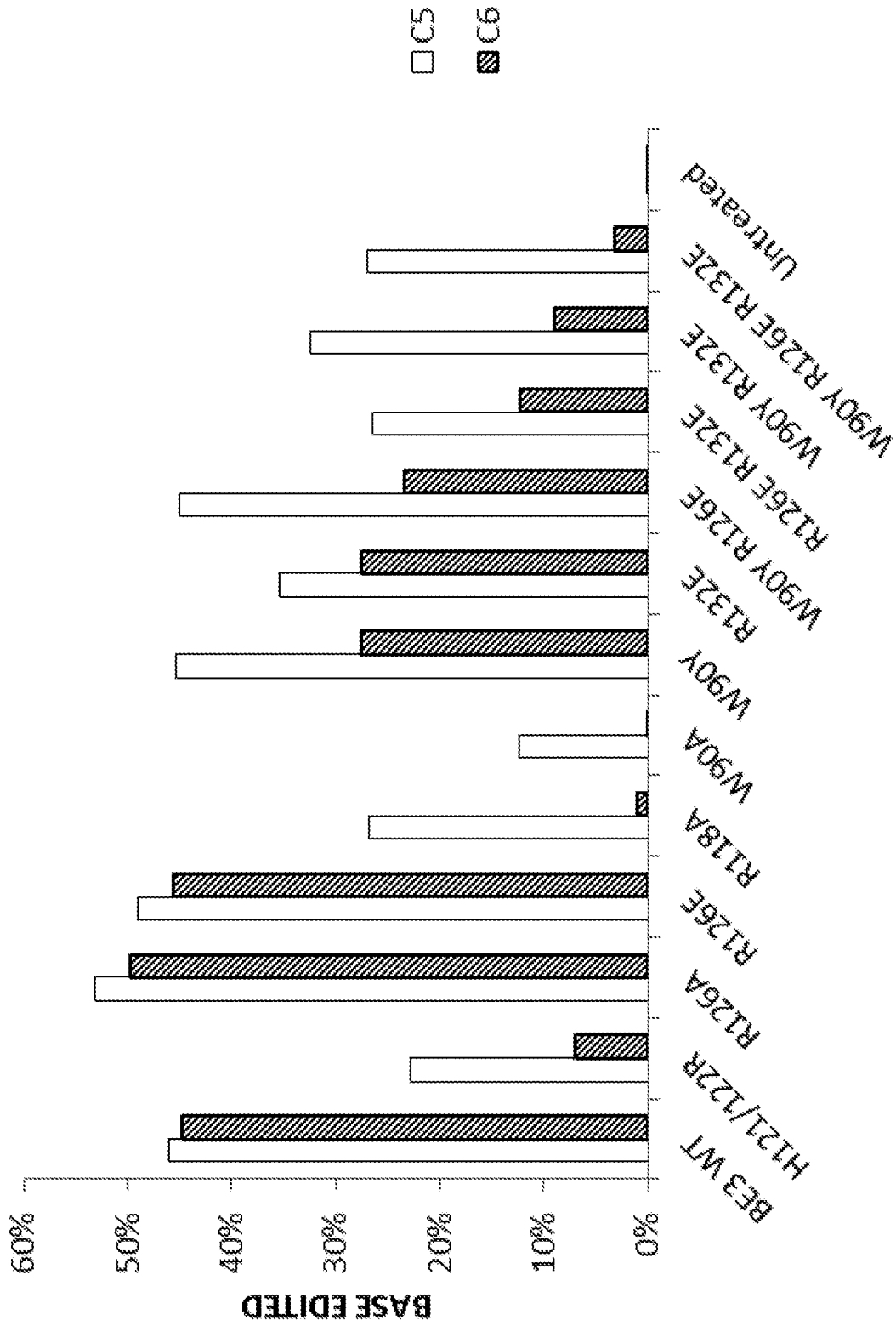


FIGURE 70

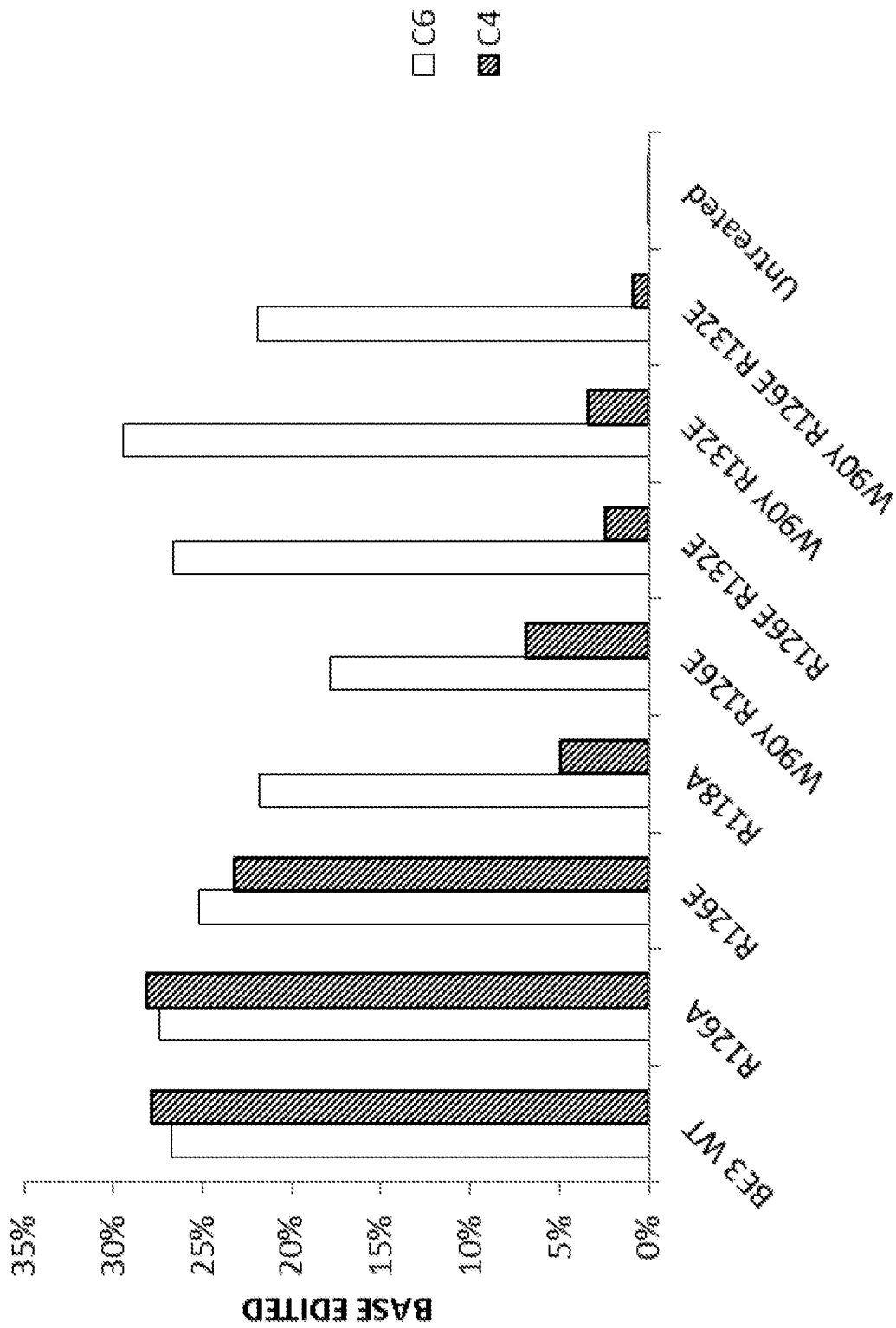


FIGURE 71

BE3

Base	Lys			Arg → Cys			Leu → Leu			Ala			Tyr			Gln				
	A	A	G	C	G	C	C	T	G	C	A	A	G	T	G	T	A	C		
	G	A	G	C	G	C	C	T	G	C	A	A	G	T	G	T	A	C		
A	0.2	99.9	99.8	0.0	1.0	0.1	1.8	1.2	0.0	0.0	0.1	0.1	99.8	0.0	0.0	0.1	0.1	99.9	0.1	0.0
C	0.0	0.0	0.1	0.0	38.1	0.0	49.8	52.3	0.0	0.0	0.0	99.7	0.1	0.0	0.1	0.0	0.0	0.0	99.9	99.9
G	99.8	0.0	0.0	99.9	1.8	99.8	1.3	0.7	0.1	99.9	99.8	0.0	0.1	99.9	0.1	99.9	0.1	0.0	0.0	0.0
T	0.0	0.0	0.1	0.1	59.2	0.1	47.0	45.8	99.9	0.1	0.1	0.2	0.0	0.0	99.8	0.0	99.8	0.0	0.1	0.1

BE3 W90Y R132E

Base	Lys			Arg → Cys			Leu → Leu			Ala			Tyr			Gln				
	A	A	G	C	G	C	C	T	G	C	A	A	G	T	G	T	A	C		
	G	A	G	C	G	C	C	T	G	C	A	A	G	T	G	T	A	C		
A	0.0	99.9	99.9	0.0	0.5	0.1	0.3	0.1	0.0	0.0	0.0	0.1	99.8	0.0	0.0	0.0	0.1	99.9	0.1	0.0
C	0.0	0.0	0.0	0.1	78.0	0.0	94.8	98.9	0.0	0.0	0.0	99.8	0.1	0.0	0.0	0.0	0.0	0.0	99.9	99.9
G	100.0	0.0	0.0	99.9	0.5	99.9	0.1	0.1	0.0	99.9	99.9	0.0	0.1	100.0	0.1	99.9	0.0	0.0	0.0	0.0
T	0.0	0.0	0.0	0.0	21.0	0.0	4.8	0.9	99.9	0.0	0.0	0.1	0.0	0.0	99.8	0.0	99.9	0.0	0.0	0.0

FIGURE 72

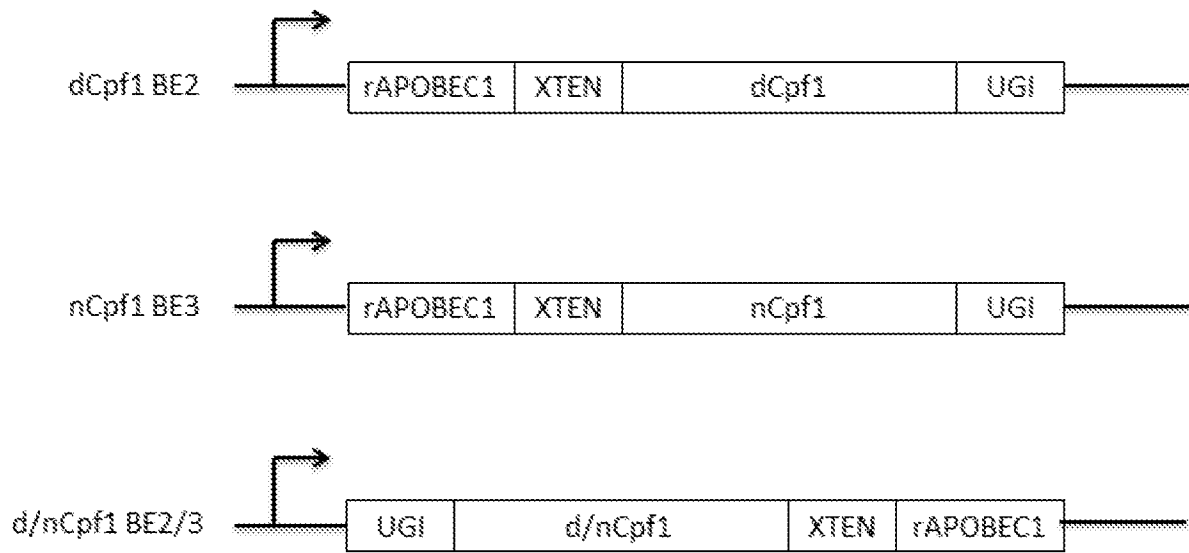
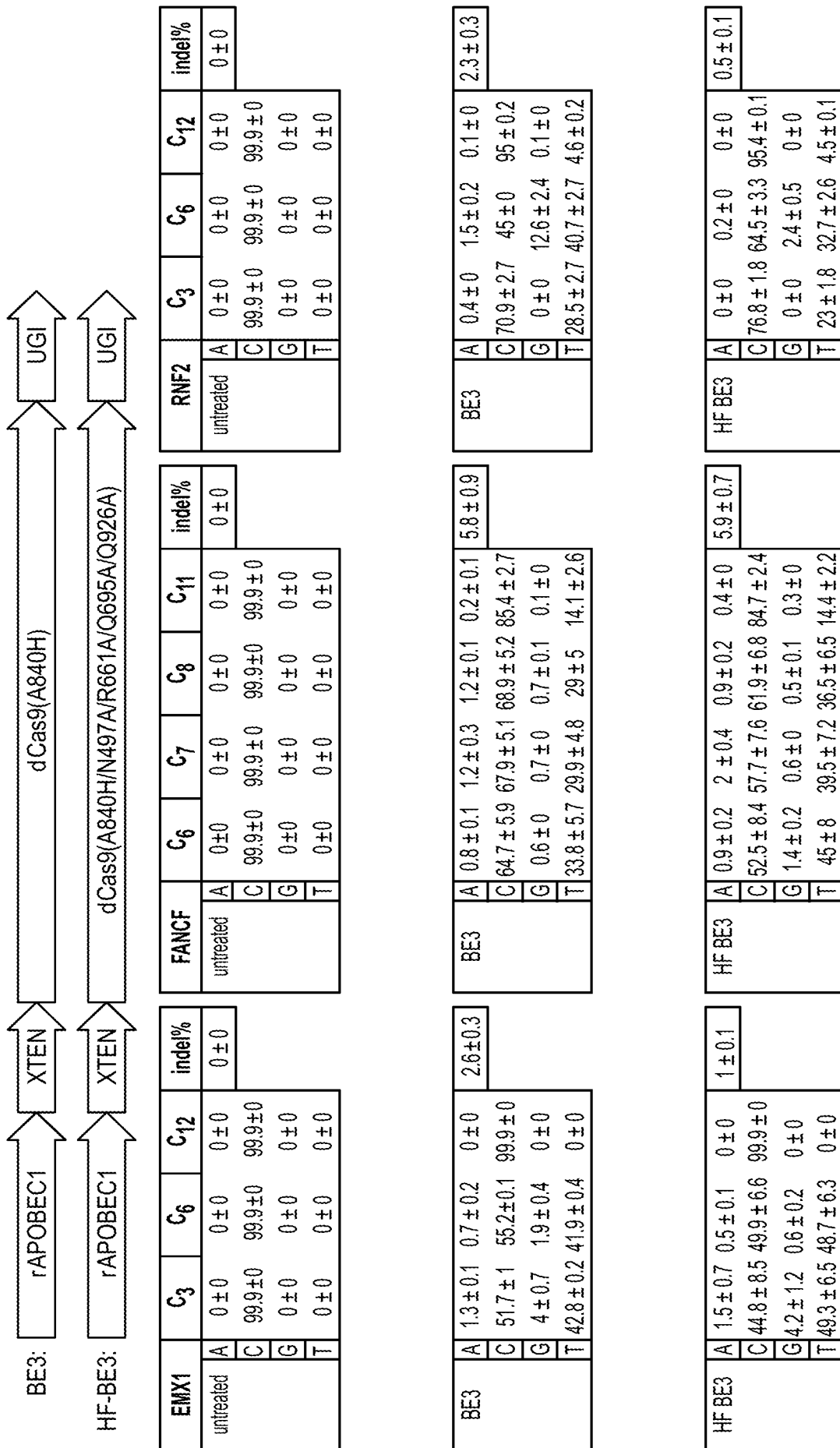
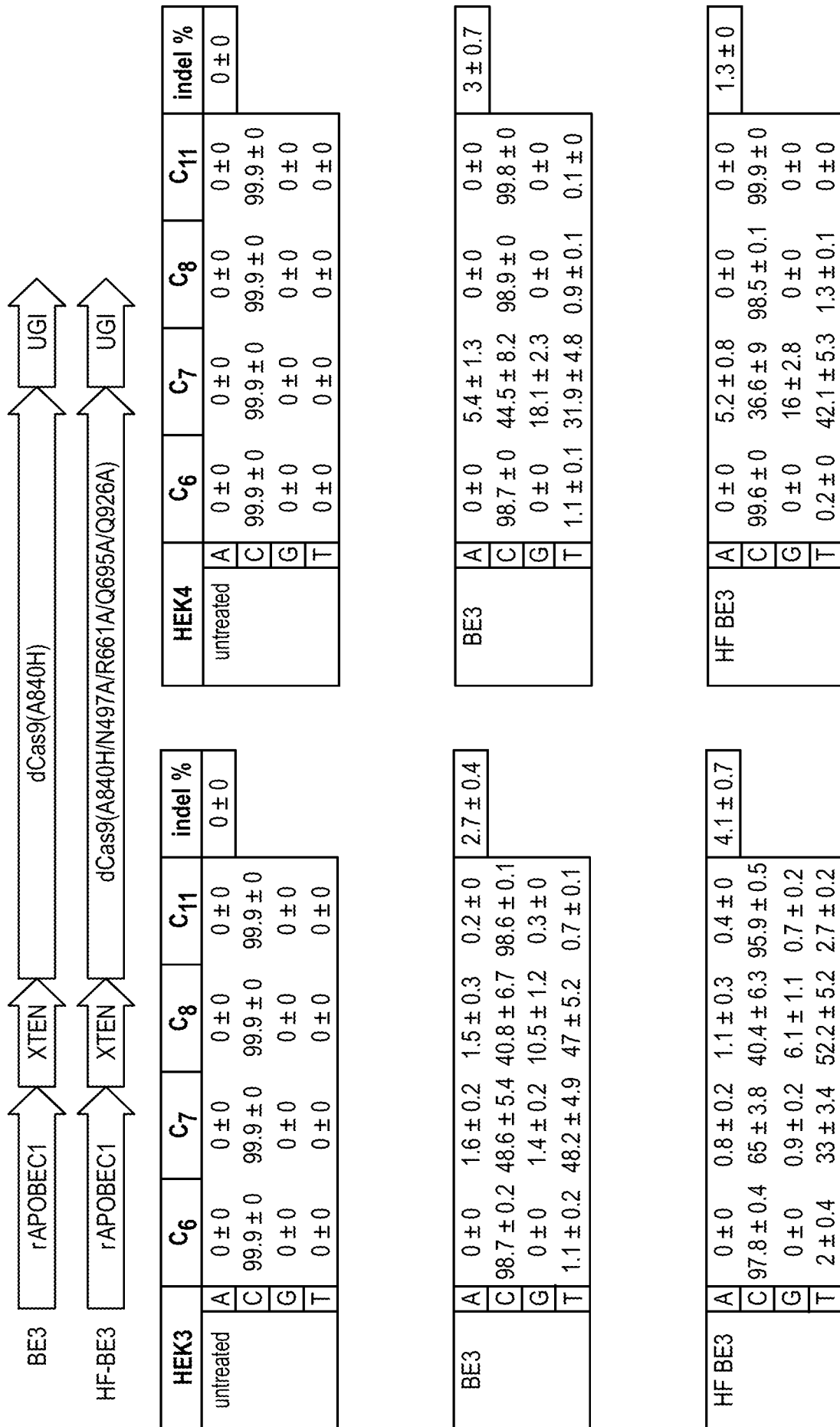


FIGURE 73



numbers are $\mu \pm \sigma$ from three independent replicates

FIGURE 74



numbers are $\mu \pm \sigma$ from three independent replicates

FIGURE 75

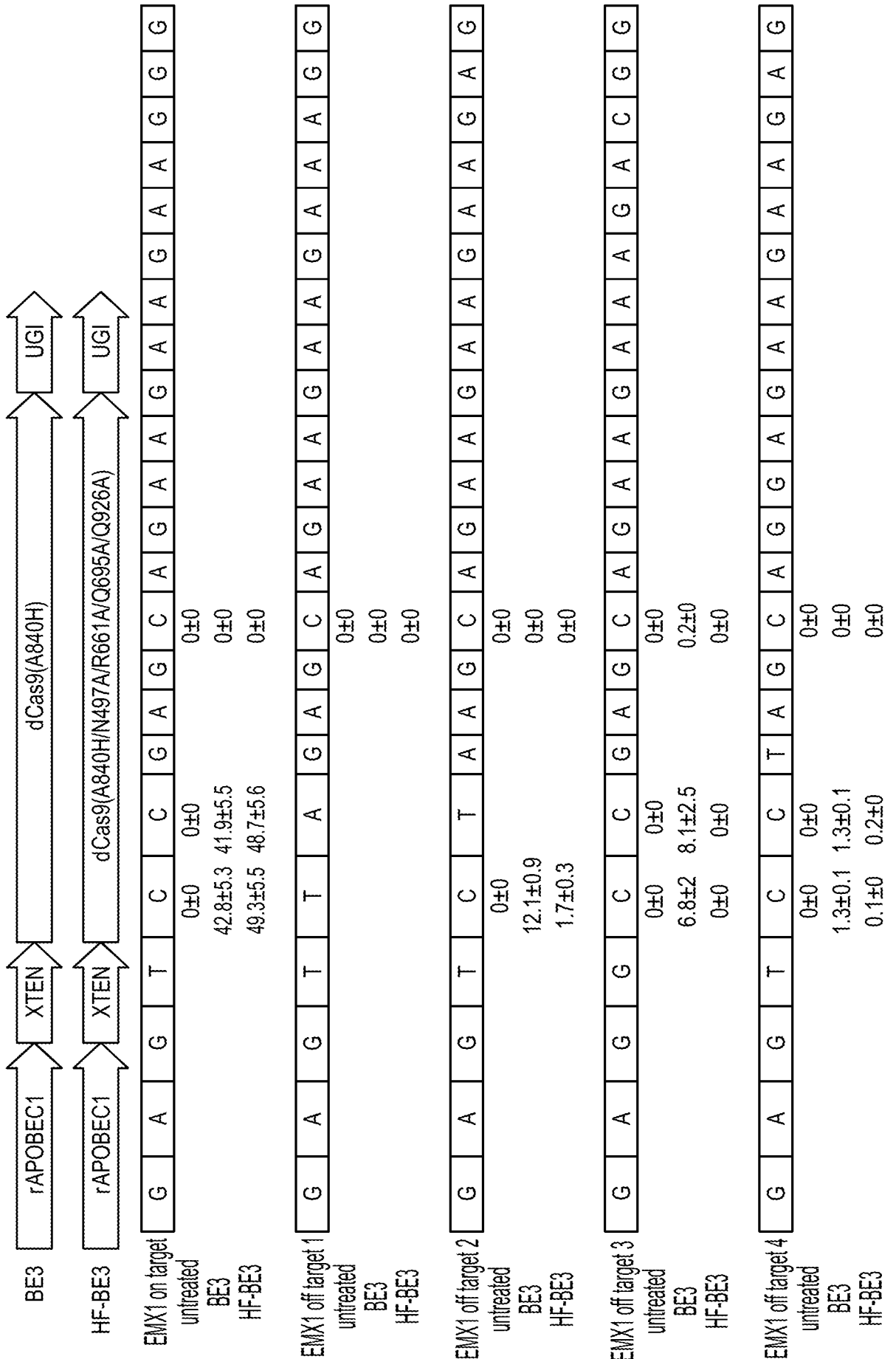


FIGURE 76

EMX1 off target 6	G	A	G	T	C	C	C	G	G	G	A	A	G	A	A	G	A	A	G	G
untreated																				
BE3					0±0															
HF-BE3					0.4±0															
					0±0															
					0±0															
EMX1 off target 7	G	A	G	C	C	G	G	A	G	C	A	A	G	A	A	G	G	A	G	G
untreated																				
BE3					0±0															
HF-BE3					0±0															
					0±0															
					0±0															
EMX1 off target 8	A	A	G	T	C	C	G	A	G	G	A	A	G	A	A	G	A	A	G	G
untreated																				
BE3					0±0															
HF-BE3					0±0															
					0±0															
					0±0															
EMX1 off target 9	G	A	A	T	C	C	A	A	G	C	A	G	A	G	A	A	G	A	G	A
untreated																				
BE3					0±0															
HF-BE3					0.1±0															
					0±0															
					0±0															
EMX1 off target 10	A	C	G	T	C	T	G	A	G	C	A	A	G	A	A	G	A	A	T	G
untreated																				
BE3					0±0															
HF-BE3					1.5±0.1															
					1.1±0.2															
					0±0															

numbers are $\mu \pm \sigma$ from three independent replicates

FIGURE 76 (CONTINUED)

