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ganon: precise metagenomics classification against large and up-to-date sets of reference sequences

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Abstract

The exponential growth of assembled genome sequences greatly benefits metagenomics 12 studies, providing a broader catalog of reference organisms on a variety of environments. 13 However, currently available methods struggle to manage the increasing amount of sequences 14 and their frequent updates. Indexing the current RefSeq is no longer possible on standard 15 infrastructures and it can take days and hundreds of GB of memory on large servers. Few 16 methods address these issues thus far, and even though many can theoretically handle large 17 amounts of references, time/memory requirements are prohibitive in practice. As a result, 18 many studies that require sequence classification use the available tools in conjunction with 19 often outdated and almost never truly up-to-date indices. This also means that the taxonomic 20 composition of the reference database is not being adjusted based on the study performed. 21 These factors can lead to unnecessary performance problems in the sequence classification. 22 Motivated by those limitations we created ganon, a k-mer based read classification tool that 23 uses Interleaved Bloom Filters in conjunction with a taxonomic clustering and a k-mer count-24 ing/filtering scheme. Ganon provides an efficient method for indexing references, keeping them 25 updated. It requires less than 55 minutes to index the complete RefSeq of bacteria, archaea, 26 fungi and viruses. The tool can further keep these indices up-to-date in a fraction of the time 27 necessary to create them, allowing researchers to always work with the most recent references. 28 Ganon makes it possible to query against very large reference sets and therefore it classifies 29 significantly more reads and identifies more species than similar methods. When classifying a 30 high-complexity real dataset from the CAMI challenge against complete genomes from RefSeq. 31 ganon shows strongly increased precision while exhibiting equal or better sensitivity compared 32 with state-of-the-art tools. When classifying the same dataset against the complete RefSeq, 33 ganon improved the F1-Score by 65% at the genus level. Ganon supports taxonomy- and 34 assembly-level classification as well as multiple indices and hierarchical classification. The 35 software is open-source and available at: https://gitlab.com/rki bioinformatics/ganon 36

37 1 Introduction

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Reference- and taxonomy-based short read classification is a fundamental task in metagenomics.
Defining the origin of each read from an environmental sample, which can be done during [1] or
after sequencing, is usually the first step prior to abundance estimation, profiling and assembly.
Over the last years many tools have been specifically developed for this task [2, 3, 4, 5, 6] with
different strategies to achieve good performance classifying a large amount of short reads against
a predefined and static set of reference sequences. Many of those approaches are taxonomy-based
[7] and use this classification to better understand the composition of samples.

*PiroV@rki.de [†]RenardB@rki.de The amount of complete or draft genomic sequences in public repositories is rapidly growing (Figure 1) due to advances in genome sequencing, improvements in read quality, length and coverage and also better algorithms for genome assembly. In addition, many partial and complete genome sequences come directly from metagenome-assembled genomes [8, 9, 10], a technique that boosts the growth of public repositories. This considerable amount of references poses a sizeable challenge for current tools that, in general, are not designed to deal with such amounts of data [11]. They also increase the already high computational cost of assigning millions of short reads to taxonomic targets





Figure 1: Number of available sequences in NCBI repositories from June 2007 to December 2018 on a logarithmic scale. Microbial stands for Archaeal and Bacterial organisms and CG stands for Complete Genomes. RefSeq Microbial has an uninterrupted and linear growth on a logarithmic scale. Data collected from: https://ftp.ncbi.nlm.nih.gov/refseq/release/release-statistics/ and https://www.ncbi.nlm.nih.gov/genbank/statistics/

Figure 1 shows the amount of reference sequences available over the last 11 years in the Gen-53 Bank [12] and RefSeq [13] repositories from NCBI. The growth is exponential. RefSeq sequences 54 from Archaeal and Bacterial genomes are highlighted for being a commonly used reference set for 55 classification in metagenomics. Within an interval of two and a half years (from June 2015 to 56 December 2018) the RefSeq Microbial of Complete Genomes (CG) grew more than four times, 57 with 2.5 times more species represented in the most recent set (1529 to 3850). Looking at the same 58 data point (end of 2018), the complete RefSeq Microbial has >12 times base pairs and >5 times 59 species compared to the CG set. These data exemplify that databases are growing fast and the 60 variation among them is significant. These repositories are becoming too big to be analyzed by 61 standard hardware and if the observed growth continues, all this wealth of data will be constrained 62 to just a few groups with resources available to process them. 63