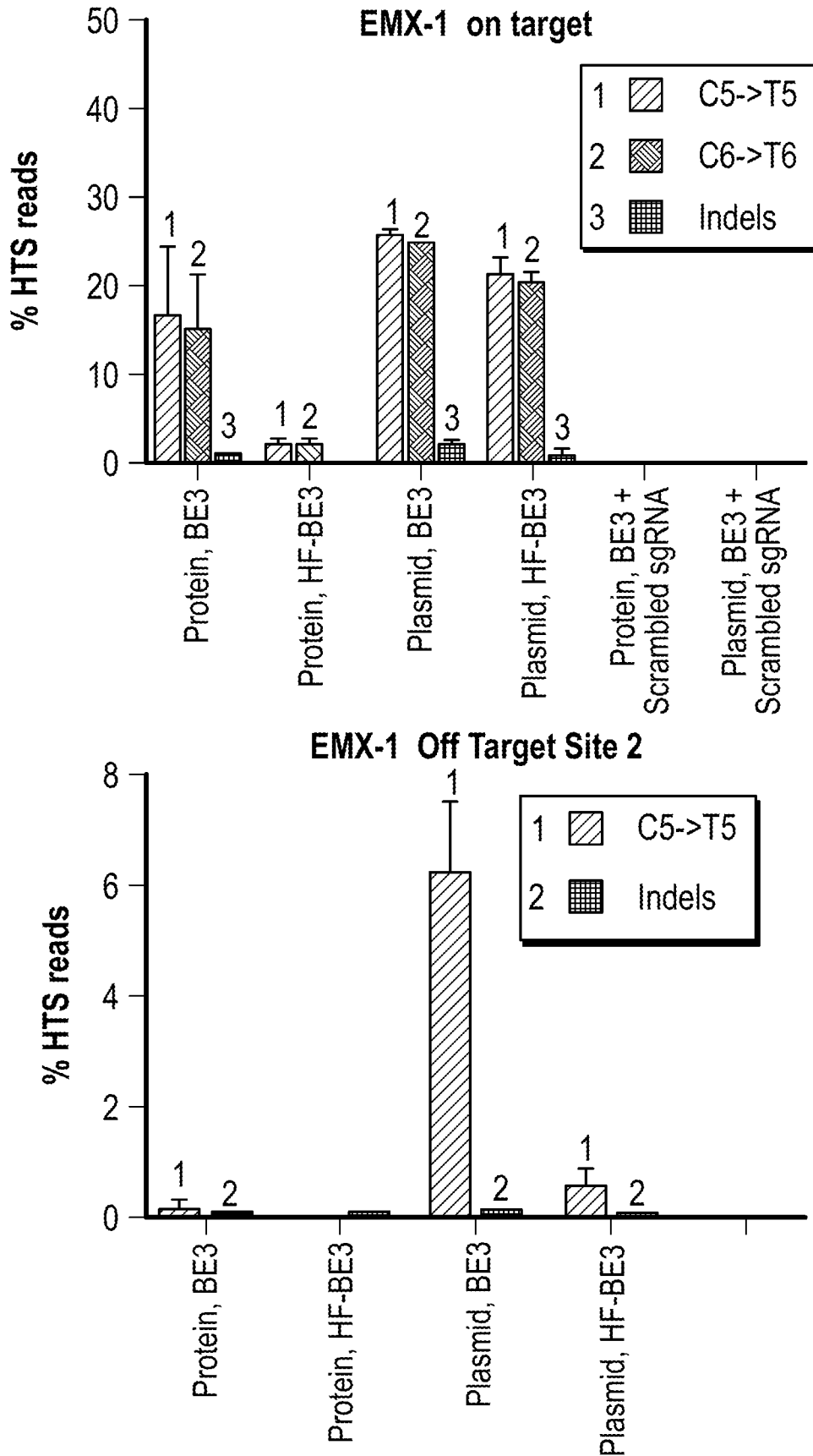


FIGURE 77

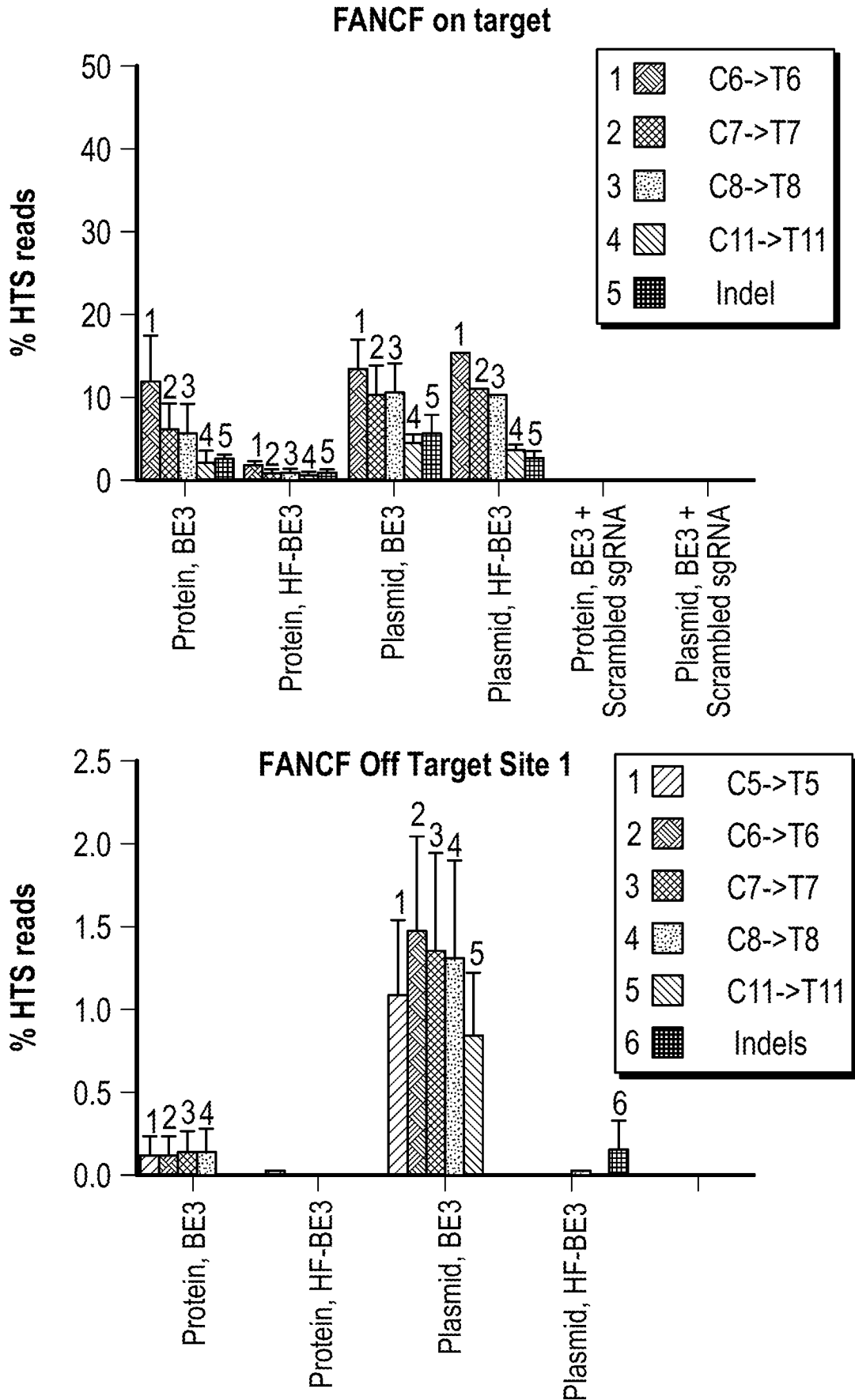


FIGURE 78

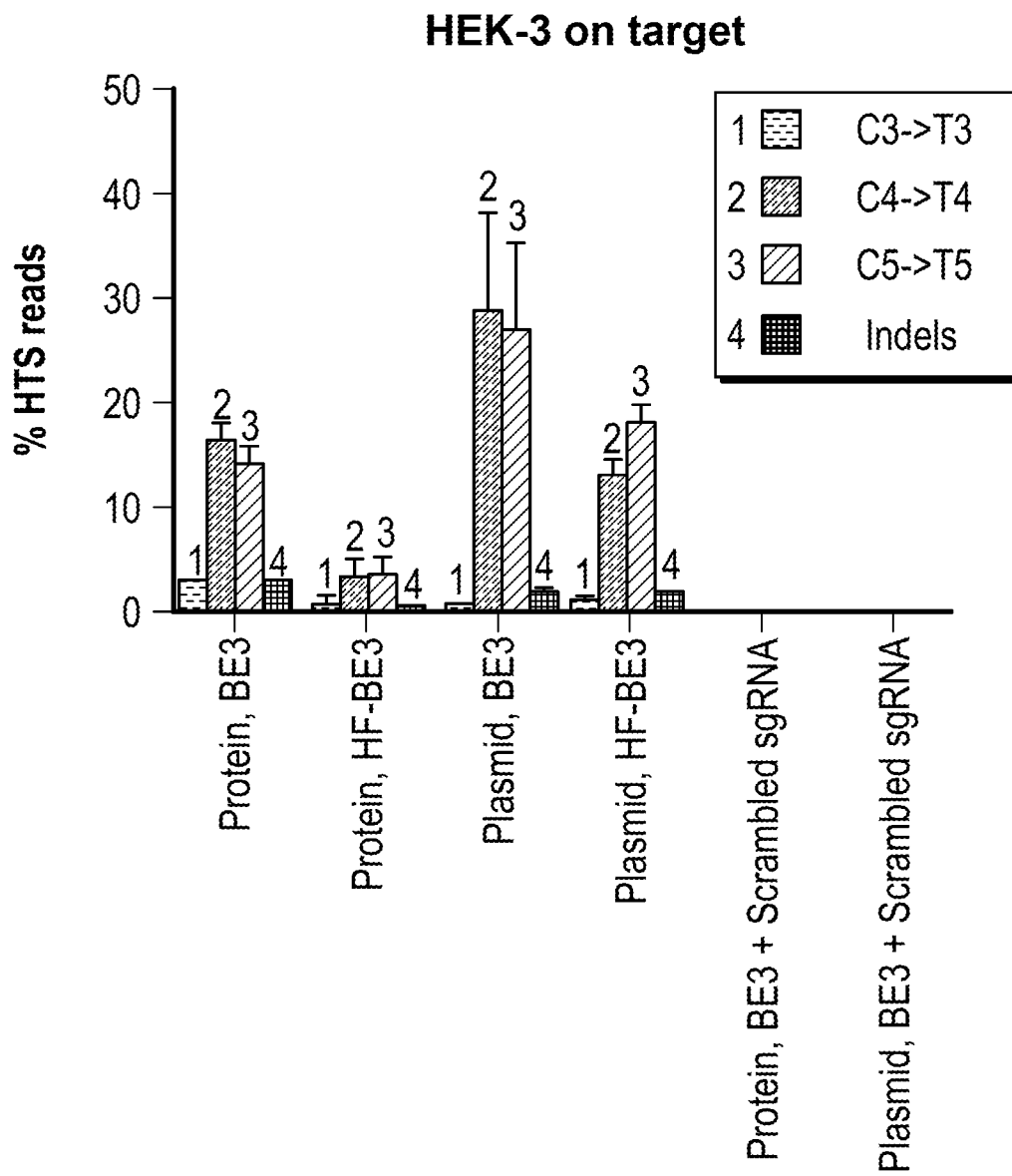


FIGURE 79

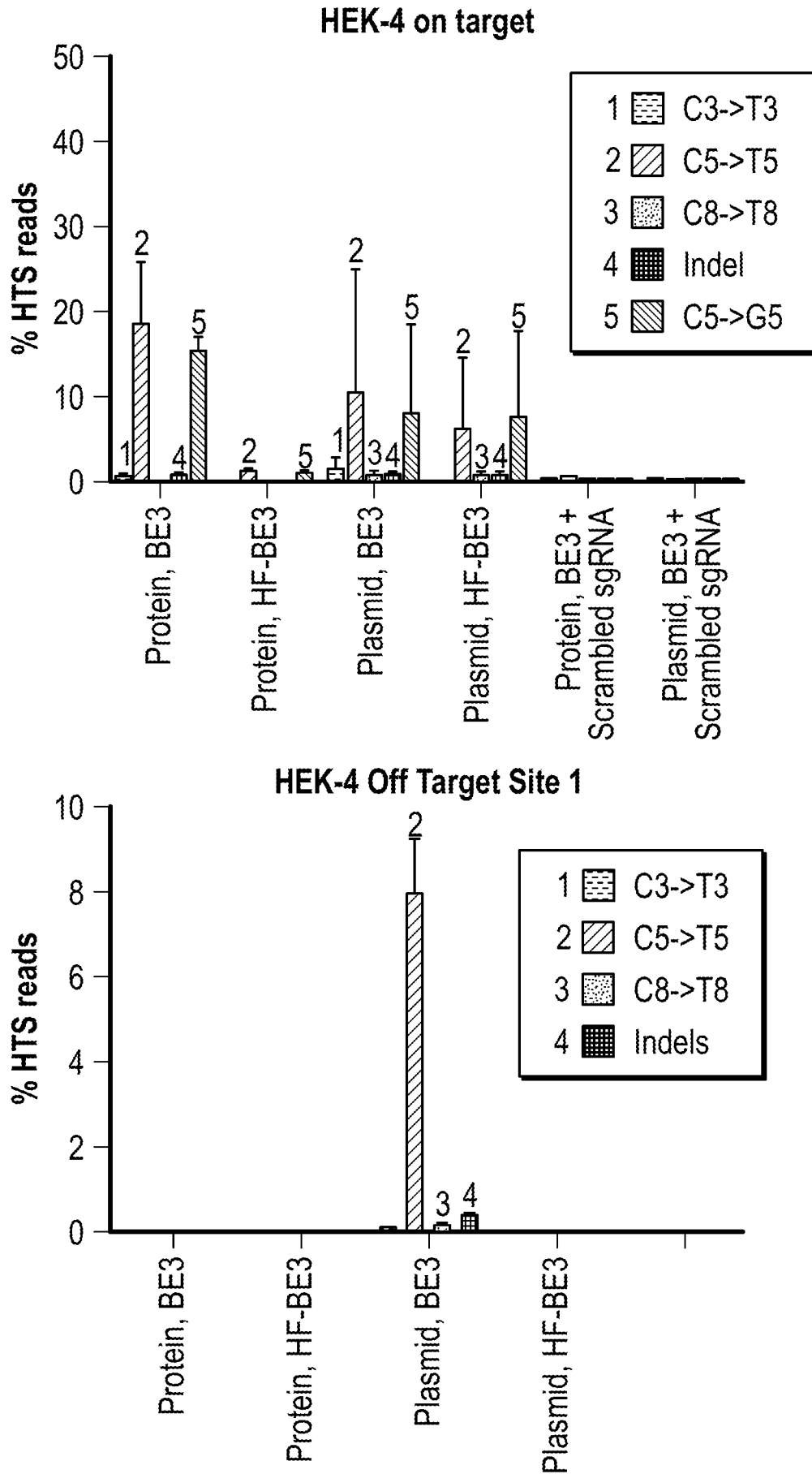
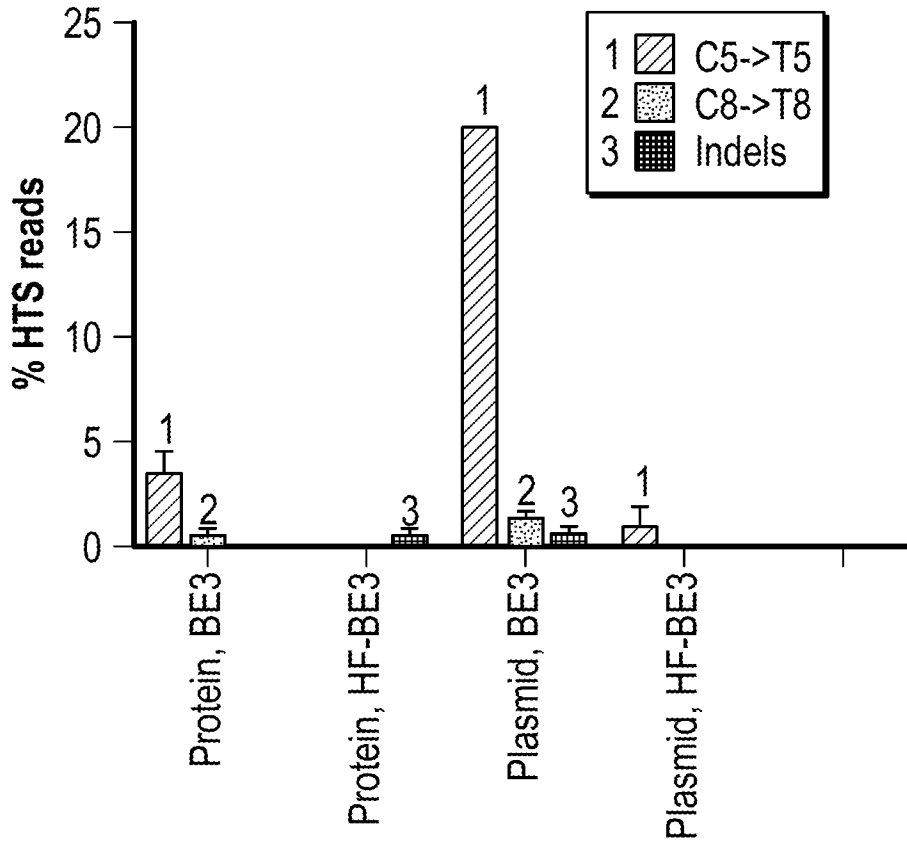


FIGURE 80

HEK-4 Off Target Site 3



HEK-4 Off Target Site 4

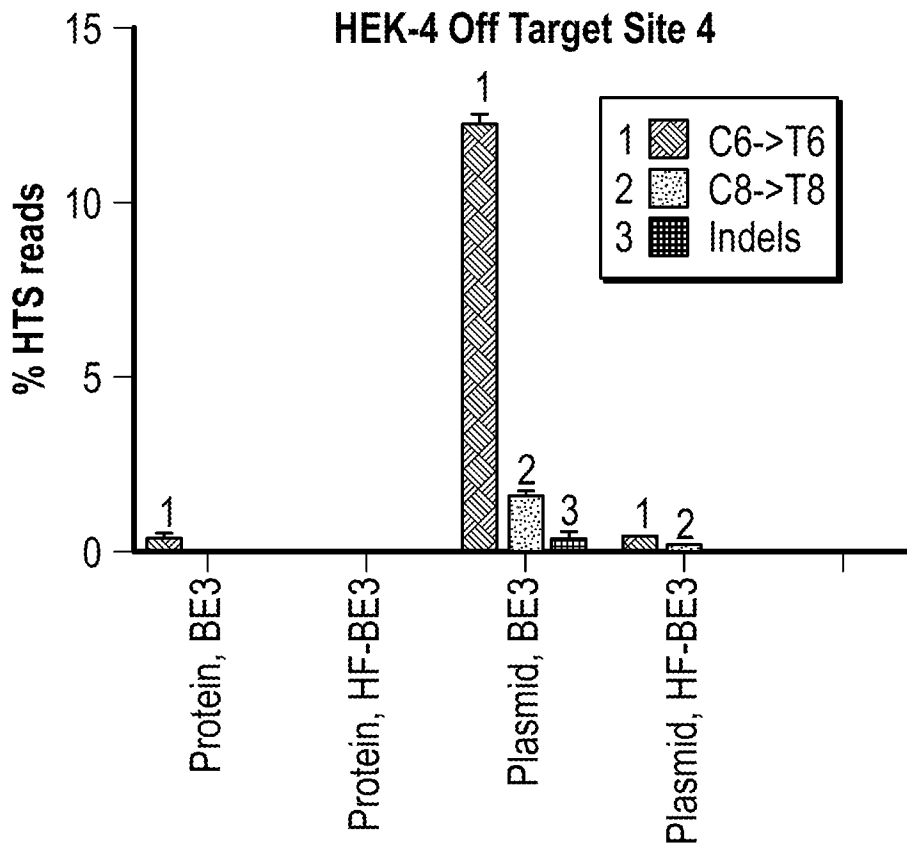


FIGURE 80 (CONTINUED)

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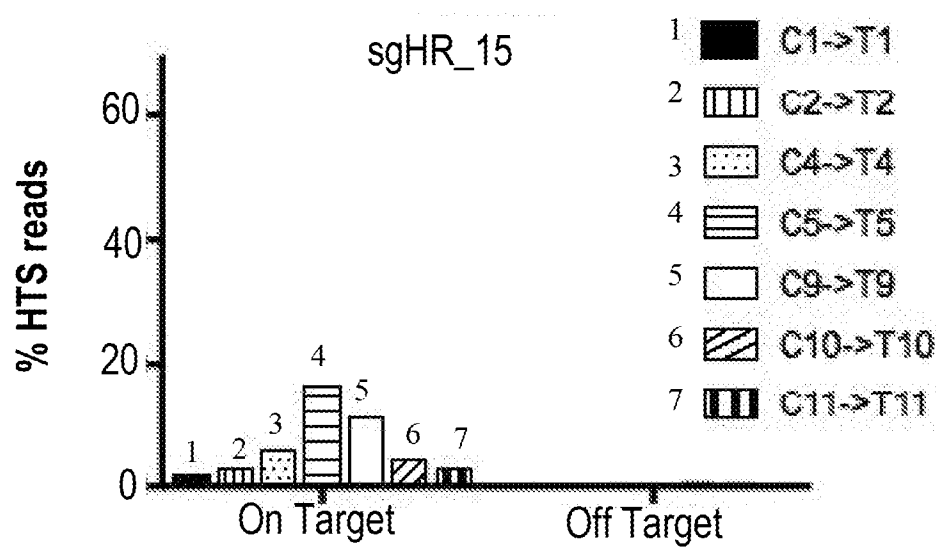
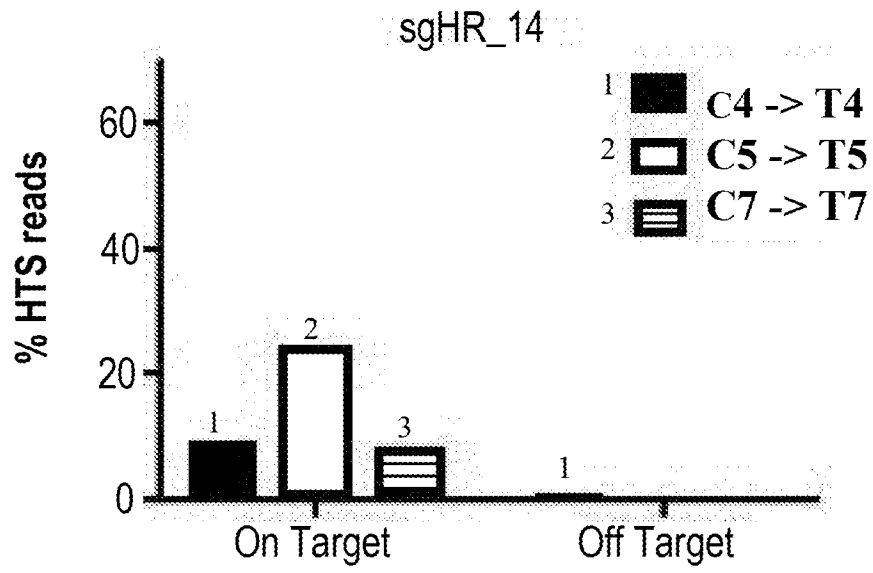
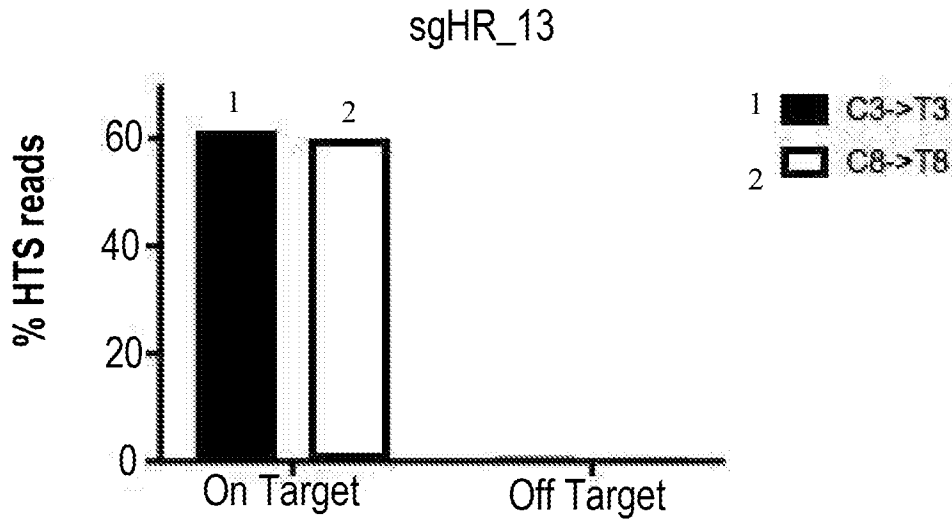


FIGURE 81

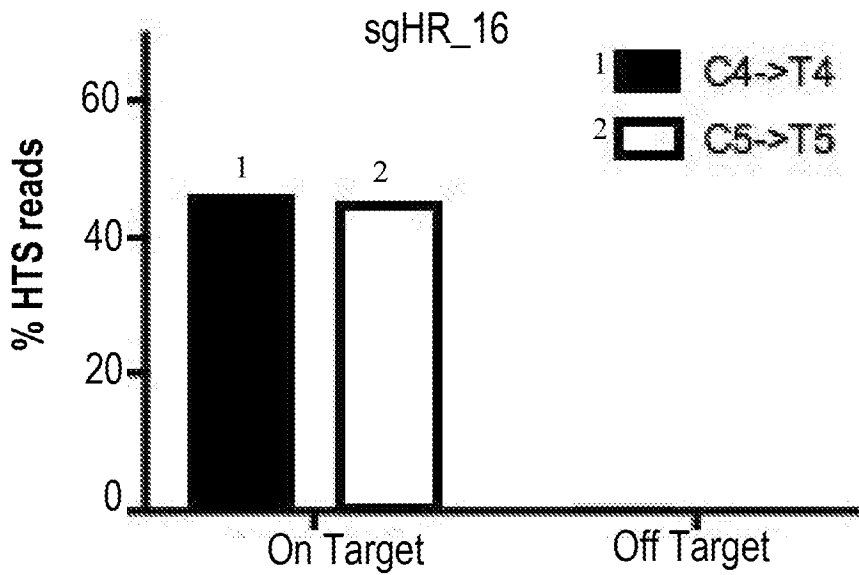
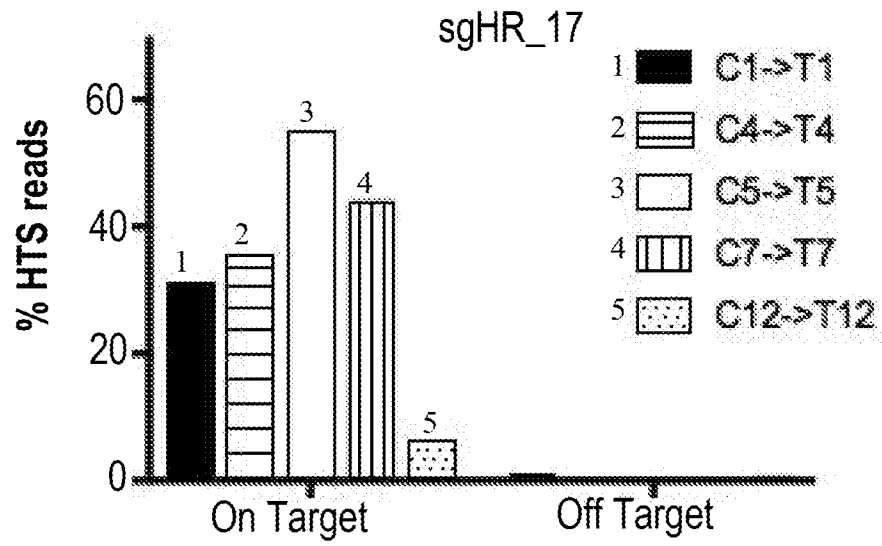


FIGURE 82

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sgHR_13

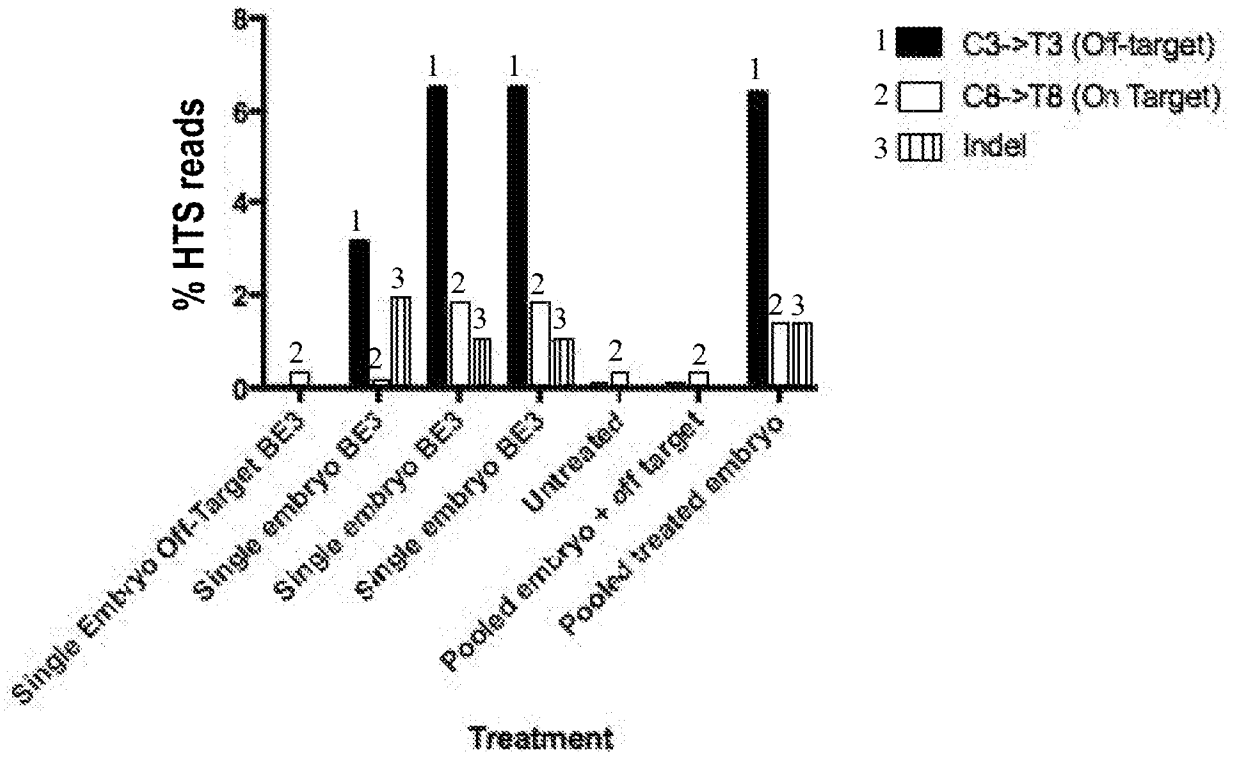


FIGURE 83

sgHR_16

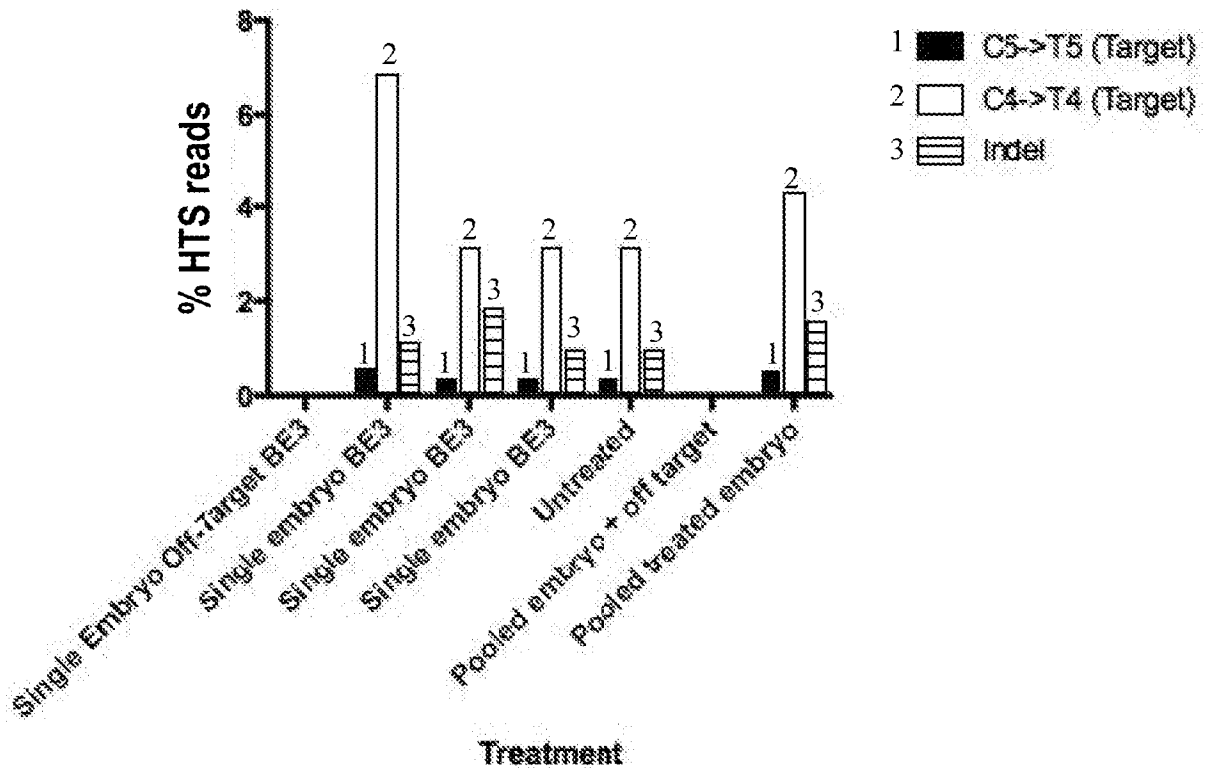


FIGURE 84

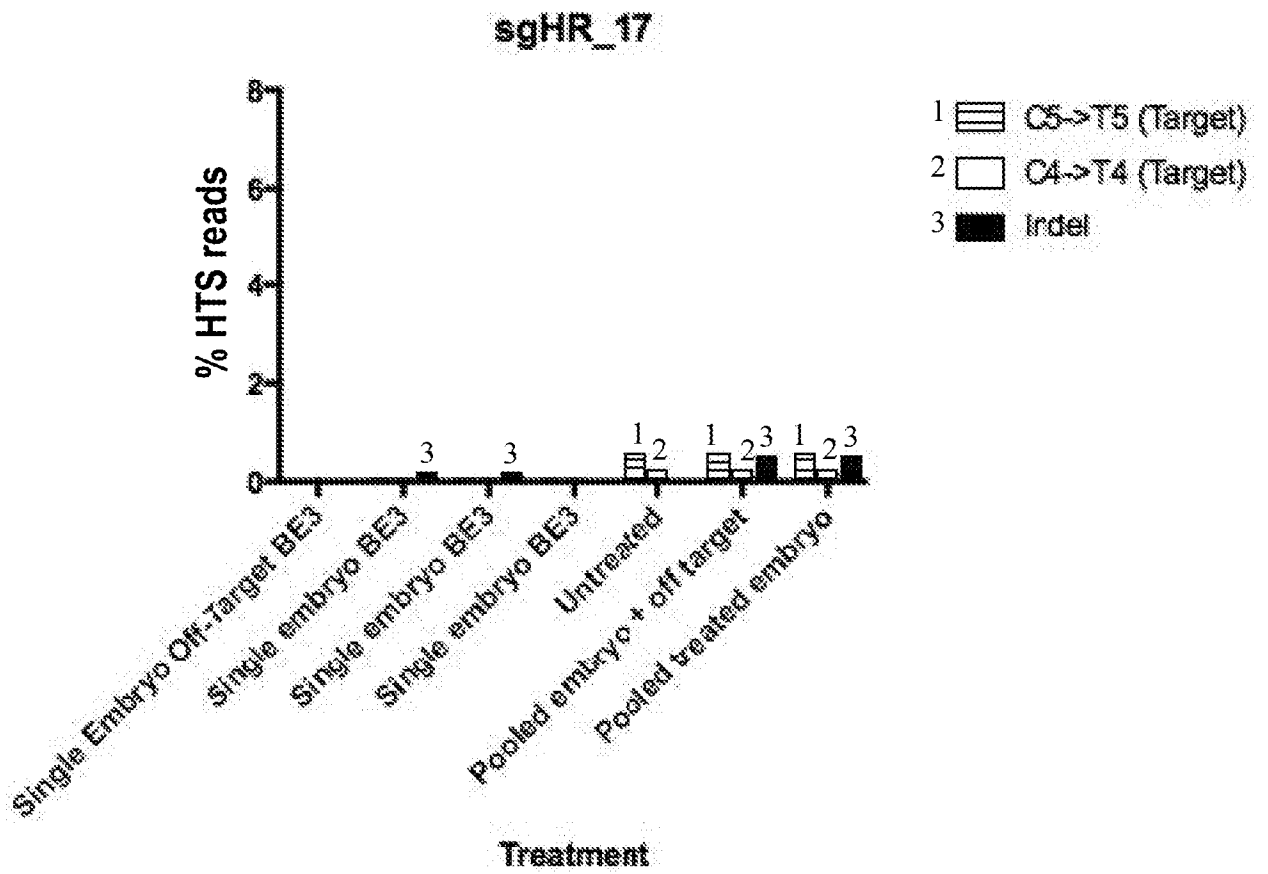


FIGURE 85

Possible Changes Using C→T Base Editors

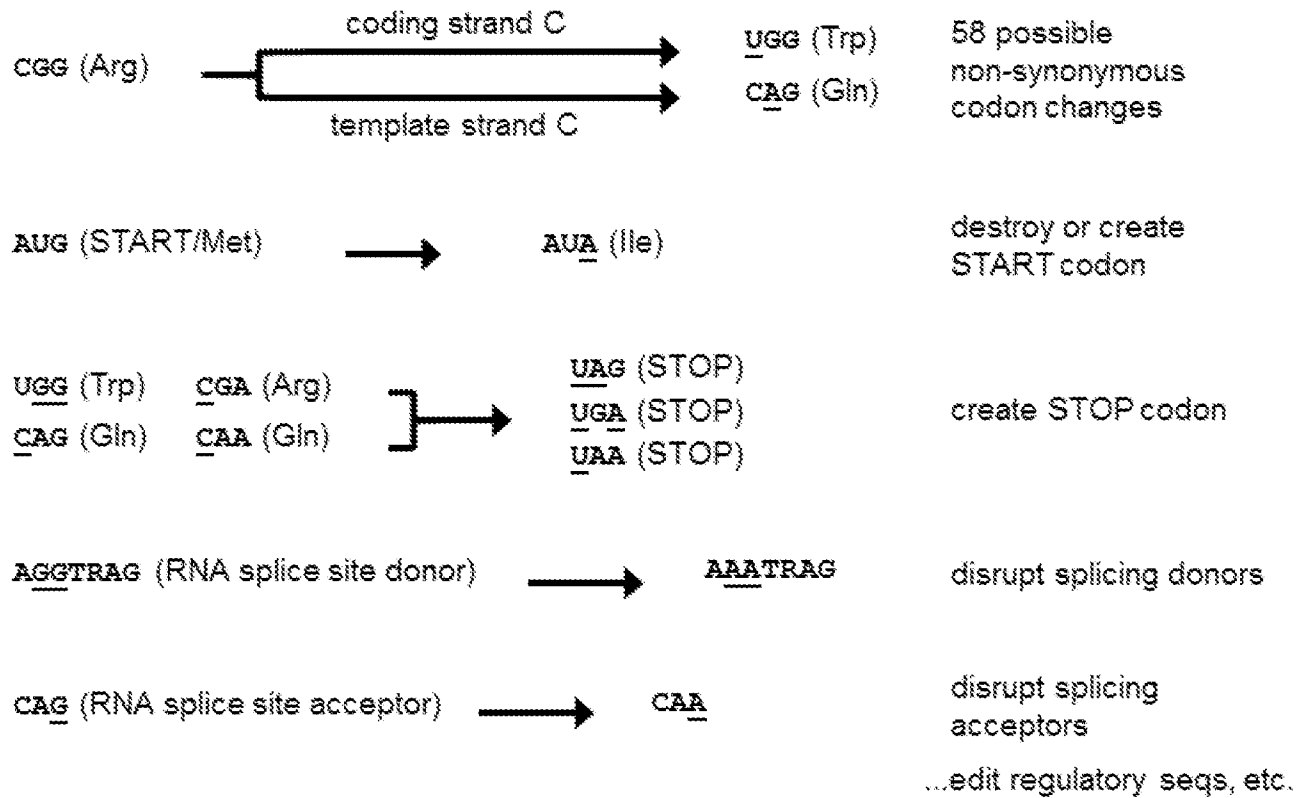
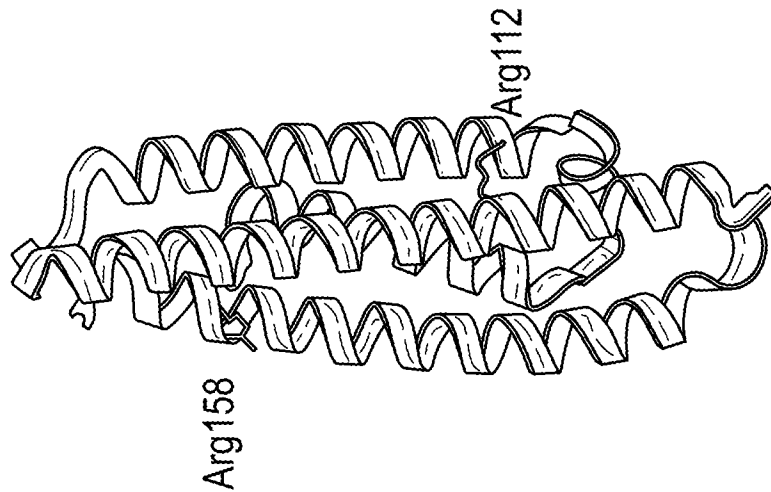


FIGURE 86

Base Editing APOE4 to APOE3r



- Apolipoprotein E has four isoforms that differ at amino acids 112 and 158
- APOE4 is the largest and most common genetic risk factor for late-onset Alzheimer's disease

allele	residue 112	residue 158	AD risk (known or presumed)
APOE2	Cys TGC	Cys TGC	low
APOE3	Cys TGC	Arg CGC	neutral
APOE3r	Arg CGC	Cys TGC	neutral
APOE4	Arg CGC	Arg BE3 CGC	high

FIGURE 87

untreated APOE4/C158R	Lys			Arg 152			Leu			Ala			Val			Tyr			Gln			indel % 0.0
	A	A	G	C ₆	G	C	C	T	G	C	G	A	G	T	G	A	C	C	A	C	A	
	G	A	A	G	C	C	C	T	G	G	C	A	T	G	G	T	A	C	A	C	A	
A	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	0.0	0.0	0.0	100.0	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	100.0	0.0	
G	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.9	
T	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	

Cas9 + HDR APOE4/C158R	Lys			Arg 152 → Cys			Leu			Ala			Val			Tyr			Gln			indel% 26.1
	A	A	G	C ₆	G	C	C	T	G	C	G	A	G	T	G	A	C	C	A	C	A	
	G	A	A	G	C	C	C	T	G	G	C	A	T	G	G	T	A	C	A	C	A	
A	0.0	100.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	99.4	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	0.0	0.0	0.0	99.7	0.0	99.9	99.9	0.0	0.0	0.0	100.0	0.5	0.0	0.0	0.0	0.0	0.0	0.0	100.0	100.0	0.0	
G	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	100.0	99.9	0.0	0.0	99.6	0.6	99.9	0.2	0.0	0.0	0.0	0.0	100.0	
T	0.0	0.0	0.0	0.3	0.0	0.1	0.1	100.0	0.0	0.0	0.0	0.1	0.4	99.3	0.1	99.8	0.0	0.0	0.0	0.0	0.0	