The choice of the data to perform reference-based classification is an important step and a known 64 issue in metagenomics [14]. As a rule of thumb, the more sequences the better the classification. 65 But even complete sets of sequences are not evenly distributed throughout the taxonomic tree, such 66 that different taxa are represented in different levels of quantity and quality. In addition, most 67 of the taxa are still unknown and do not have any genomic sequence or entry in the taxonomic 68 tree. This requires the tools to consistently remain up to date with the latest releases of public 69 repositories, a task that is not trivial when dealing with very large amounts of sequences. Most of 70 the tools lack the ability to update their own indices and databases, and currently many analyses 71 are performed with outdated resources. 72

For example, the RefSeq Microbial repository from the beginning of 2018 is 10% less taxonomic
diverse than it is today (mid. 2019). An even older RefSeq release from June 2015 lacks 27% of
today's taxonomic diversity. Further, a commonly used subset of RefSeq, the microbial complete

genomes, covers only 15% of the available diversity of the full repository (December 2018). As 76 an example, the latest release of kraken's [15] MiniKraken database (as of 18-Oct-2017) based 77 on complete bacterial, archaeal, and viral genomes, although helpful to obtain fast insights on 78 community composition, comprises only 11% of the total taxonomic diversity available on the 79 latest RefSeq release from January 4th 2019. Metagenomics analyses based on those releases 80 are prone to underperform and miss potential species of interest. However, the use of outdated 81 references or "pre-built" indices is still common practice [16]. Most methods are able to build 82 custom databases but unable to update them. Weekly or daily updates with the most recent data 83 are almost impossible given the time requirements to re-build those indices 84

The sequence classifiers MetaPhlAn [17] and Kaiju [18] created alternatives to cover most of the diversity contained in public sequence repositories by selecting a subset of marker genes and protein sequences, respectively. On one hand, those methods are very powerful, such that they provide fast and precise community profiles given their reduced index sizes. On the other hand, when analyzing whole genome sequences of complex environments, organisms with low abundance are easily missed due to their lack of complete genomic coverage. In addition, current methods using complete genome sequences struggle with the present amount of available data [11].

Given these limitations, we developed ganon, a new reference and taxonomy-based short read 92 classification tool for metagenomics. Ganon uses Interleaved Bloom Filters (IBF) [19] to represent 93 very large amounts of sequences into a searchable index. This enables the indexing of large sets of references (e.g. complete RefSeq) in faster time and with low memory consumption, consequently 95 improving read classification for whole metagenomics sequencing experiments. Ganon also provides 96 updatable indices, which can incorporate new released sequences in short time. The classification 97 method, which is based on the k-mer counting lemma and a progressive filtering step, improves 98 the precision of the classification without harming sensitivity when compared to state-of-the-art 99 tools. Ganon was developed in C++ using the SeqAn library [20] and Python. The code is open 100 source and freely available from: https://gitlab.com/rki bioinformatics/ganon 101

$_{102}$ 2 Methods

Ganon classifies reads against a set of reference sequences to find their exact or closest taxonomic 103 origin. The method can also work in a further specialized level (e.g. assembly). Clustering and 104 indexing steps are necessary before classification, where the reference sequences will be grouped into 105 taxonomic groups and processed into a searchable index. Ganon indices store all k-mers present 106 in the reference sequences into a specialized type of Bloom filter. Once the index is created, ganon 107 classifies the reads based on the k-mer counting lemma together with a post-filtering step providing 108 a unique or multiple classifications for each read. Multiple classifications are solved optionally with 109 the lowest common ancestor (LCA) algorithm [21]. The following sections will further explain each 110 of these steps in detail. 111

112 2.1 Indexing

Ganon indices are based on the k-mer content of the reference sequences, in other words, it uses 113 all possible substrings of length k of the given sequences. Instead of using standard methods for 114 k-mer storage, which can have high memory and space consumption when k is high (> 15), we 115 opted for Bloom filters [22], a space-efficient probabilistic data structure. Since the goal of the tool 116 is to classify sequences based on their taxonomic origin, multiple Bloom filters would be necessary 117 to represent each distinct group of sequences belonging to a certain taxonomic level (e.g. species). 118 This approach provides a straightforward but impractical solution since it requires classification 119 against multiple filters. This is solved by interleaving the Bloom filters, a technique previously 120 described for the DREAM-Yara tool [19] and also part of the SeqAn library [20]. TaxSBP is used 121 to separate sequences into taxonomic groups and to distribute them better into equal-sized clusters. 122