BE3-treated APOE4/C158R	Lys			Arg 152 → Cys			Leu → Leu			Ala			Val			Tyr			Gln			indel% 4.6
	A	A	G	C ₆	G	C ₇	C ₂	T	G	C	G	A	G	T	G	A	C	C	A	C	A	
	G	A	A	G	C	C	C	T	G	G	C	A	T	G	G	T	A	C	A	C	A	
A	0.1	100.0	100.0	0.0	0.5	0.0	1.3	0.9	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
C	0.0	0.0	0.0	23.7	0.0	47.4	43.5	0.0	0.0	0.0	0.0	99.9	0.0	0.0	0.0	0.0	0.0	0.0	100.0	100.0	0.0	
G	99.9	0.0	0.0	0.0	100.0	0.0	0.0	99.9	1.1	0.7	0.0	100.0	100.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	100.0	
T	0.0	0.0	0.0	74.9	0.1	50.2	55.0	100.0	0.0	0.0	0.1	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	

FIGURE 88

BE3-treated PRMP R37X	Gly			Ser → Ser			Arg 27 → Stop			Tyr			Pro			Gly			Glu			
	G	G	C	A	G	C ₇	C ₈	G	A	T	A	C	C	C	G	G	G	C	A	G	G	
A	0.0	0.0	0.1	100.0	0.0	1.1	0.3	0.1	100.0	0.0	100.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0
C	0.0	0.0	98.9	0.0	0.0	54.9	57.8	0.0	0.0	0.0	0.0	99.8	99.8	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0	0.0
G	100.0	100.0	0.0	0.0	100.0	2.1	0.6	99.9	0.0	0.0	0.0	0.0	0.0	0.0	100.0	100.0	100.0	100.0	0.0	0.0	100.0	100.0
T	0.0	0.0	1.1	0.0	0.0	41.9	41.2	0.0	0.0	100.0	0.0	0.2	0.2	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0

FIGURE 89

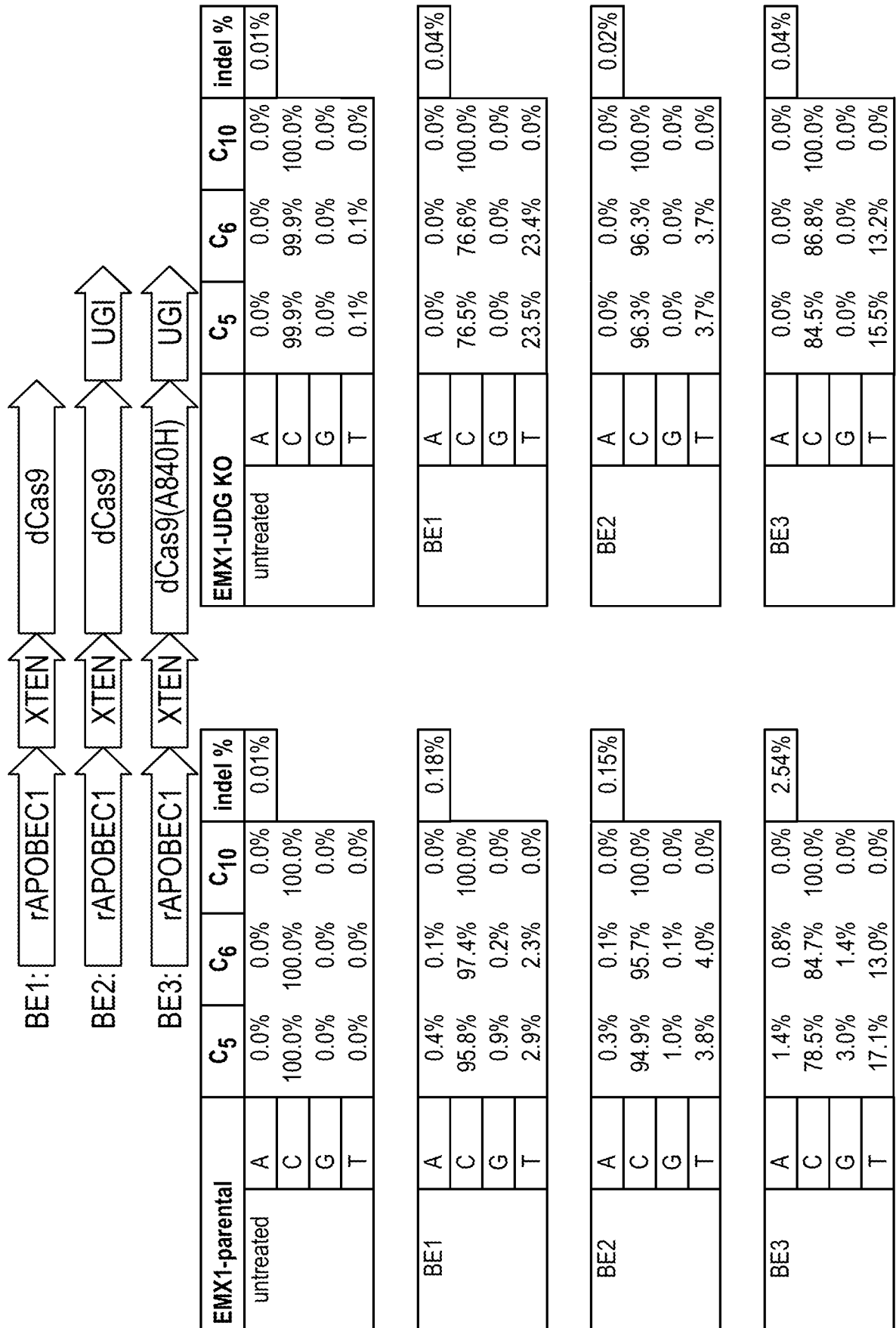


FIGURE 90

FANCF-parental		C6	C7	C8	C11	indel %
untreated	A	0.0%	0.0%	0.0%	0.0%	0.03%
	C	99.9%	99.9%	99.9%	99.9%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	0.1%	0.1%	0.1%	0.0%	

FANCF-UDG KO		C6	C7	C8	C11	indel %
untreated	A	0.0%	0.0%	0.0%	0.0%	0.02%
	C	99.9%	99.9%	100.0%	100.0%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	0.1%	0.0%	0.0%	0.0%	

BE1		C6	C7	C8	C11	indel %
BE1	A	0.0%	0.1%	0.0%	0.0%	0.09%
	C	60.9%	61.1%	61.3%	94.7%	
	G	0.1%	0.1%	0.1%	0.0%	
	T	39.0%	38.8%	38.6%	5.2%	

BE2		C6	C7	C8	C11	indel %
BE2	A	0.0%	0.0%	0.0%	0.0%	0.02%
	C	88.6%	88.6%	88.7%	98.9%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	11.4%	11.4%	11.3%	1.1%	

BE3		C6	C7	C8	C11	indel %
BE3	A	0.0%	0.1%	0.1%	0.0%	0.35%
	C	47.9%	48.5%	48.8%	55.8%	
	G	0.0%	0.1%	0.3%	0.2%	
	T	52.1%	51.3%	50.8%	44.0%	

BE1		C6	C7	C8	C11	indel %
BE1	A	0.5%	0.5%	0.3%	0.0%	0.13%
	C	94.2%	97.8%	97.9%	99.8%	
	G	0.7%	0.0%	0.0%	0.0%	
	T	4.7%	1.6%	1.8%	0.2%	

BE2		C6	C7	C8	C11	indel %
BE2	A	0.3%	0.4%	0.2%	0.0%	0.25%
	C	95.3%	97.3%	97.6%	99.8%	
	G	0.4%	0.1%	0.0%	0.0%	
	T	4.0%	2.3%	2.1%	0.1%	

BE3		C6	C7	C8	C11	indel %
BE3	A	2.4%	3.2%	2.2%	0.9%	18.88%
	C	60.3%	72.6%	73.8%	86.6%	
	G	1.2%	0.6%	0.4%	0.3%	
	T	36.2%	23.6%	23.5%	12.3%	

FIGURE 90 (CONTINUED)

HEK3-JDGD KO		C3	C4	C5	C9	indel %
untreated	A	0.0%	0.0%	0.0%	0.0%	0.00%
	C	100.0%	100.0%	100.0%	100.0%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	0.0%	0.0%	0.0%	0.0%	

BE1		C3	C4	C5	C9	indel %
BE1	A	0.0%	0.0%	0.0%	0.0%	0.03%
	C	96.0%	50.3%	41.1%	96.3%	
	G	0.0%	0.0%	0.1%	0.1%	
	T	4.0%	49.6%	58.8%	3.6%	

BE2		C3	C4	C5	C9	indel %
BE2	A	0.0%	0.0%	0.0%	0.0%	0.00%
	C	99.0%	80.2%	73.2%	98.8%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	1.0%	19.7%	26.8%	1.1%	

BE3		C3	C4	C5	C9	indel %
BE3	A	0.0%	0.0%	0.0%	0.0%	0.09%
	C	98.5%	55.6%	40.3%	99.4%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	1.4%	44.3%	59.7%	0.6%	

HEK3-parental		C3	C4	C5	C9	indel %
untreated	A	0.0%	0.0%	0.0%	0.0%	0.00%
	C	100.0%	99.9%	99.9%	100.0%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	0.0%	0.0%	0.0%	0.0%	

BE1		C3	C4	C5	C9	indel %
BE1	A	0.0%	0.4%	0.3%	0.1%	0.07%
	C	99.9%	96.3%	94.4%	99.8%	
	G	0.0%	0.1%	1.9%	0.1%	
	T	0.1%	3.2%	3.4%	0.1%	

BE2		C3	C4	C5	C9	indel %
BE2	A	0.0%	0.2%	0.3%	0.1%	0.05%
	C	100.0%	97.7%	96.1%	99.8%	
	G	0.0%	0.0%	1.2%	0.1%	
	T	0.0%	2.1%	2.4%	0.1%	

BE3		C3	C4	C5	C9	indel %
BE3	A	0.1%	2.9%	2.4%	0.3%	3.27%
	C	99.4%	62.6%	55.9%	98.8%	
	G	0.0%	1.5%	10.4%	0.3%	
	T	0.6%	33.0%	31.3%	0.6%	

FIGURE 90 (CONTINUED)

HEK4-JDUG KO		C3	C5	C8	C11	indel %
untreated	A	0.0%	0.0%	0.0%	0.0%	0.00%
	C	100.0%	100.0%	99.9%	100.0%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	0.0%	0.0%	0.1%	0.0%	

BE1	A	0.0%	0.0%	0.0%	0.0%	0.00%
	C	99.9%	67.5%	99.0%	100.0%	
	G	0.0%	0.0%	0.1%	0.0%	
	T	0.1%	32.5%	1.0%	0.0%	

BE2	A	0.0%	0.0%	0.0%	0.0%	0.00%
	C	100.0%	91.9%	99.7%	100.0%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	0.0%	8.1%	0.2%	0.0%	

BE3	A	0.0%	0.0%	0.0%	0.0%	0.03%
	C	99.9%	65.9%	99.7%	100.0%	
	G	0.0%	0.3%	0.0%	0.0%	
	T	0.1%	33.8%	0.3%	0.0%	

HEK4-parental		C3	C5	C8	C11	indel %
untreated	A	0.0%	0.0%	0.0%	0.0%	0.00%
	C	100.0%	100.0%	99.9%	100.0%	
	G	0.0%	0.0%	0.0%	0.0%	
	T	0.0%	0.0%	0.1%	0.0%	

BE1	A	0.0%	1.4%	0.0%	0.0%	0.08%
	C	100.0%	95.5%	99.9%	100.0%	
	G	0.0%	2.2%	0.0%	0.0%	
	T	0.0%	0.9%	0.1%	0.0%	

BE2	A	0.0%	0.4%	0.0%	0.0%	0.07%
	C	100.0%	97.7%	99.9%	100.0%	
	G	0.0%	1.1%	0.0%	0.0%	
	T	0.0%	0.7%	0.1%	0.0%	

BE3	A	0.0%	7.1%	0.0%	0.0%	6.05%
	C	99.9%	58.5%	99.7%	100.0%	
	G	0.0%	17.4%	0.0%	0.0%	
	T	0.1%	17.0%	0.2%	0.0%	

FIGURE 90 (CONTINUED)

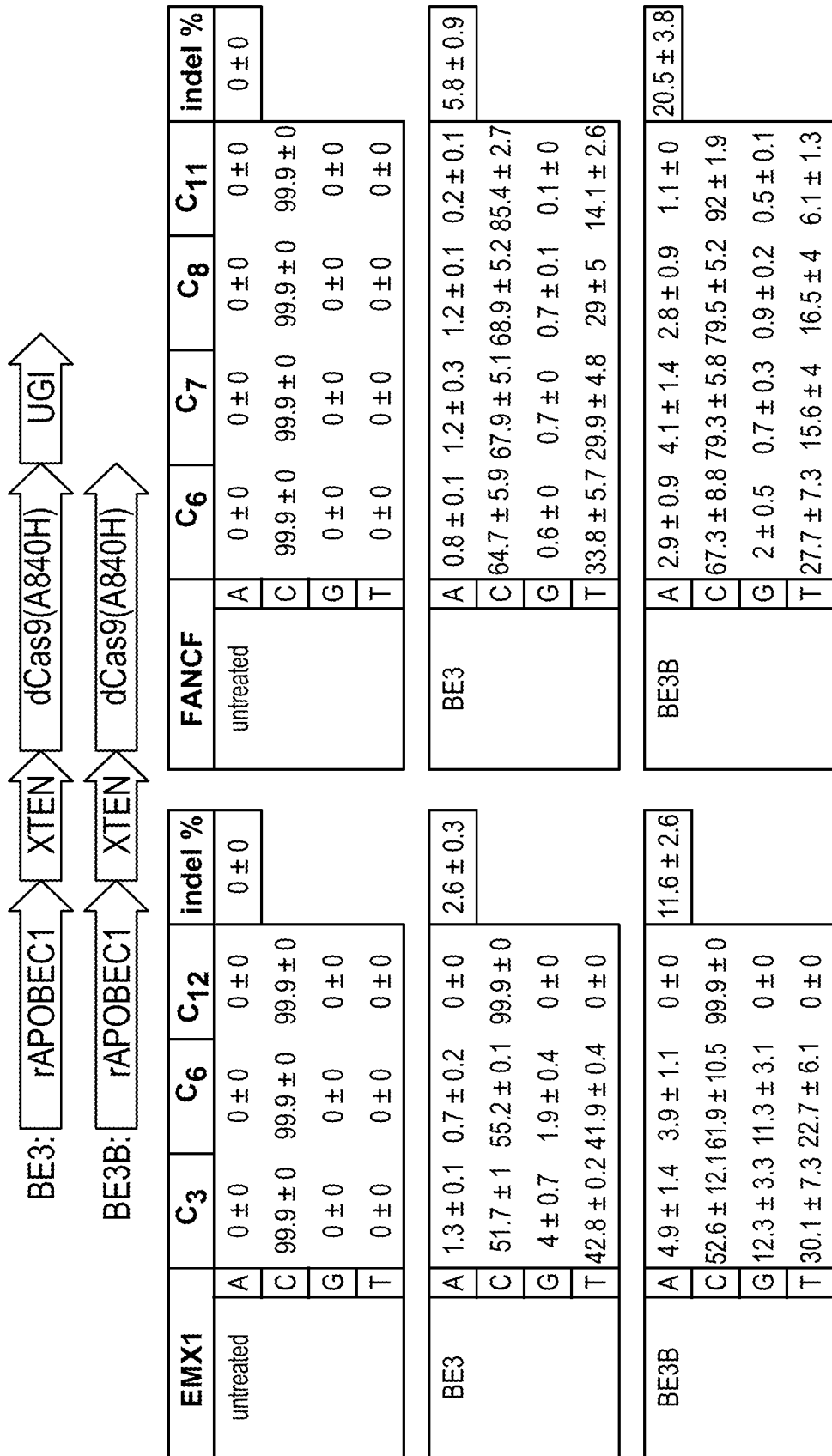
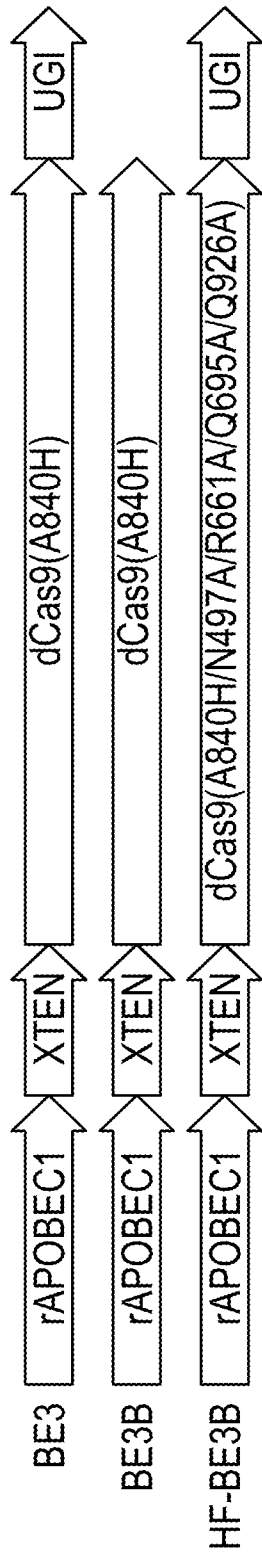


FIGURE 91



HEK3	C6	C7	C8	C11	indel %	HEK4	C6	C7	C8	C11	indel %	
untreated	A	0 ± 0	0 ± 0	0 ± 0	0 ± 0	untreated	A	0 ± 0	0 ± 0	0 ± 0	0 ± 0	
	C	99.9 ± 0	99.9 ± 0	99.9 ± 0	99.9 ± 0		C	99.9 ± 0	99.9 ± 0	99.9 ± 0	99.9 ± 0	
	G	0 ± 0	0 ± 0	0 ± 0	0 ± 0		G	0 ± 0	0 ± 0	0 ± 0	0 ± 0	
	T	0 ± 0	0 ± 0	0 ± 0	0 ± 0		T	0 ± 0	0 ± 0	0 ± 0	0 ± 0	
BE3	A	0 ± 0	1.6 ± 0.2	1.5 ± 0.3	0.2 ± 0	BE3	A	0 ± 0	5.4 ± 1.3	0 ± 0	0 ± 0	3 ± 0.7
	C	98.7 ± 0.2	48.6 ± 5.4	40.8 ± 6.7	98.6 ± 0.1		C	98.7 ± 0	44.5 ± 8.2	98.9 ± 0	99.8 ± 0	
	G	0 ± 0	1.4 ± 0.2	10.5 ± 1.2	0.3 ± 0		G	0 ± 0	18.1 ± 2.3	0 ± 0	0 ± 0	
	T	1.1 ± 0.2	48.2 ± 4.9	47 ± 5.2	0.7 ± 0.1		T	1.1 ± 0.1	31.9 ± 4.8	0.9 ± 0.1	0.1 ± 0	
BE3B	A	0.1 ± 0	4.9 ± 0.8	3.8 ± 0.8	0.8 ± 0.1	BE3B	A	0 ± 0	10.6 ± 2.6	0.3 ± 0	0 ± 0	7.6 ± 0.7
	C	99.4 ± 0.1	57.6 ± 6.3	44.4 ± 8.4	97.7 ± 0.3		C	99.6 ± 0	45 ± 9.9	99.2 ± 0.1	99.9 ± 0	
	G	0 ± 0	3.3 ± 0.7	21.2 ± 2.8	0.7 ± 0.1		G	0 ± 0	32.8 ± 5.8	0 ± 0	0 ± 0	
	T	0.4 ± 0.1	33.9 ± 4.7	30.4 ± 4.7	0.6 ± 0		T	0.2 ± 0	11.4 ± 1.6	0.4 ± 0.2	0 ± 0	

FIGURE 91 (CONTINUED)

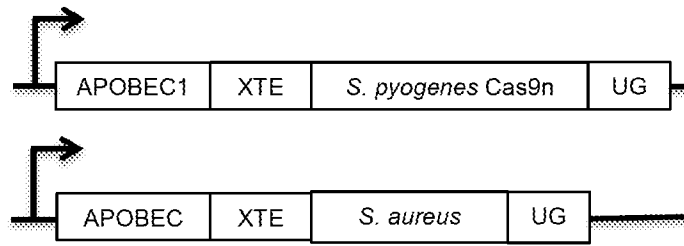


FIGURE 92A

Species	PAM	Base editor	Reference
<i>S. pyogenes</i>	NGG	BE3	Wild-type
	NGA	VQR, EQR BE3	Ref #7
	NGCG	VRER BE3	Ref #7
<i>S. aureus</i>	NNGRRT	SaBE3	Wild-type
	NNNRRT	SaKKHBE3	Ref #8