123 2.1.1 TaxSBP

TaxSBP [https://github.com/pirovc/taxsbp] uses the NCBI Taxonomy database [23] to generate clusters of sequences that are close together in the taxonomic tree. It does this based on an implementation of the approximation algorithm for the hierarchically structured bin packing problem [24]. As defined by Codenotti et al. this clustering method "[...] can be defined as the problem of

distributing hierarchically structured objects into different repositories in a way that the access to 128 subsets of related objects involves as few repositories as possible", where the objects are sequences 129 assigned to taxonomic nodes of the taxonomic tree. Sequences are clustered together into groups 130 limited by a maximum sequence length size of its components. Splitting sequences into smaller 131 chunks with overlapping ends is supported. TaxSBP supports one level of specialization after the 132 leaf nodes of the tree, making it possible to further cluster sequences by strain or assembly infor-133 mation that is not directly contained in the NCBI Taxonomy database (Figure 2 A). TaxSBP can 134 also pre-cluster members of a certain taxonomic level, preventing them to be split among clusters. 135 It can further generate clusters with exclusive ranks, which are guaranteed to be unique in their 136 cluster. The tool was developed alongside the distributed indices concept [19] and supports the 137 update of pre-generated clusters. Since TaxSBP uses the "pre-clustered" taxonomic tree informa-138 tion, the algorithm is very efficient and requires very few computational resources, thus having 139 potential use in many other bioinformatics applications. 140

141 2.1.2 IBF

A Bloom filter is a probabilistic data structure that comprises a bit vector and a set of hash functions. Each of the functions maps a key value (k-mer in our application) to one of the bit positions in the vector. Collisions in the vector are possible, meaning that distinct k-mers can be set to the same bit positions in the vector. Those overlaps can be avoided with a larger bit vector, thus reducing the probability of false positives.

An Interleaved Bloom Filter (IBF) is a combination of several (b) Bloom filters of the same size (n) with the same hash functions into one bit vector (Figure 2 D). Each *i*-th bit of every Bloom filter is interleaved, resulting in a final IBF of size b * n. Querying in this data structure is possible by retrieving the sub-bit vectors for every hash function and merging them with a logical AND operation, which will result in a final bit vector indicating the membership for the query, as depicted in Figure 2 in the DREAM-Yara manuscript by [19].

Aiming at the classification based on taxonomic levels (e.g. species, genus, ...) or assembly level, TaxSBP is set to cluster the input sequences into exclusive groups (Figure 2 B). Every group will contain only sequences belonging to the same taxon or assembly unit, but the same unit can be split into several groups. Groups are limited by a predefined threshold of the sum of the base pair length of its elements and sequences can be sliced into smaller pieces to better generate equal sized clusters.

Each of those clusters will correspond to a single Bloom filter that is interleaved in a final IBF 159 (Figure 2 C-D). Here a trade-off between the number of groups, their maximum size and the k-mer 160 content of each group is important. The false positive rate of a Bloom filter depends mainly on its 161 bit vector size and the number of inserted elements. In general, the more base pairs a particular 162 cluster has, the higher the number of distinct k-mers. This requires the Bloom filter to be bigger 163 in order to achieve low false positive rates when querying. In ganon indices, the group with the 164 most unique k-mers will define the size and the maximum false positive rate of the final IBF since 165 all groups have to be equal-sized by definition. Thus the previous clustering step is crucial to 166 achieve a good trade-off between the number of groups, their sizes and k-mer content. The lower 167 the taxonomic level, the more fragmented the clusters. For example: if a reference set has 2000 168 species groups, there will be at least the same number of clusters when building at the species 169 level. The higher the taxonomic level, the fewer the number of clusters, since they can be grouped 170 together, thereby producing smaller filters. This trade-off and parameterization is automatically 171 calculated by ganon, with a single option to define the maximum memory available to build an 172 index. 173

The IBF has an inherent capability of updating since it is fragmented into many sub-parts. Adding new sequences to a previously generated IBF is as easy as setting the bit positions of the k-mers from the new sequences to their known clusters or appending new clusters to the existing filter. To remove sequences from the IBF, all bit positions of the updated cluster are set to zero and the cluster is re-created from the updated content.