FIGURE 92B

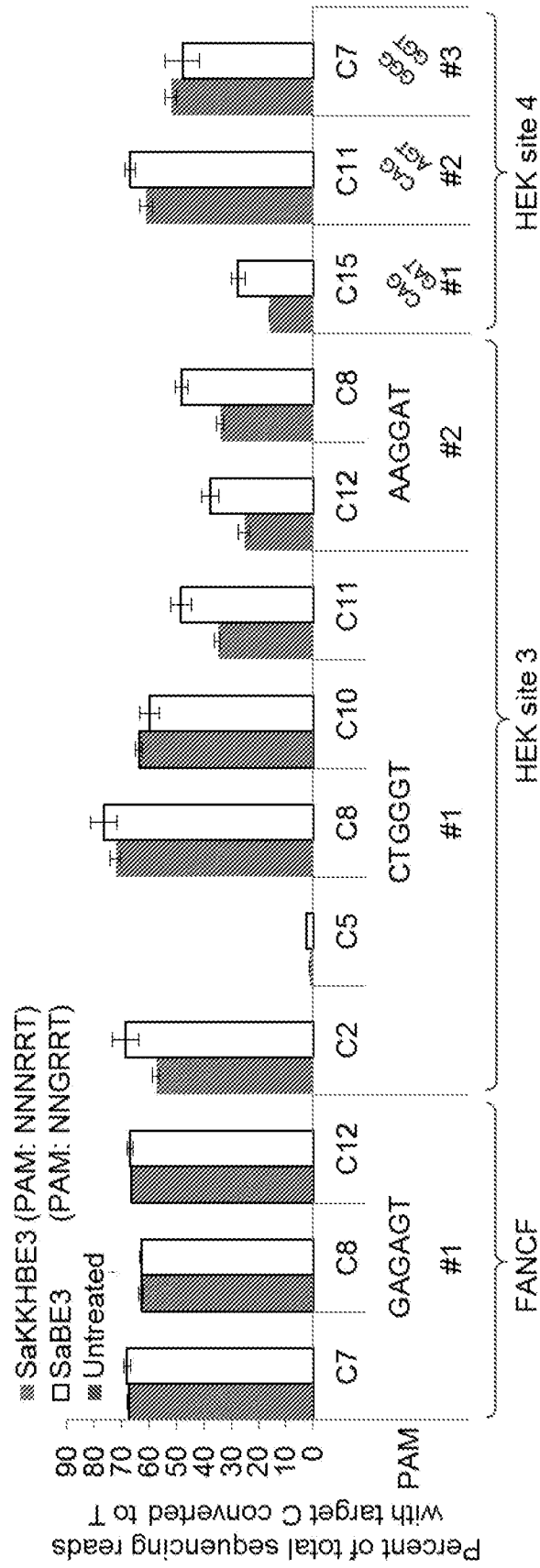


FIGURE 92C

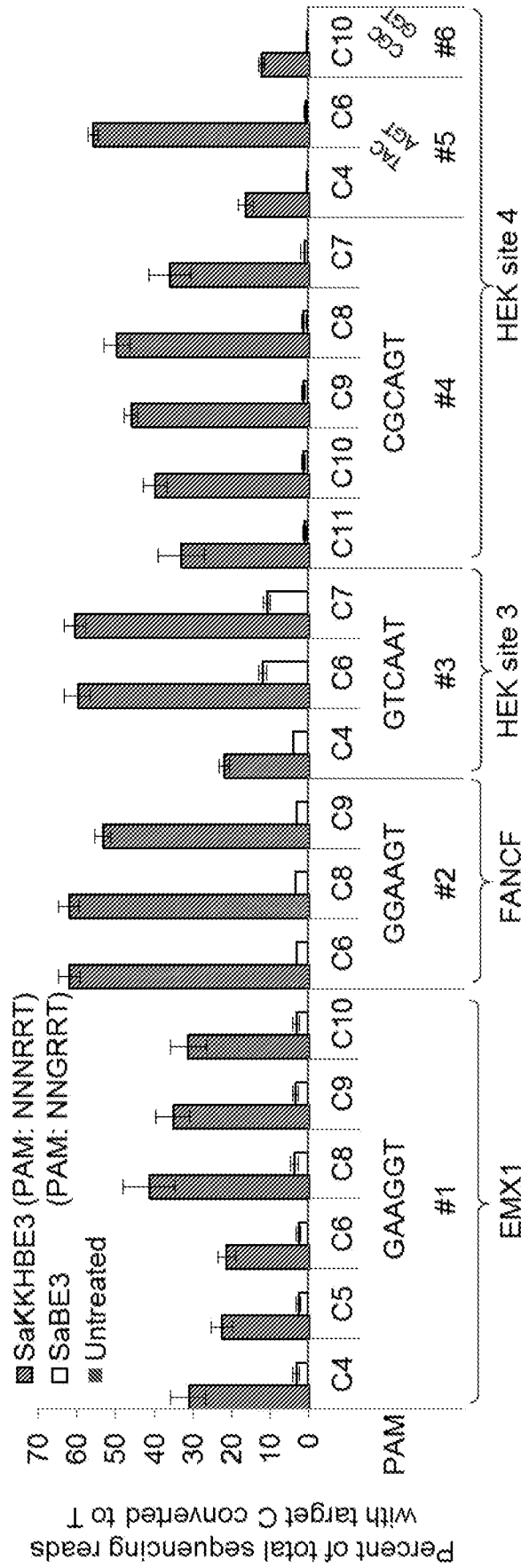


FIGURE 92D

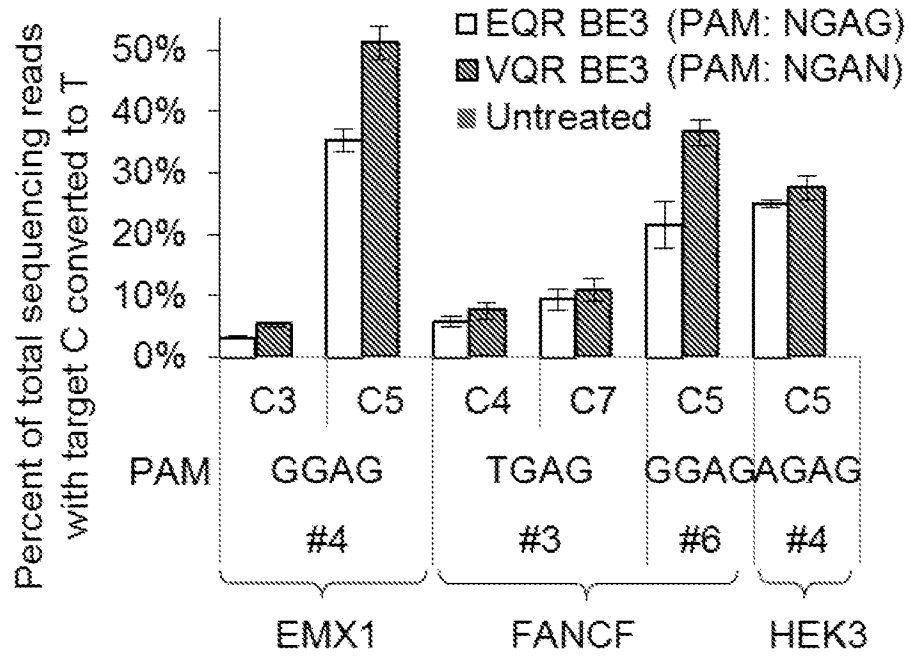


FIGURE 92E

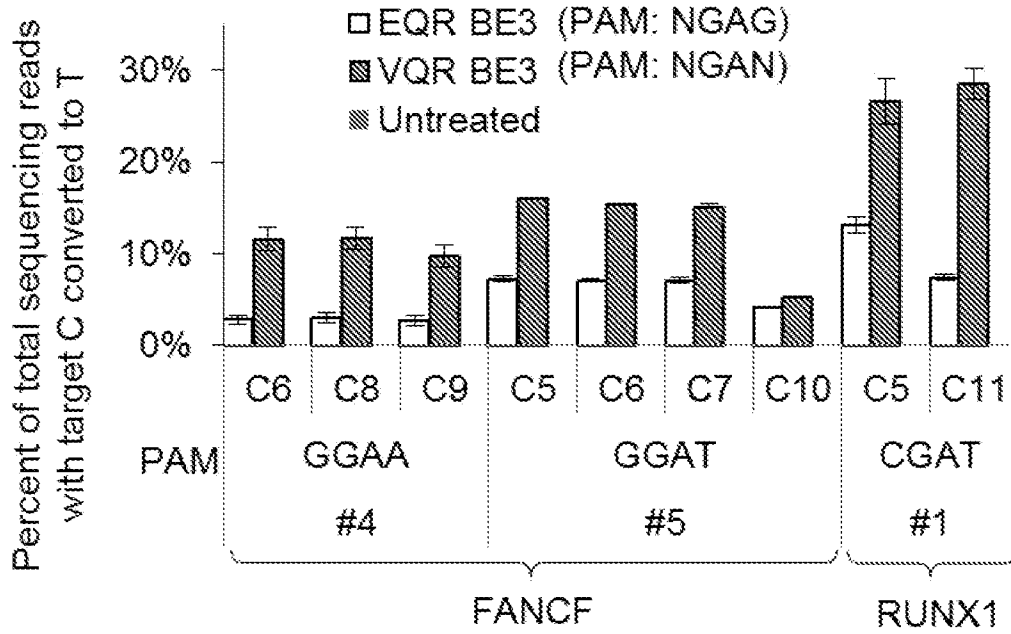


FIGURE 92F

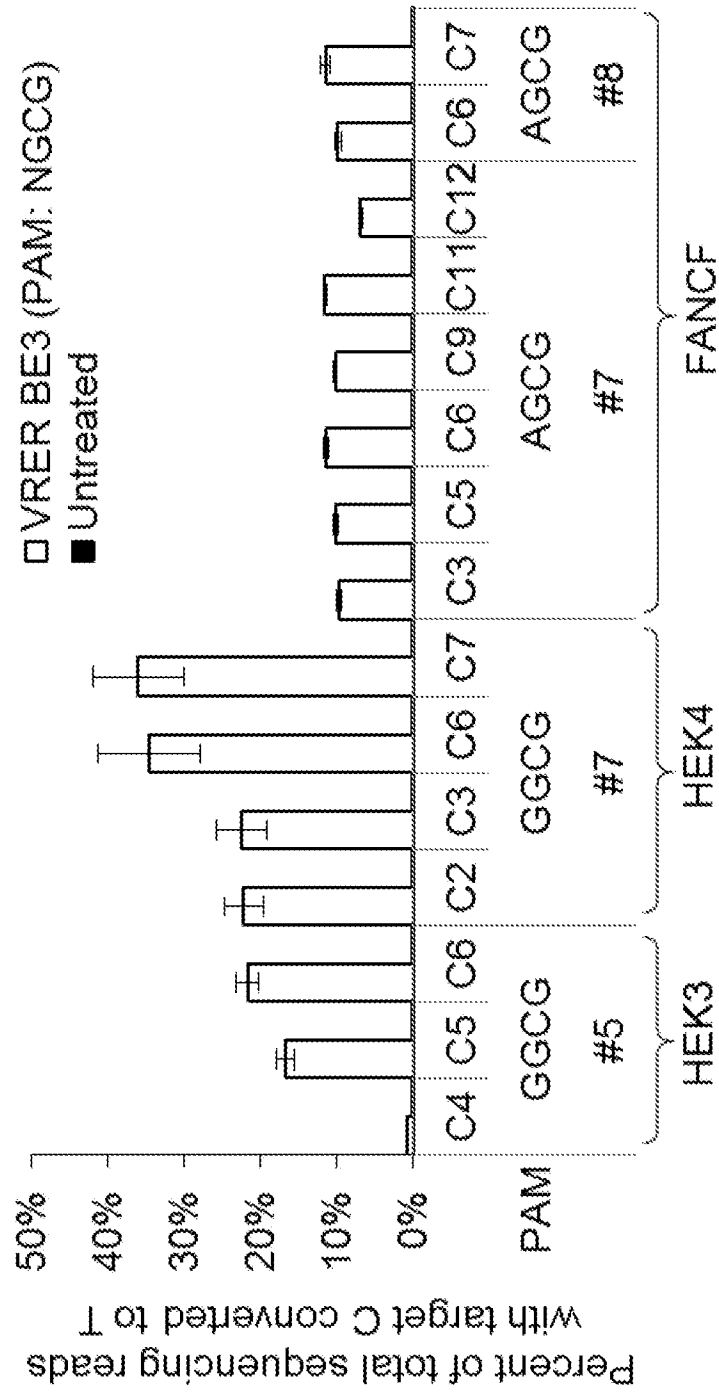
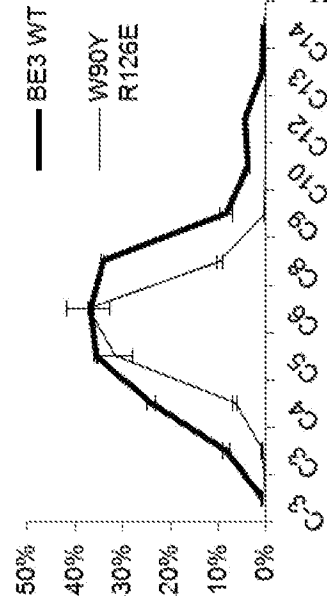
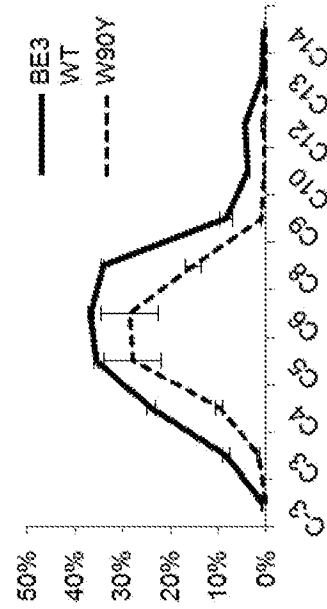
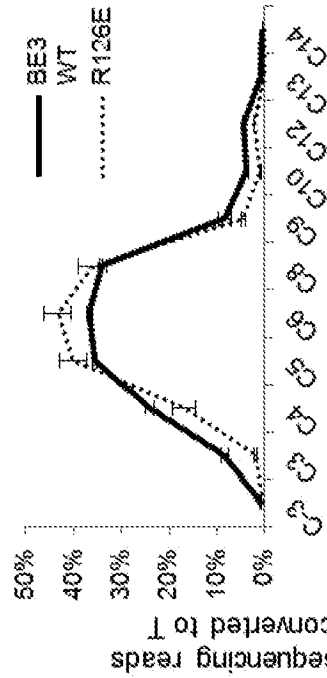


FIGURE 92G

site A: TGC₃C₄C₅C₆C₇C₈C₉C₁₀TC₁₂C₁₃C₁₄TGGCCCAGG



site B: AGAGC₅C₆C₇C₈C₉C₁₀C₁₁TC₁₃AAAGAGAGGGG

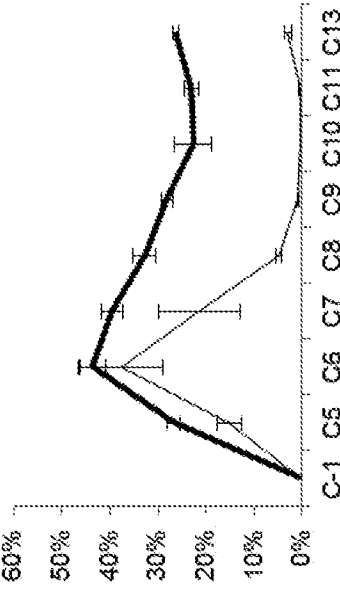
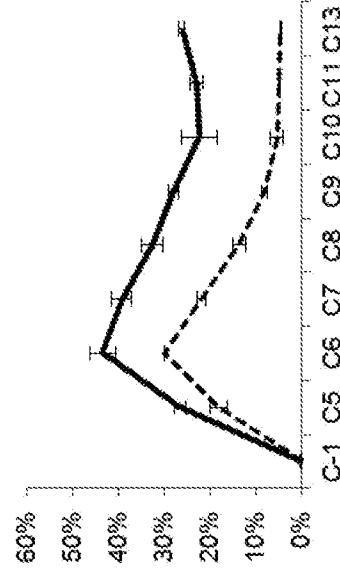
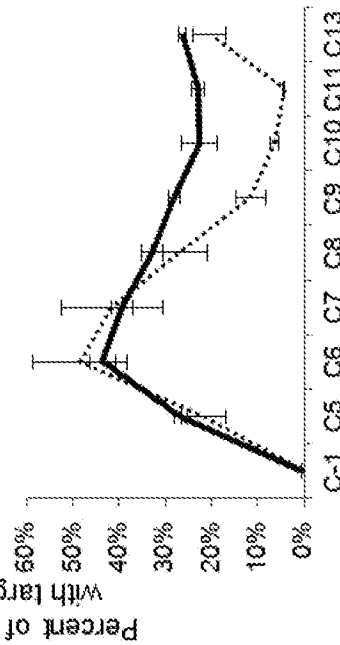