The IBF is the main data structure for ganon indices to perform alignment-free classification while DREAM-Yara, the tool that originally proposed the IBF, is a read mapper that uses the same data structure to filter reads to further perform distributed alignment. At the end of the building process, the ganon index will consist of an IBF based on a maximum classification level chosen (taxonomic rank or assembly) and auxiliary files for the classification step.



Figure 2: Classification levels, taxonomic distribution and Interleaved Bloom Filter A) Empty circles are inner nodes of the tree; circles marked with an "x" are leaf nodes (also referenced in this manuscript as "taxid" nodes); full lines represent taxonomic relations, dotted lines represent the extension of the taxonomic classification to the assembly and sequence level. Species+ represents all taxonomic groups that are more specific than species with species in the lineage (e.g. subspecies, species group, no rank). B) A toy example of sequences clustered by species into equal-sized groups, performed by TaxSBP C) Sequences are fragmented into k-mers and with a given number of hash functions, those k-mers are inserted into equal-sized bit-vectors (Bloom Filters) D) The Interleaved Bloom Filter, representing the previously generated bit-vectors with each bit interleaved.

184 2.2 Classifying

The read classification is based on the well-studied k-mer counting lemma [25, 26]. All k-mers from given reads are looked up on the indices previously generated. If a minimum number of matches between the read and the reference is achieved, a read is considered classified. Based on incremental error rates, multiple classifications for each read are filtered out and only the best ones are selected. When the filtering cannot define a single origin for a read, an optional LCA step is applied to join multiple matching reads into their lowest common ancestor node in the taxonomic tree.

¹⁹² 2.2.1 K-mer counting lemma

The k-mer counting lemma can be defined as the minimum number of k-mer sequences of a read that should match against reference k-mers in order to be considered present in a set with a certain number of errors allowed. Given a read, R, with length l, the number of possible k-mers with length k in this read can be defined as:

$$kmers_R = l_R - k + 1 \tag{1}$$

An approximate occurrence of R in a set of references has to share at least

$$kcount_R = kmers_R - k \cdot e \tag{2}$$

k-mers, where e is the maximum number of errors/mismatches allowed.

199 2.2.2 Filtering

A read can be assigned to multiple references with different error rates, thus a filtering step is necessary to decrease the number of false assignments. The applied k-mer counting lemma provides k-mer counts for each read against the reference sequences. From this count it is possible to estimate the number of mismatches a read has. For example: for k = 19 and length = 100, a read with 50 19-mers matching a certain reference will optimally have 2 mismatches. This calculation can be achieved by solving the Equation 2 equation for e.

Assuming that reads with fewer mismatches have a higher chance of being correct, the following 206 filtering is applied: first, only matches against references with no mismatches are kept (all k-mers 207 matching). If there are no such assignments, matches with only 1 error are kept. If there are none, 208 matches with only 2 errors are kept and so on up to the maximum number of errors allowed (e in 209 Equation 2). Similar filtration methods (also known as mapping by strata) were previously used in 210 read mappers such as Yara [19]. If a read is classified in several references within the same range 211 of errors, they are all reported since it is not possible to define which one has a higher chance of 212 being correct based on the k-mer count information. Given our clustering approach, some groups 21 3 can share the same identification target (e.g. one species was split in two or more clusters due to 214 a large amount of sequences). These cases are treated specially by reporting only the match with 215 more k-mer similarities since they belong to the same classification group. 216

Ganon also provides a way to further filter the unique classifications with a different error rate for reads that matched against just one reference group. This filter will be applied after the standard filtration and will re-classify a low scored read to its parent taxonomic level if it scores below a certain threshold. This can be applied for filtering at low levels (e.g. assembly) since the classification in those levels should be more precise with less mismatches. This feature is also useful to avoid classifications that only happen due to a lack of related genomes (e.g. a low score match on the only representative species of a lineage).