FIGURE 93A

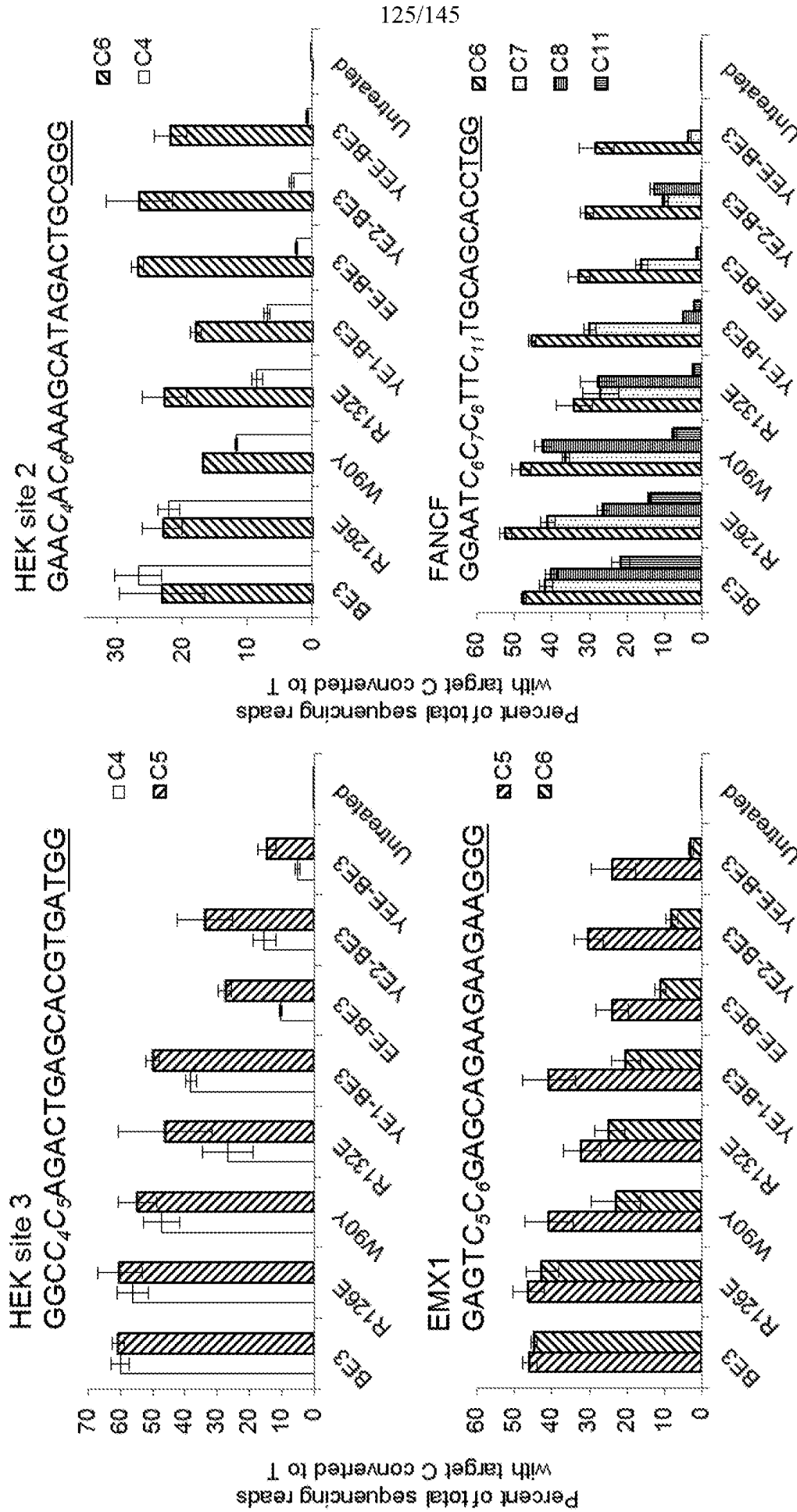


FIGURE 93B

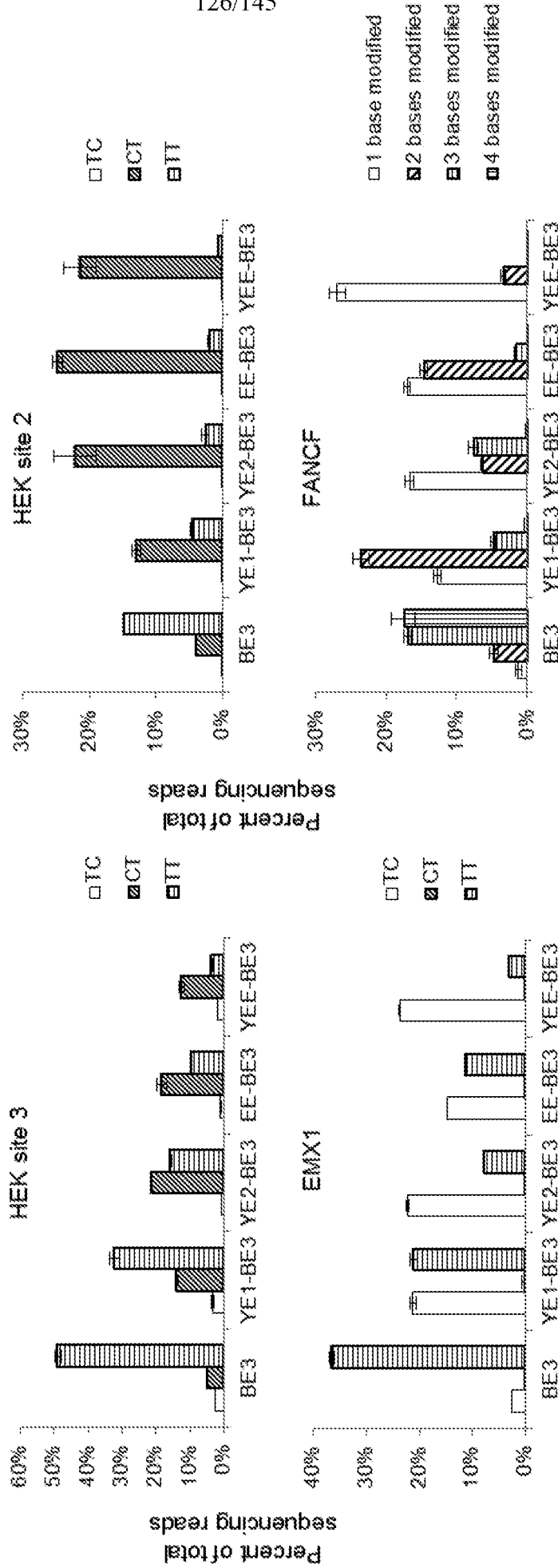


FIGURE 93C

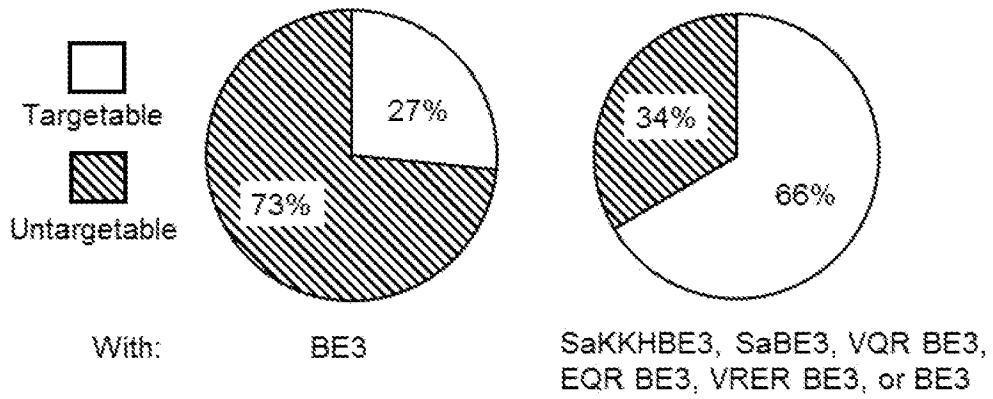


FIGURE 94A

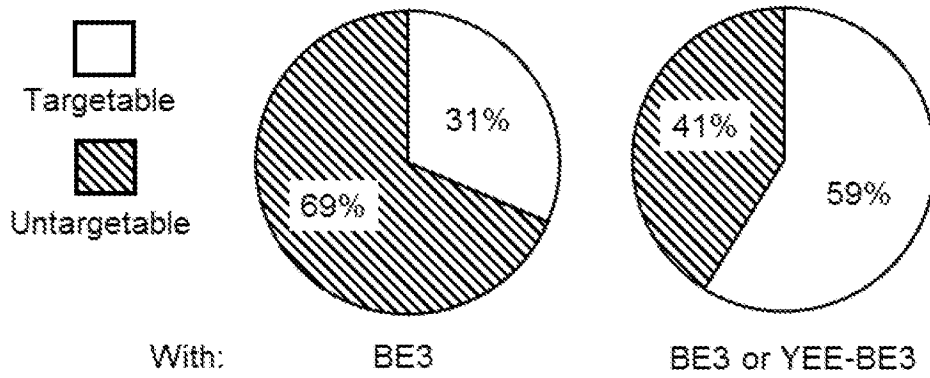


FIGURE 94B

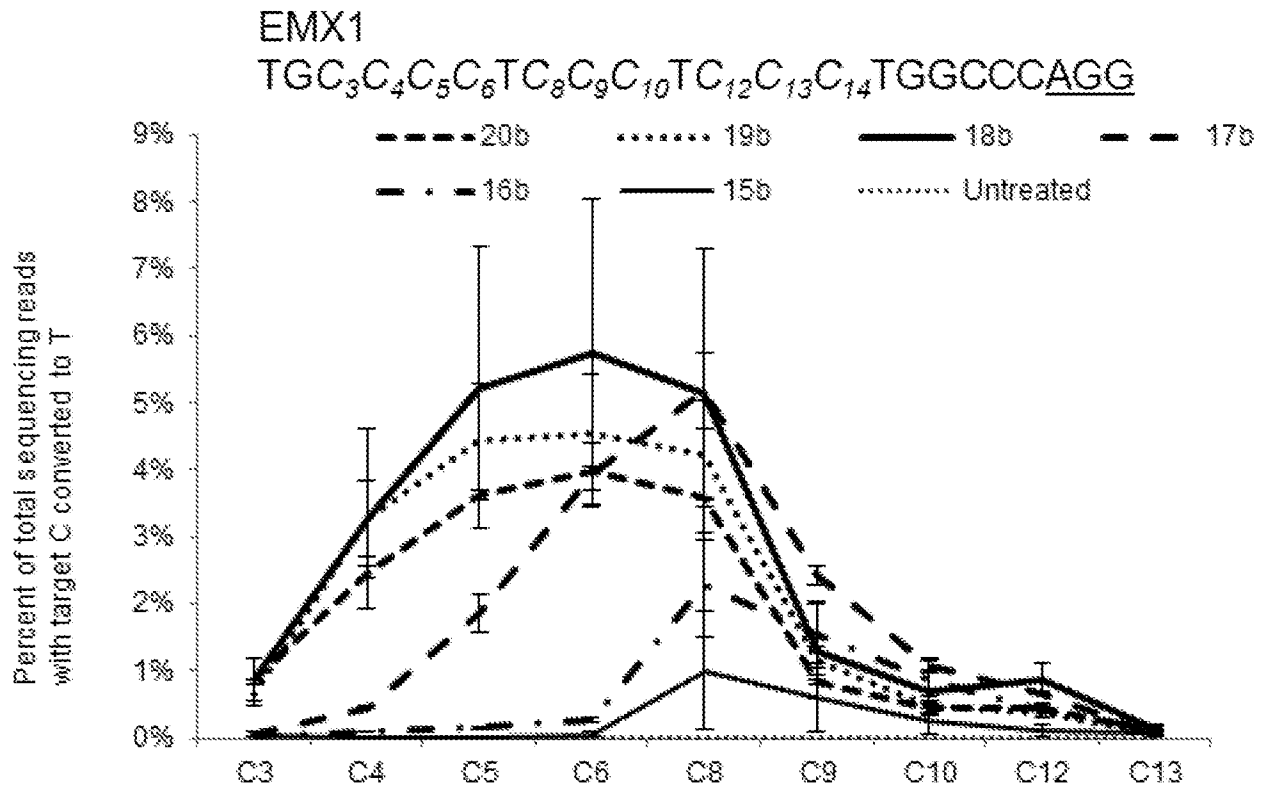
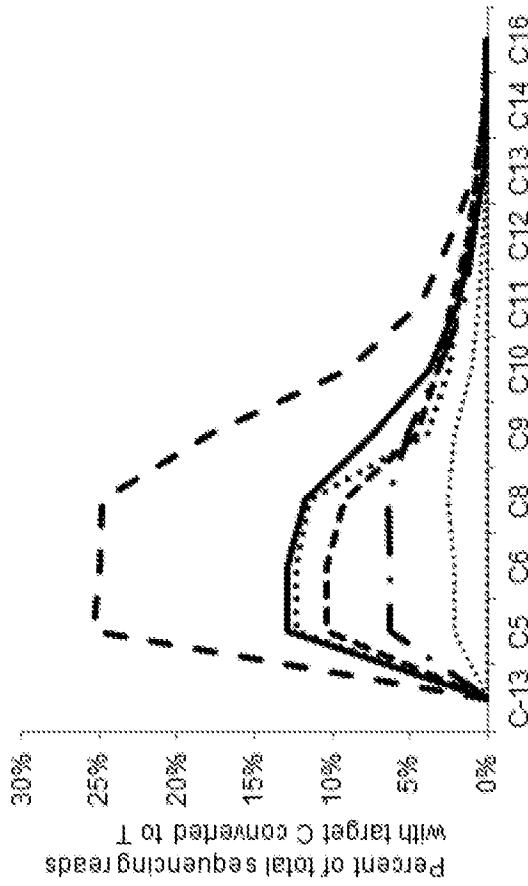


FIGURE 95A

HEK site 3
TTTTTC₅C₆TC₈C₉C₁₀C₁₁C₁₂C₁₃C₁₄AC₁₆AGGTAGA



HEK site 4
GC₂C₃C₄C₅C₆C₇C₈TC₁₀AAAGAGAGGGTGG

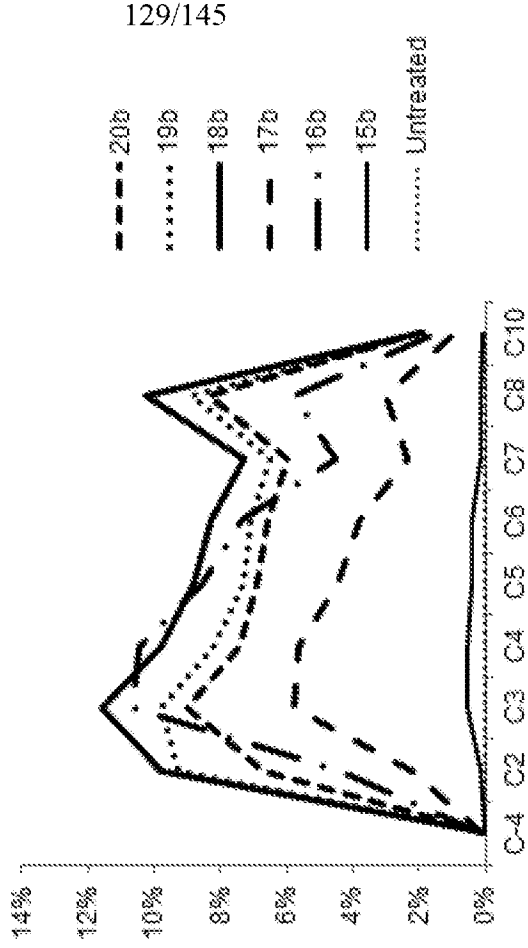


FIGURE 95B

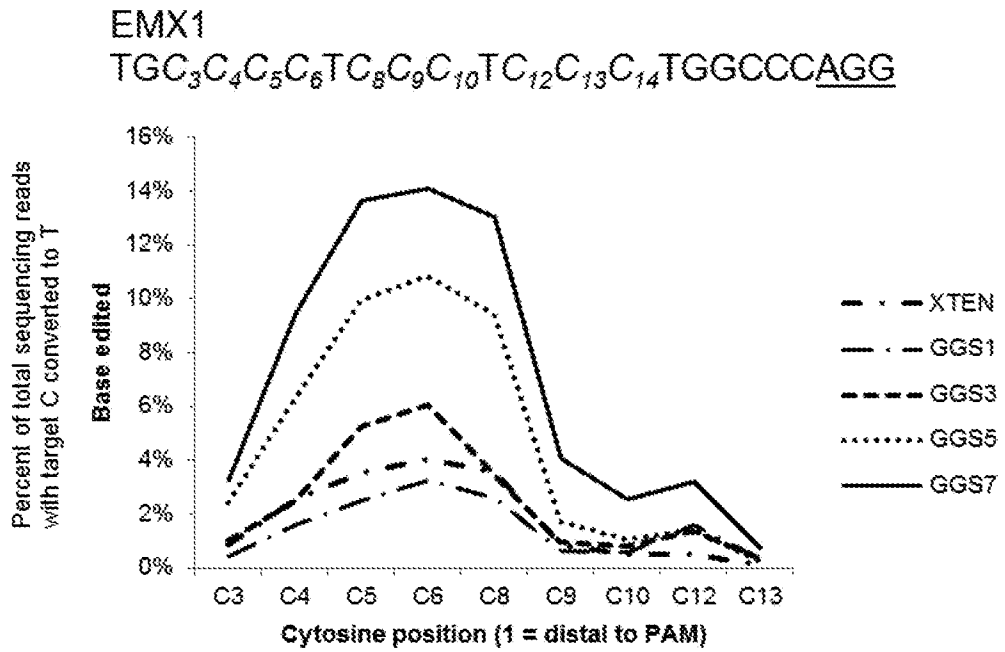


FIGURE 96

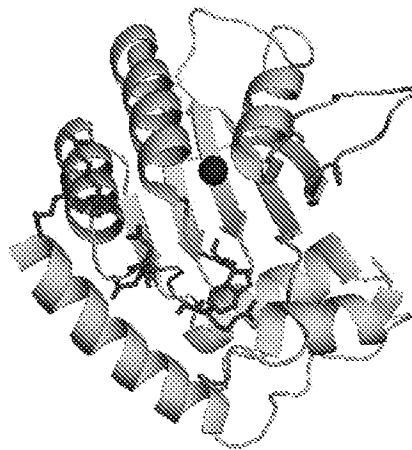


FIGURE 97A

APOBEC1 mutation	APOBEC3G mutation	Reference
R126A	R320A	#9,10
R126E	R320E	#9,10
W90A	W285A	#9,10
W90Y	W285Y	This work
R132E	R326E	This work

FIGURE 97B

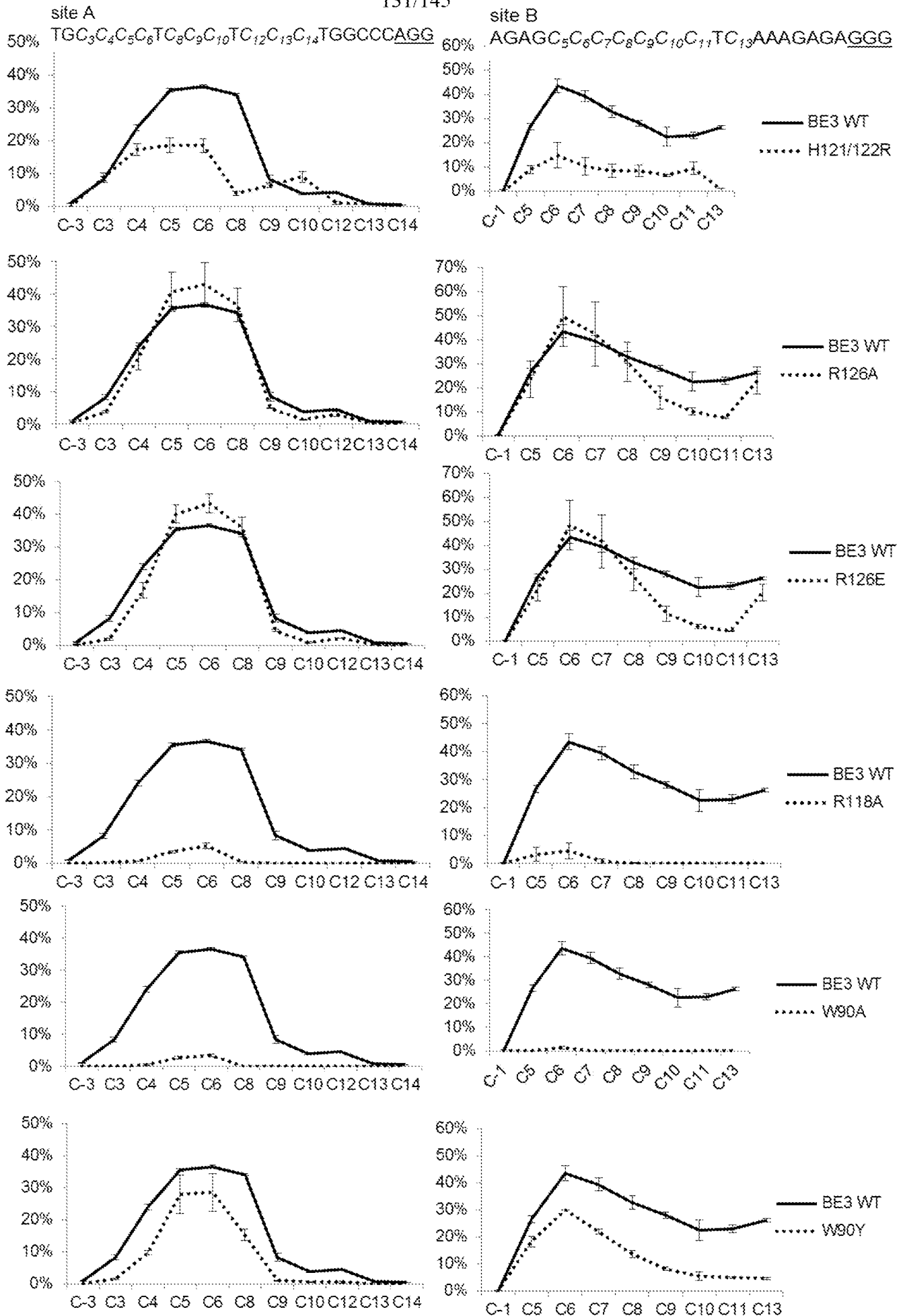
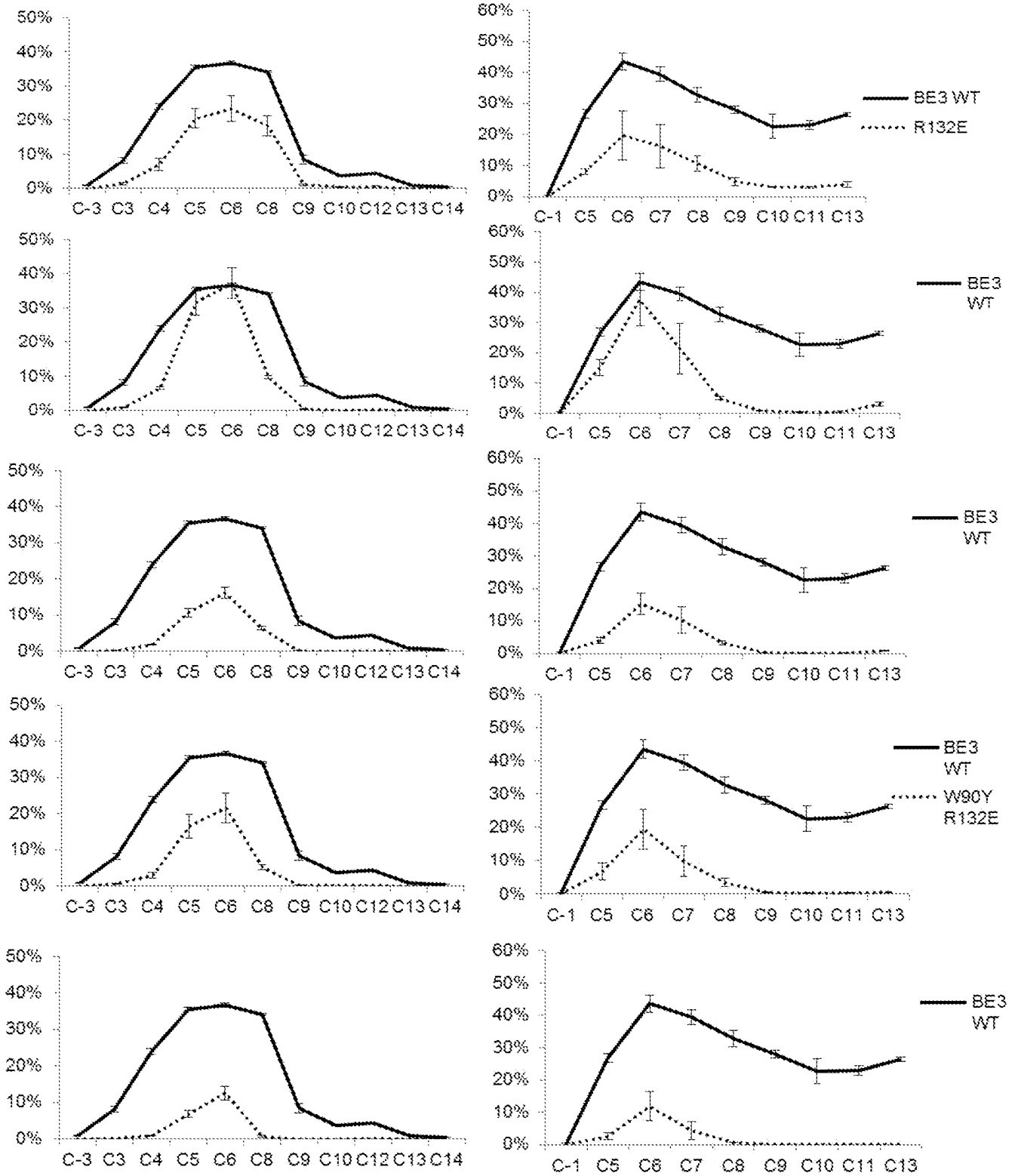


FIGURE 97C
SUBSTITUTE SHEET (RULE 26)

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Editing window width

	Site A	Site B		Site A	Site B
BE3 WT	5.0	6.1	W90Y	3.8	4.9
H121/122R	4.2	7.4	R132E	4.0	3.0
R126A	4.4	3.4	W90Y R126E	2.9	3.0
R126E	4.2	3.1	R126E R132E	2.9	3.0
R118A	2.4	3.6	W90Y R132E	2.7	2.8
W90A	2.5	1.1	W90Y R126E R132E	2.1	1.4

FIGURE 97C (CONTINUED)
SUBSTITUTE SHEET (RULE 26)