In summary, ganon indices represent groups of reference sequences clustered by taxonomy or 224 assembly group. All k-mers from the reads are extracted and compared against an index by 225 applying the k-mer counting lemma to select candidates. This is done based on a user defined 226 optimal number of errors. All matches within the error rate are filtered and one or more matches 227 are reported. At the end, an optional LCA method can be applied for reads with multiple matches 228 with a more conservative and less precise taxonomic classification, thus resulting in one match for 229 each read. Additionally, ganon supports classification based on multiple indices in a user-defined 230 hierarchy, with independent error rates for each index (Supplementary Material 1 - Section 2.4). 231

232 **3** Results

We evaluated ganon against a set of well-established methods from recent benchmarks [3, 4, 6] that 233 performs short read classification and supports indexing of large sets of reference sequences. The 234 aim here is to compare in equal conditions the methods regarding input data, reference sequences 235 and taxonomy. We compared ganon against kraken [15], one of the most used k-mer based methods 236 for metagenomics short read classification. We also included krakenuniq [27], which uses the basic 237 kraken algorithm and also allows classification on more specific levels after taxonomic assignments 238 (e.g. up to assembly or sequence level). We further compare the results against centrifuge [28] that 239 uses the Burrows-Wheeler transform (BWT) and the Full-text index in Minute space (FM-)index 240 for indexing and aims to reduce the index size by compressing similar sequences together. Clark 241 [29], another k-mer approach that uses common k-mers between reference sequences was also eval-24.2 uated. Diamond [30] an alignment tool for short DNA sequencing reads against protein reference 24 3 databases was also included. Here we consider only the direct read classification capabilities of 244 the tools. Further functionalities such as the estimation of a presence of a certain organism or 245 abundance estimation were not covered. All steps performed in the evaluation were compiled in a 246 benchmark pipeline (version 1.0.0) available from https://github.com/pirovc/ganon benchmark. 247

Ganon and the other evaluated tools are reference-based, meaning all classifications are made 248 based on previously generated sequence data. The choice of the underlying database is therefore 249 crucial. We use the same sequences and taxonomic database version for all tools when creating 250 their indices to guarantee a fair comparison. The NCBI RefSeq repository was the chosen source 251 of sequences since it provides a curated and very extensive set of references. Two subsets of RefSeq 252 were extracted: a set of only complete genomes from the groups Archaea, Bacteria, Fungi and Viral 253 (RefSeq-CG) and a complete set of all genomes from the same groups (RefSeq-ALL) both dating 254 from 19-December-2018 (Table 1). Genomic DNA data was obtained for all tools. Protein sequence 255 data from annotated genome assemblies was obtained for diamond (Supplementary Material 1 -256 Table 5). Taxonomic information was obtained at the same dates as the sequences. Additionally, 257

	Base pairs	# assemblies	# sequences
RefSeq-OLD	$9,\!632,\!441,\!987$	$3,\!042$	5,242
RefSeq-CG	$46,\!986,\!899,\!184$	$19,\!623$	33,029
$\operatorname{RefSeq-ALL}$	$587,\!607,\!072,\!429$	147,713	$15,\!201,\!684$

an old set of only Bacterial complete genomes from 02-June-2015 (RefSeq-OLD) was included to evaluate the tool's performance on an outdated and less diverse set of references.

Table 1: Genomic DNA of reference sequences used for evaluations. Protein data information can be found in the Supplementary Material 1 - Table 5. Detailed information of each dataset can be found in the Supplementary Material 1 - Section 1.5.1. Data was downloaded using https://github.com/pirovc/genome_updater

The selected reference sets contain over-represented taxonomic groups with several assemblies 260 for a single species. For example: the *Escherichia coli* species group is represented by 634 assem-261 blies, accounting for almost 7% of all base pairs in the RefSeq-CG. This is even more pronounced 262 on RefSeq-ALL, with 13,259 E. Coli assemblies representing more than 11% of the base pairs in 263 the whole set. In RefSeq-CG, the 92 most over-represented species have as many base pairs as 264 the remaining 11,372 species. In RefSeq-ALL this ratio is 14 to 29,047 (Supplementary Material 265 1 - Figure 1). This unbalanced distribution of references may not only bias analysis but also in-266 troduces redundancy to the set when aiming classification at taxonomic levels. Therefore, when 267 not classifying at assembly level, we removed over-represented assemblies from our reference set, 268 keeping only the 3 biggest assemblies of each taxonomic group (Table 2, Supplementary Material 269 1 - Figure 2). 270

	Base pairs	# species	# leaf taxids	# assemblies	# sequences
RefSeq-CG-top-3 RefSeq-ALL-top-3	$2.9e10~(62\%) \ 2.1e11~(36\%)$	$11,464\ (100\%)\ 29,061\ (100\%)$	$14,071\ (100\%)\ 51,292\ (100\%)$	$15,171 \ (77\%) \ 56,805 \ (38\%)$	$\begin{array}{r} 24,290 \ (74\%) \\ 4,400,402 \ (29\%) \end{array}$

Table 2: Reference sequences after over-representation filtering. Percentages in brackets show the amount of data left compared to the original set (Table 1). Protein data information can be found in the Supplementary Material 1 - Table 5.

For classification we used reads from the first CAMI Challenge [6]. Sets of simulated and real 271 datasets mimicking commonly used environments and settings were obtained, representing multiple 272 closely related strains, plasmid and viral sequences. These samples were divided into 3 categories: 273 low, medium and high complexity with increasing number of organisms and different sequencing 274 profiles providing a well-produced and challenging dataset to analyze. The simulated reads were 275 generated based on public data (NCBI, SILVA46) and an exact ground truth assignment is provided 276 for each read down to sequence level. The real dataset was obtained from newly sequenced genomes 277 of 700 microbial isolates and 600 circular elements and a ground truth is provided at taxonomic 278 levels. Here we used one high complexity sample from both categories to perform evaluations and 279 benchmark the tools (Supplementary Material 1 - Table 6). 280

The classification results were evaluated in terms of sensitivity and precision in two different 281 ways: cumulative- and rank-based. Details on their differences can be found in the Supplementary 282 Material 1 - Section 1.7. In short, the cumulative-based evaluation will compare how well tools 283 perform up to a certain taxonomic level, considering only the taxon of their final classification 284 level. The rank-based evaluation considers the full lineage of each classification. For example: 285 in a cumulative-based evaluation, values of sensitivity and precision at family level will account 286 cumulatively for all sequences classified at subsequent taxonomic levels (genus, species, species+) 287 up to (and including) the family level. In a rank-based evaluation, family level sensitivity and 288 precision values are calculated based on the family assignment from the lineage of the classified 289 sequences. The cumulative-based evaluation provides a better way to compare tools and their 290 ability to correctly classify sequences to their targets. The rank-based approach will better compare 291 how tools perform at a specific taxonomic level. In this work we will use both methodologies 292 to compare the results of the evaluated methods. Additionally, we evaluated all scenarios with 293 AMBER [31], an independent tool for assessment of metagenome binners with a similar approach 294

to the rank-based evaluation. The complete cumulative-based, rank-based and amber results are in the Supplementary Material 2.

The results for the CAMI simulated and real datasets should be interpreted considering the 297 depth of classification. Most tools classify at a certain taxonomic level, either specific rank (e.g. 298 species) or any taxon. Clark provides only species assignments and it was evaluated together with 299 all other tools providing results at any taxonomic level (centrifuge, diamond, ganon, and kraken). 300 Centrifuge, ganon and krakenuniq are also able to classify sequences at assembly level. Centrifuge 301 outputs at sequence level, thus an extra step of applying an LCA algorithm for non-unique matches 302 was necessary to generate results at assembly and taxonomic levels. Given the availability of the 303 ground truth, only simulated data was evaluated up to assembly level while real data was evaluated 304 at taxonomic levels. 305

306 3.1 Indexing

The set of reference sequences from RefSeq-OLD/CG/ALL (Table 1) and RefSeq-CG/ALL-top-3 (Table 2) were used as inputs to generate the indices for each evaluated tool. Here evaluation is

done by total run-time, memory consumption and final index size (Table 3 and 4).