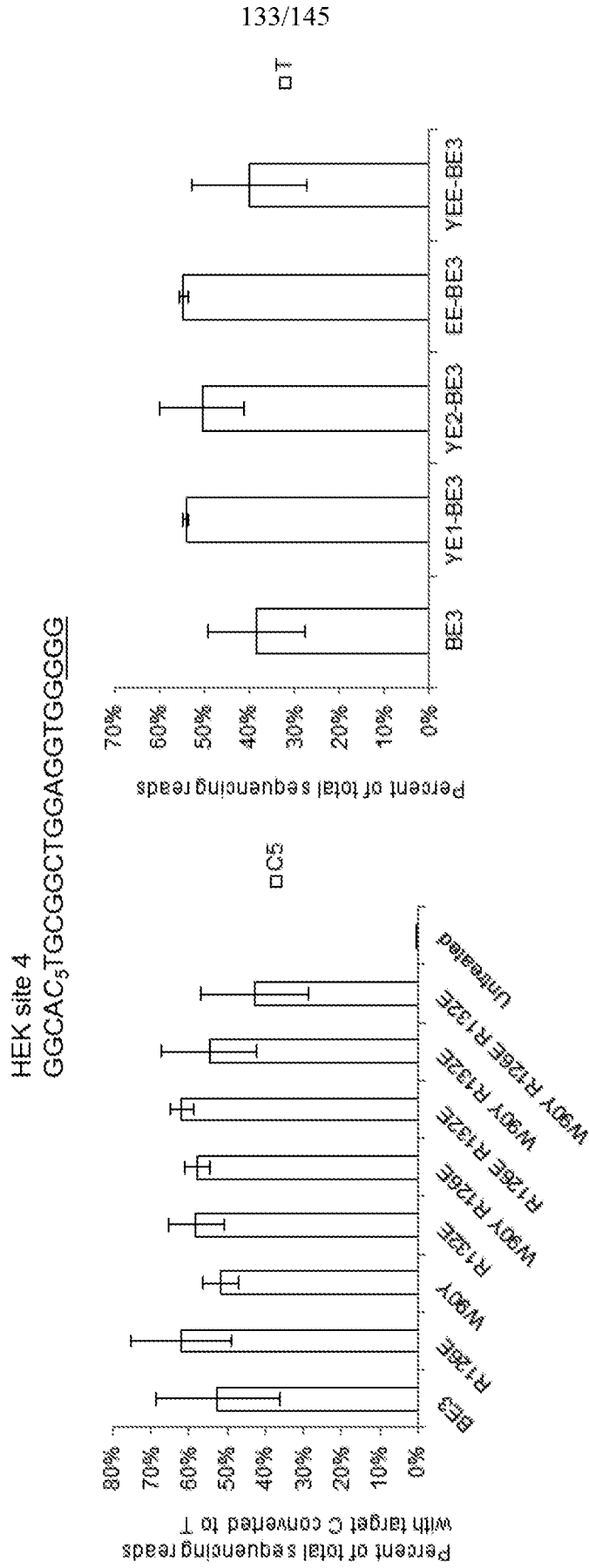


FIGURE 98

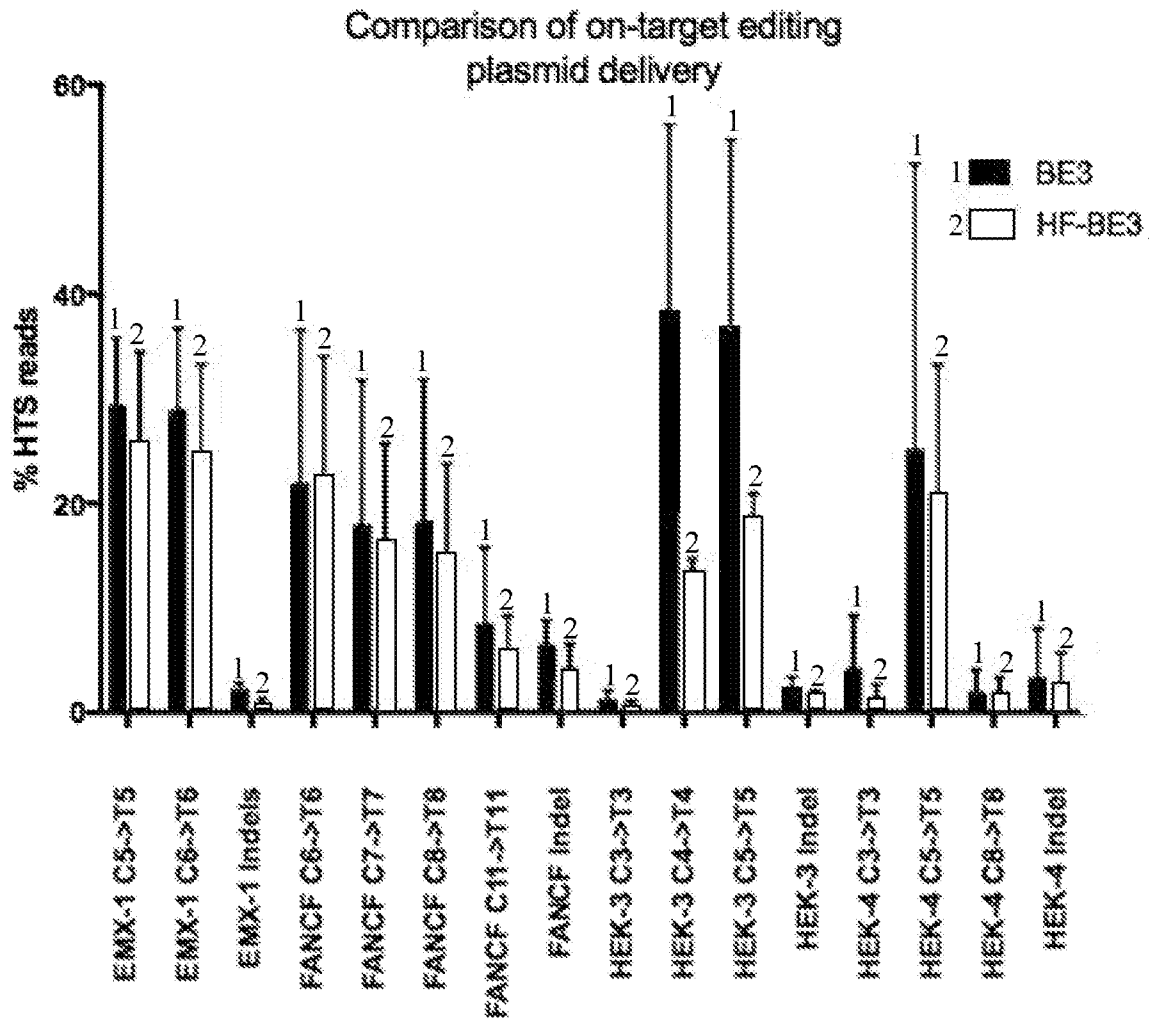


FIGURE 99

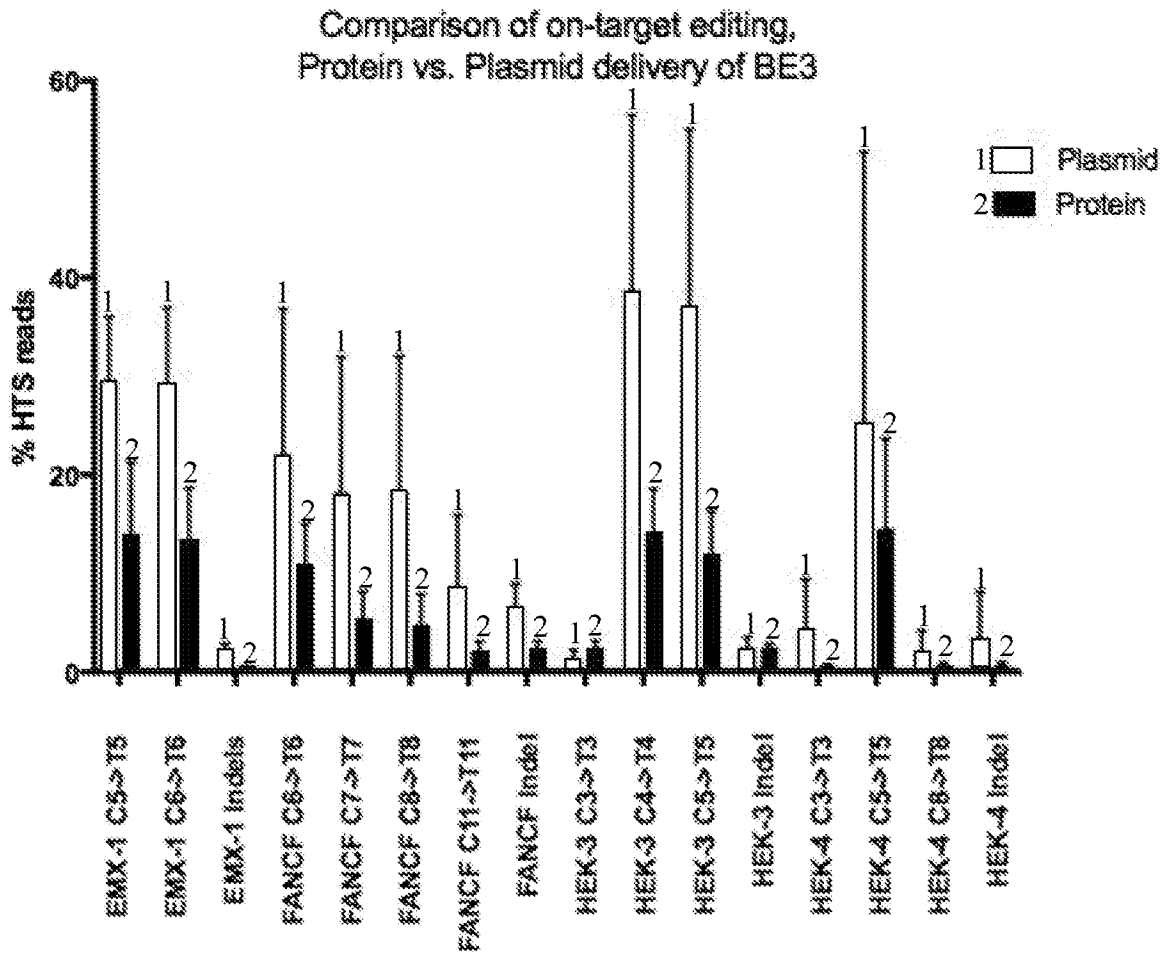


FIGURE 100

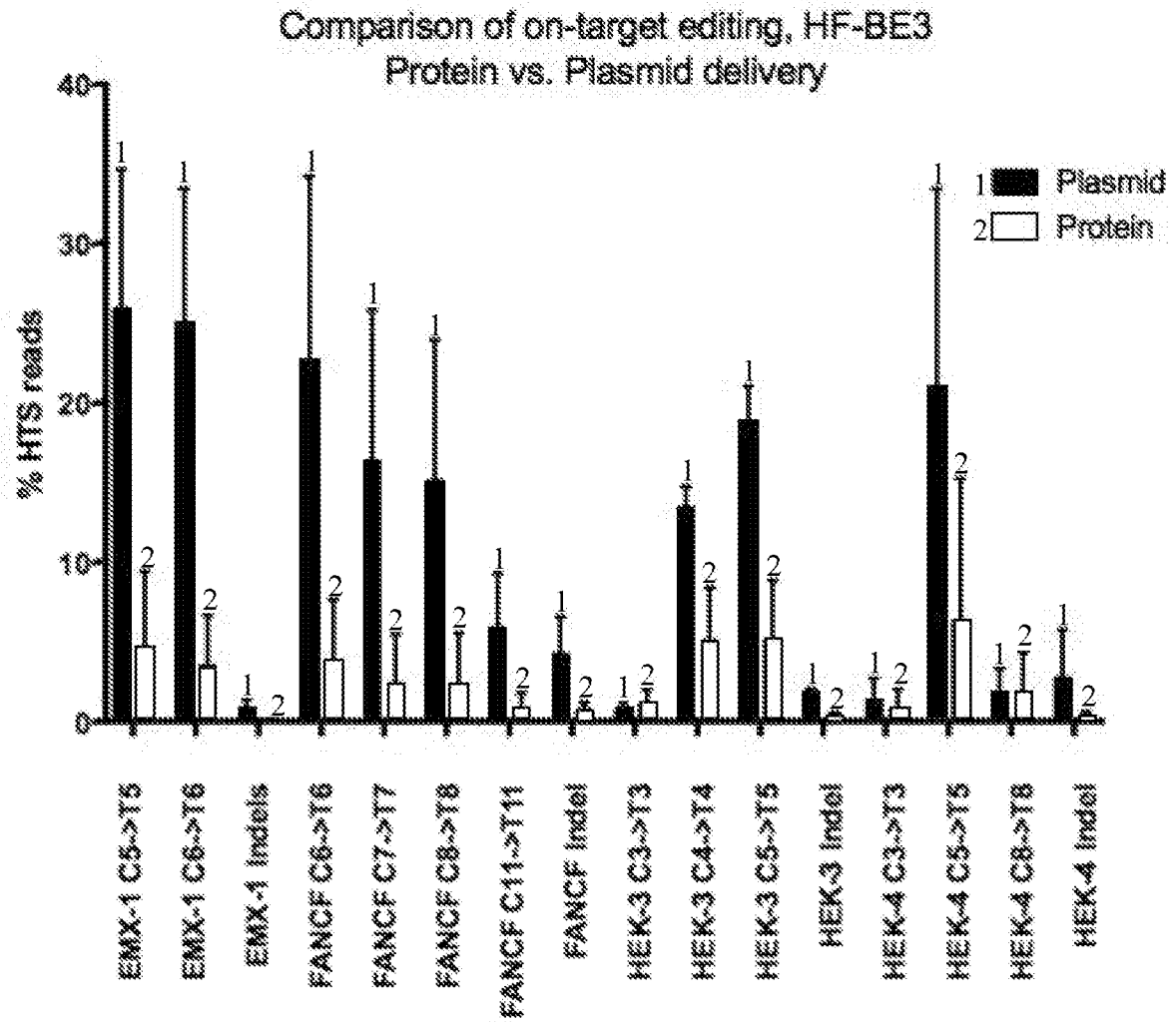


FIGURE 101

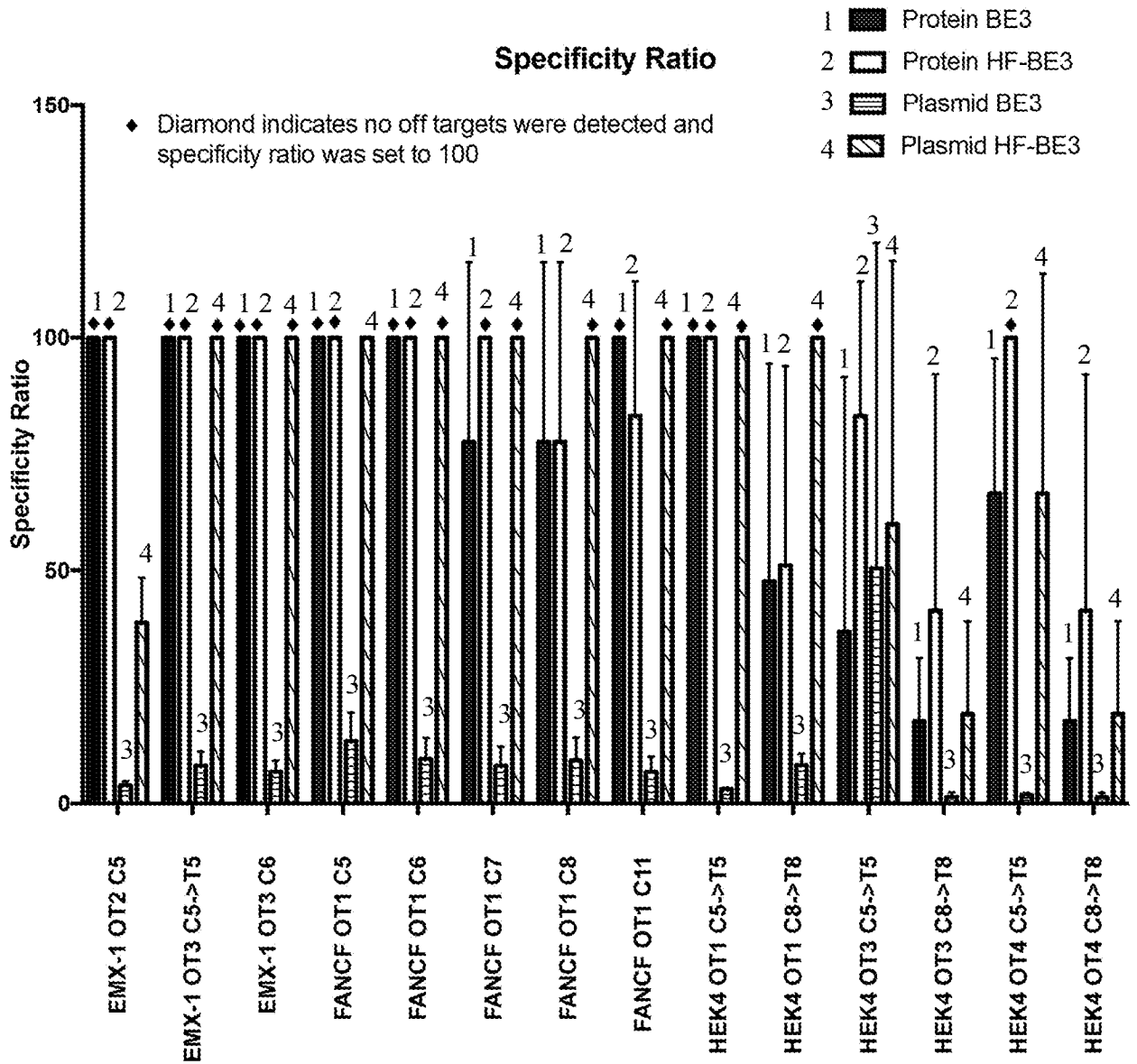


FIGURE 102

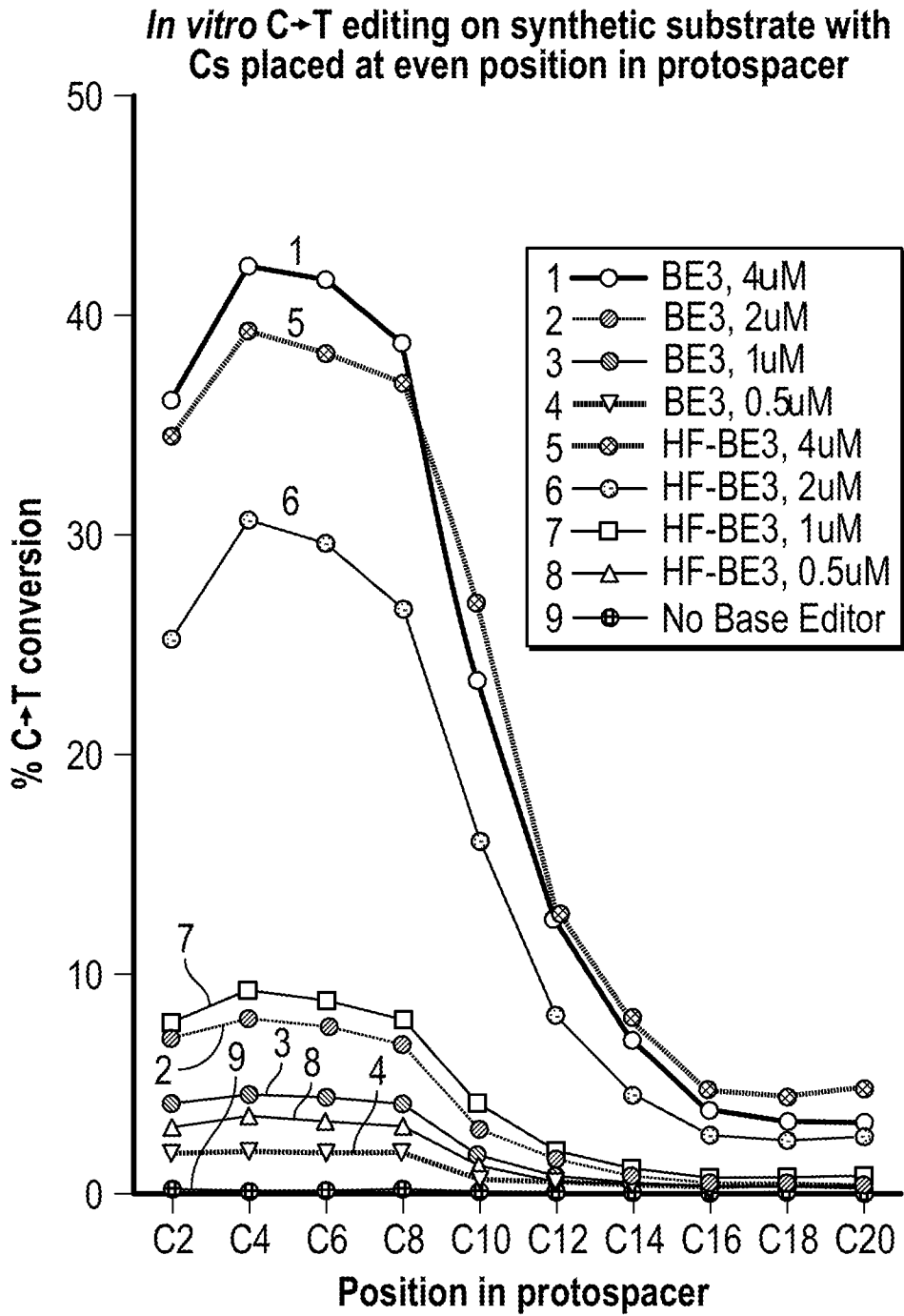


FIGURE 103

***In vitro* C→T editing on synthetic substrate with Cs placed at odd positions in protospacer**

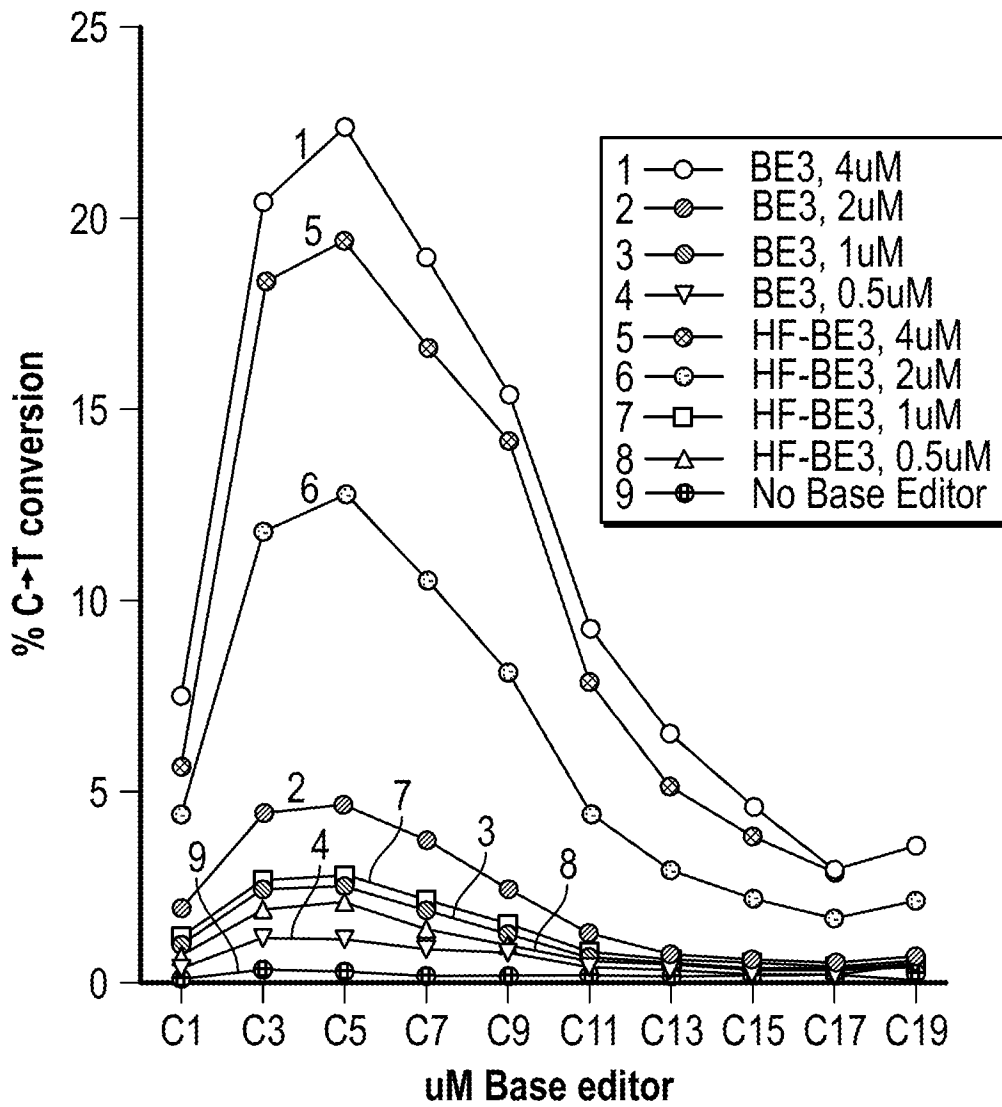


FIGURE 104

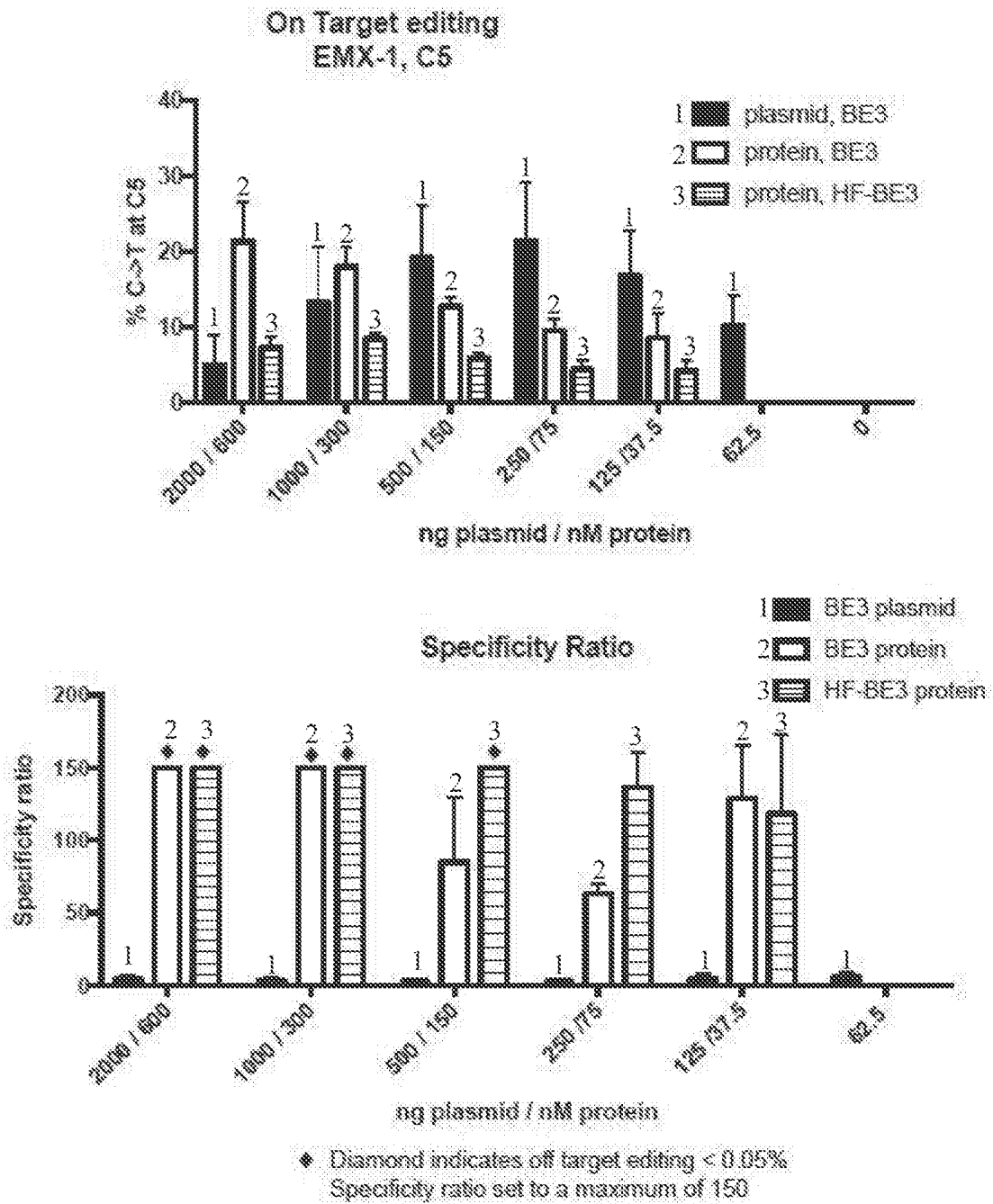


FIGURE 105