Reference	Method	time	Memory	Index size	
RefSeq-OLD	centrifuge	02:51:03	105	4	
	clark	04:07:56	161	35	
	diamond	00:08:07	30	3	
	ganon	00:02:08	24	16	
	kraken	02:04:16	93	79	
RefSeq-CG-top-3	centrifuge	06:51:25	269	12	
	clark	08:45:31	243	81	
	diamond	00:10:33	28	10	
	ganon	00:07:01	68	63	
	kraken	04:53:31	200	184	
RefSeq-ALL-top-3	diamond	00:36:23	32	76	
	ganon	00:54:48	266	267	

Table 3: Build times, memory consumption and index sizes at taxonomic level. Memory and Index size in GiB. All tools build at taxonomic leaf nodes (taxid) besides clark building at species level. Tools running more than 24 hours to build were not considered. 48 threads were used for all tools. Computer specifications and parameters used are in the Supplementary Material 1 - Section 1.1 and 1.4. Krakenuniq was not evaluated on taxonomic level since it runs exactly the same base algorithm as kraken in this configuration.

Reference	Method	time	Memory	Index size
RefSeq-OLD	centrifuge	02:51:03	105	4
	ganon	00:02:22	32	25
	krakenuniq	02:06:41	93	79
RefSeq-CG	centrifuge	12:32:08	459	21
	ganon	00:10:49	108	100
	krakenuniq	08:54:56	345	204
RefSeq-ALL	ganon	02:30:47	530	539

Table 4: Build times, memory consumption and index sizes at assembly level. Memory and Index size in GiB. Tools running more than 24 hours to build were not considered. 48 threads were used for all tools. Computer specifications and parameters used are in the Supplementary Material 1 - Section 1.1 and 1.4

When indexing the RefSeq-CG-top-3 at taxonomic levels (Table 3), the evaluated tools took between 7 minutes and 8 hours, resulting in ganon being the fastest and clark the slowest. We do

not consider runs taking more than 24 hours to build indices, given that they clearly do not scale 31 2 well enough to index high amounts of data and will not be able to keep indices up-to-date in a 31 3 reasonable amount of time for new data (Supplementary Material 1 - Section 2.2). Ganon shows 314 a significant overall reduction in memory consumption and run-time compared to the other tools 31 5 besides diamond. However, diamond is the only tool using protein data, accounting approximately 316 for a third of the volume of the genomic data. Ganon builds 41 times faster than kraken, the 317 second fastest using the same data source. Centrifuge achieves the lowest index size with the 31 8 cost of having the highest memory consumption. Additionally, ganon is able to generate smaller 31 9 indices at the cost of speed in the classification step, without harming sensitivity. Ganon indices 320 for RefSeq-CG-top-3 can be as small as 21GiB. RefSeq-ALL-top-3 was built in under an hour for 321 diamond and ganon, with diamond providing a smaller filter and lower memory consumption. We 322 could not run centrifuge, clark, kraken and krakenuniq for RefSeq-ALL on our infrastructure, given 323 computational limitations or long execution time. A recent publication [11] reported that kraken 324 and consequently krakenuniq both need 11 days to build a database for the bacterial RefSeq version 325 80, an approximate of the RefSeq-ALL here evaluated, with a more powerful server consisting of 326 64 cores of E7-8860v4 CPUs and three terabytes of memory. Estimated run times for these tools 327 in the evaluated datasets can be found in the Supplementary Material 1 - Figure 6. 328

When building indices on assembly level (Table 4), ganon took around 10 minutes to index RefSeq-CG while the second fastest tool, krakenuniq, took almost 9 hours. Given our computational and time limitations, ganon was the only tool able to build indices on assembly level for the RefSeq-ALL dataset, taking 2 hours and 30 minutes.

333 3.2 Updating

Ganon is the only tool among the evaluated ones that allows for incremental updates on previously generated indices. We evaluated this functionality on Bacterial sequences added to RefSeq-CG dating from 19-December-2018 to 21-January-2019, comprising 2.77 Gbp, 1307 sequences, 370 species from which 213 are new to the reference set and 716 new assemblies (Supplementary Material 1 - Table 4). Updating the ganon index based on RefSeq-CG with this dataset finished under 5 minutes, less than half of the time necessary to create the index (Table 4).