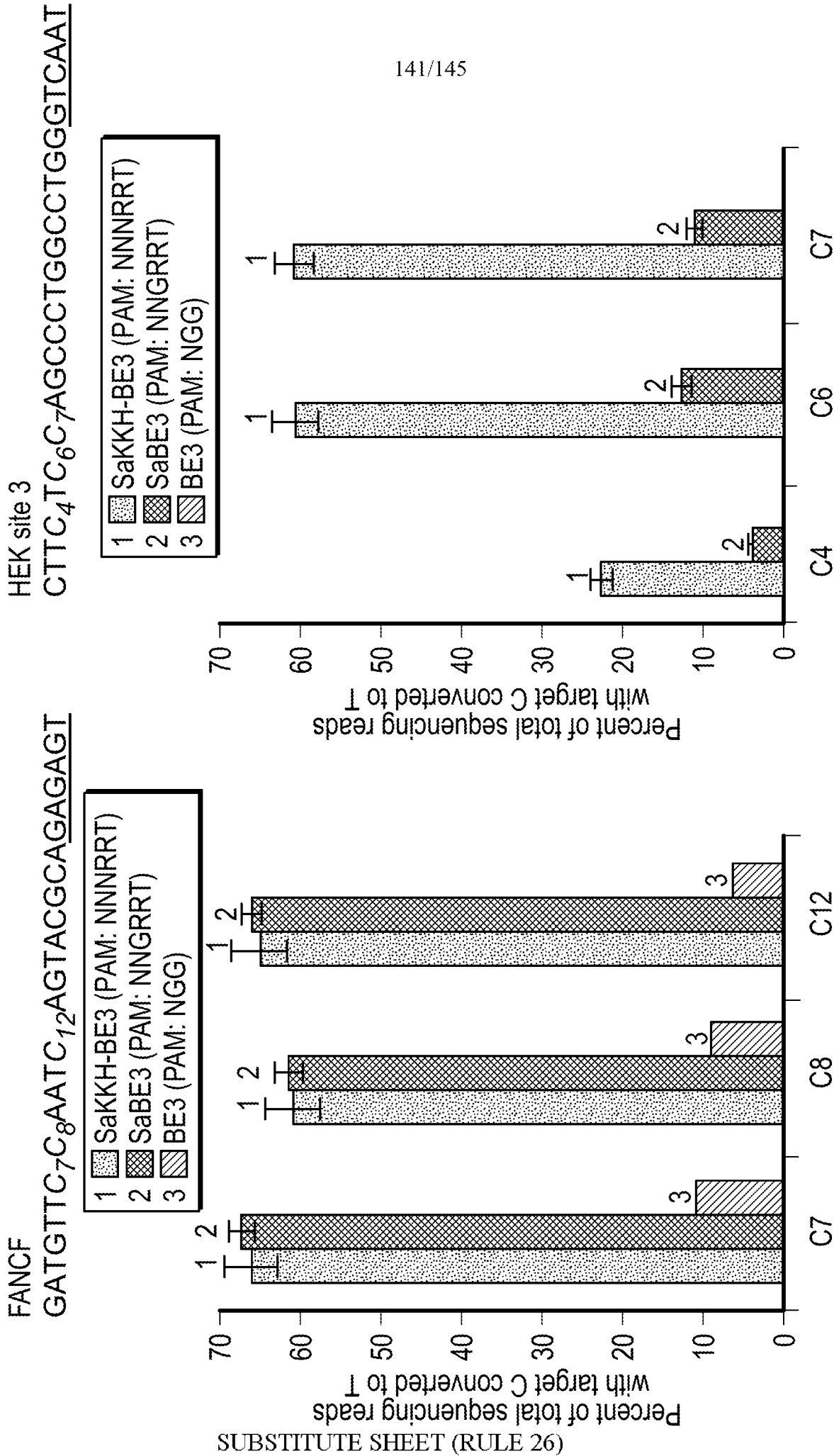
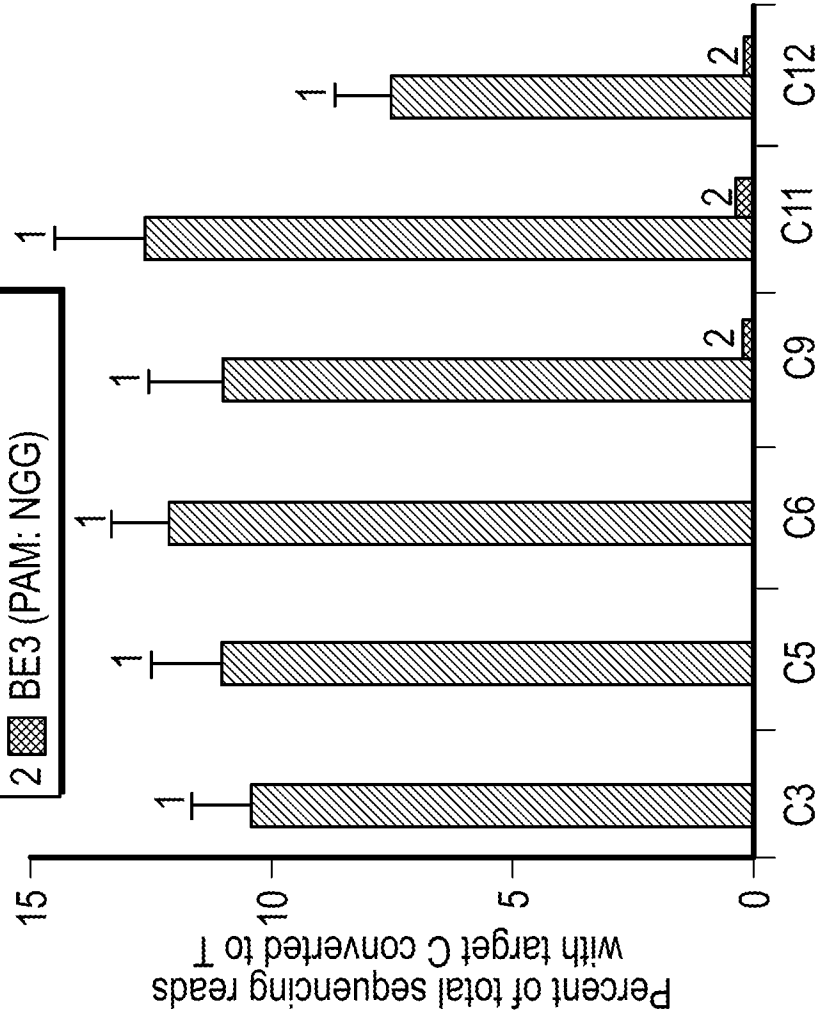


FIGURE 106A

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FANCF
 GTC₃GC₅C₆GTC₉TC₁₁C₁₂AAGGTGAAAGCGG

1 VRER-BE3 (PAM: NGCG)
 2 BE3 (PAM: NGG)



FANCF
 GCCGTC₆TC₈C₉AAGGTGAAAGCGGAA

1 VQR-BE3 (PAM: NGA)
 2 BE3 (PAM: NGG)

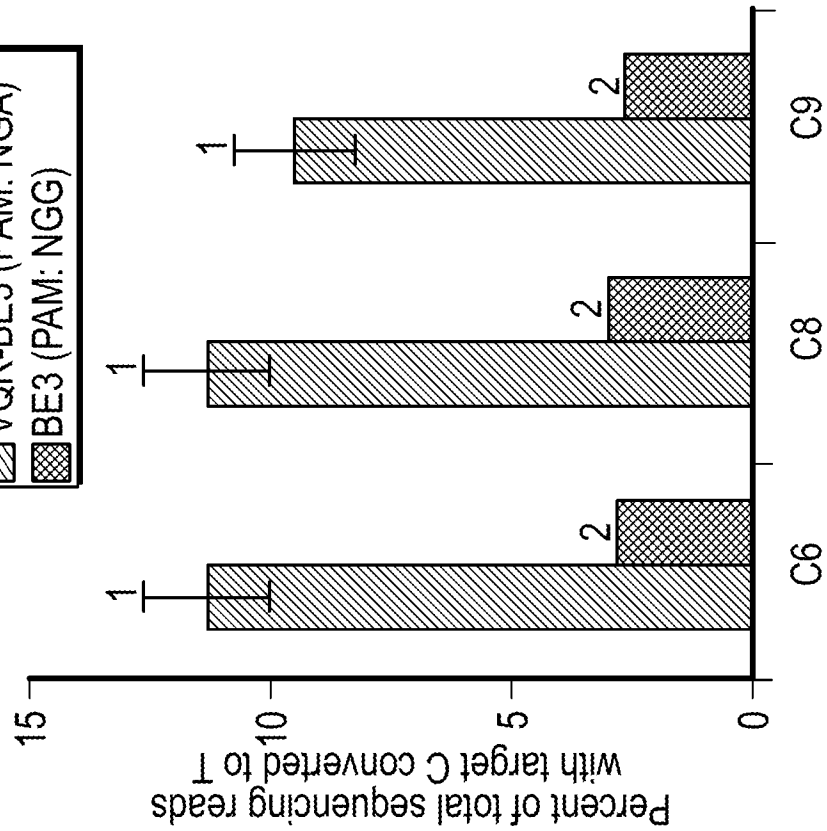


FIGURE 106B

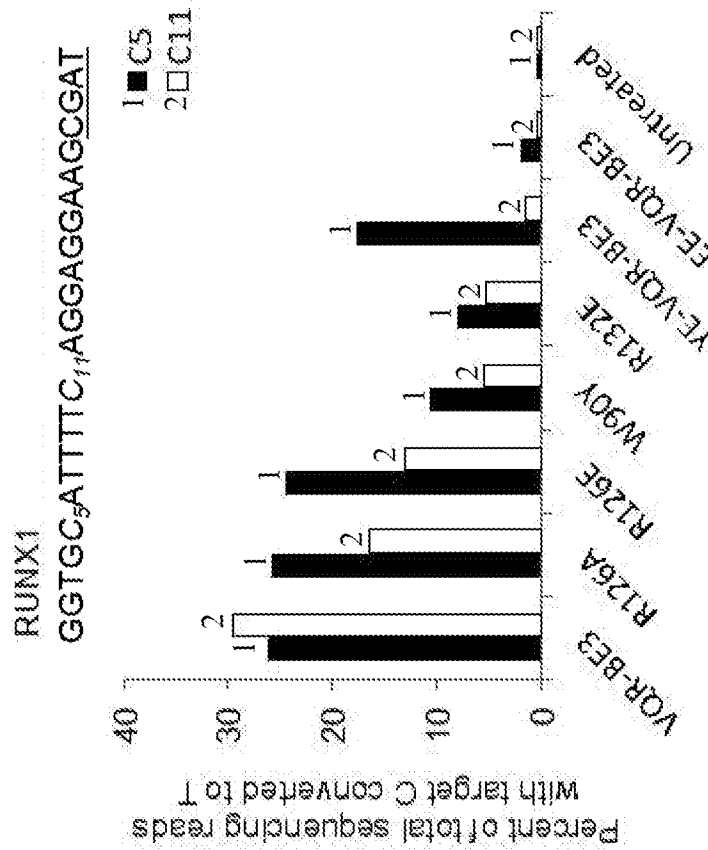
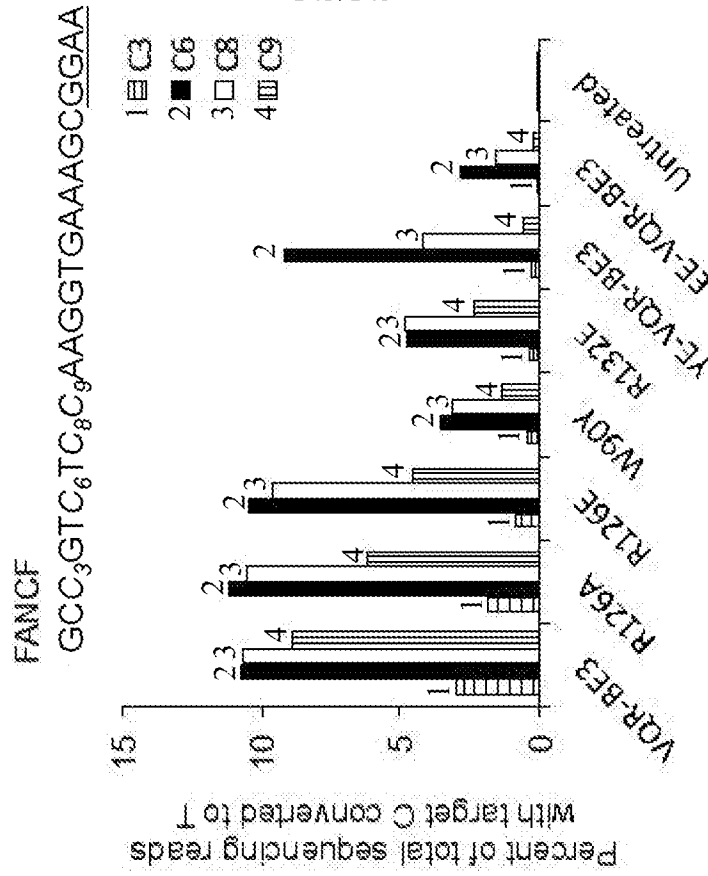


FIGURE 107A

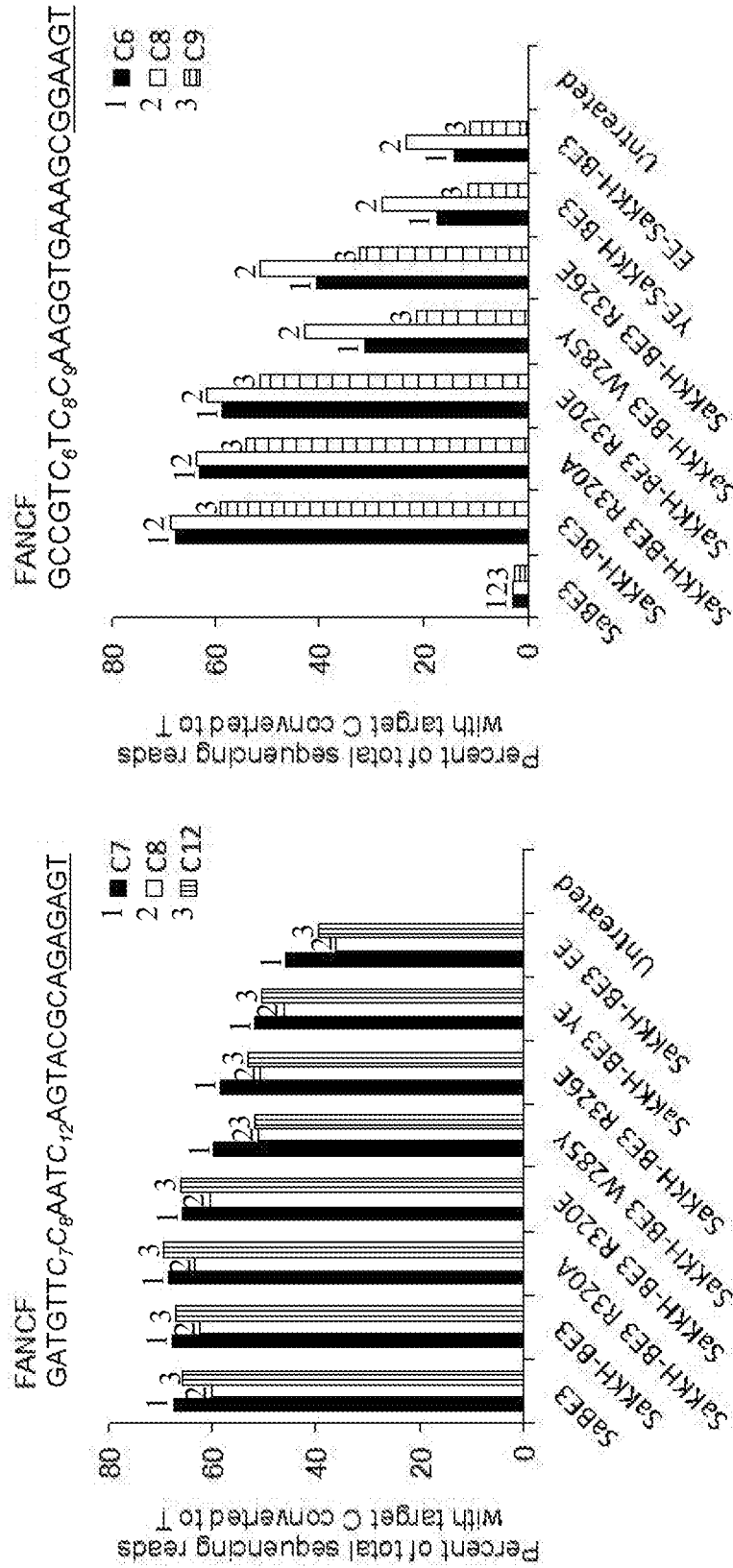


FIGURE 107B

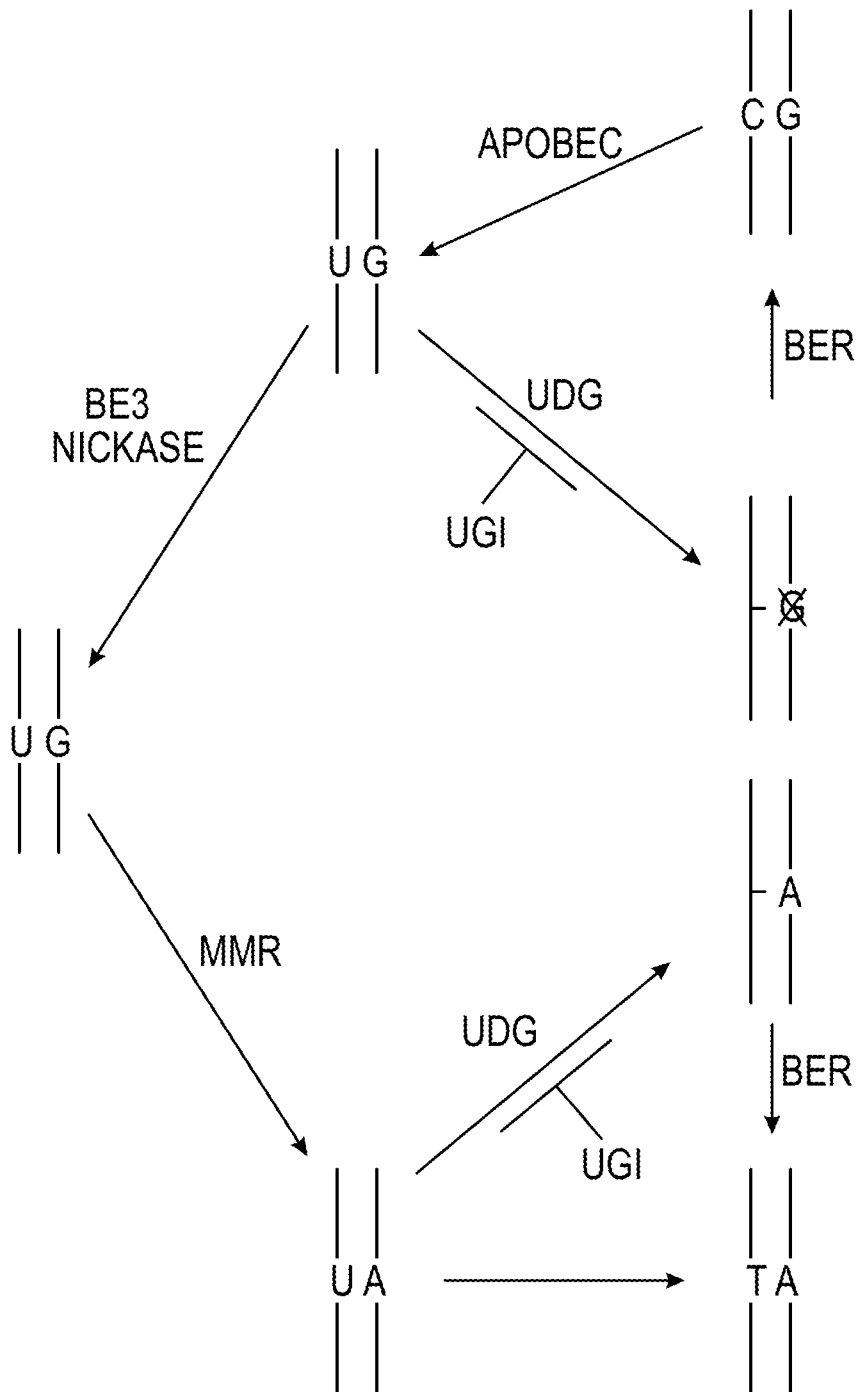


FIGURE 108