340 3.3 Classifying

Figure 3 compares in a cumulative-based fashion the results of one simulated high complexity 341 dataset (CAMI toy set) classified against the indices based on RefSeq-OLD, RefSeq-CG-top-3 **34** 2 and RefSeq-ALL-top-3. In this analysis we can observe how each method performs classifying 34 3 reads to their ground truth targets up to a certain taxonomic level. The overall improvement 344 in terms of sensitivity and precision is clear when using a more complete and up-to-date set of 34 5 references (RefSeq-ALL-top-3), since they provide higher coverage for the evaluated ground truth 34 6 targets (Supplementary Material 1 - Figure 5). The highest F1-Score at any taxonomic level is 347 achieved with ganon using RefSeq-ALL-top-3. Diamond shows an increase in performance at higher 348 taxonomic levels but performs poorly at species level. Clark classifies only at species level and has 34 9 no improvements in higher taxonomic levels. Metrics for the complete RefSeq-CG and RefSeq-ALL 350 differ slightly from the respective top-3 sets, therefore they were not included in the evaluations 351 (Supplementary Material 1 - Section 2.3.1). This indicates that over-representation filtering does 352 not affect the results but it can speed up analysis. 353

When looking at the metrics by each rank individually (Table 5, Supplementary Material 1 -Figure 7), the overall precision and sensitivity values are greater, since incorrect classifications at lower levels are not penalized in this type of evaluation. Besides diamond, which underperforms at species level, all tools have overall similar performance values using RefSeq-OLD and RefSeq-CG-top-3. However, ganon shows improvement on sensitivity on all levels with RefSeq-ALL-top-3, being 12% more sensitive at species level and reaching 99.27% precision at genus level. For lower ranks (species and species+) results were mainly limited by the availability of reference sequences (Supplementary Material 1 - Figure 5).

The same analysis was performed on real data (CAMI challenge set). This set is more challenging since most of the species in the sample are novel and, still to this date, mostly not present in the analyzed repositories of reference sequences (Supplementary Material 1 - Figure 5). As stated by the CAMI results [6], tools performed poorly in this dataset in terms of sensitivity (Figure 4). Here the impact of a larger and up-to-date set of references (RefSeq-ALL-top-3) is more evident,



Figure 3: Cumulative-based precision, sensitivity and F1-Score for the simulated reads. Colors represent different reference sets: blue = RefSeq-OLD, orange = RefSeq-CG-top-3, red = RefSeq-ALL-top-3

Reference	Method	$\mathbf{S}\mathbf{e}\mathbf{n}\mathbf{s}\mathbf{i}\mathbf{t}\mathbf{i}\mathbf{v}\mathbf{i}\mathbf{t}\mathbf{y}$	Precision	F1-Score
RefSeq-OLD	centrifuge	41.49%	79.60%	54.55%
	clark	41.13%	84.29%	55.29%
	diamond	4.96%	29.30%	8.49%
	ganon	40.78%	88.68%	55.87%
	kraken	41.23%	83.92%	55.30%
RefSeq-CG-top-3	centrifuge	43.57%	79.12%	56.19%
	clark	42.93%	82.43%	56.45%
	diamond	11.30%	74.41%	19.62%
	ganon	41.86%	87.56%	56.64%
	kraken	43.14%	82.32%	56.61%
RefSeq-ALL-top-3	diamond	13.10%	88.64%	22.82%
	ganon	$\mathbf{53.99\%}$	$\mathbf{94.91\%}$	68.83%

Table 5: Rank-based precision, sensitivity and F1-Score values for the simulated reads at species level The use of a larger reference set with RefSeq-ALL-top-3 significantly improves results. Only ganon and diamond indexed the RefSeq-ALL-top-3 in less than 24 hours. Highlighted values for the best results with this dataset at species level. Results for all taxonomic levels are in the Supplementary Material 1 - Figure 7 and Supplementary Material 2

thus significantly improving the results on both sensitivity and precision. The same trend from the simulated data analysis is present, with ganon achieving best results up to species level and diamond improving classifications at higher levels but having poor resolution at lower ranks.

diamond improving classifications at higher levels but having poor resolution at lower ranks.
In the rank-based analysis (Table 6, Supplementary Material 1 - Figure 8) ganon has 10% higher
F1-Score compared to diamond with the RefSeq-ALL-top-3 as species level. Sensitivity has a peak
of 10% and 25% at species+ and species levels, respectively, which are not far from the maximum
possible using this reference set (12% and 32% respectively). Similar results can be seen in amber
evaluation (Figure 5). Comparing results between RefSeq-CG-top-3 and RefSeq-ALL-top-3, genus
level sensitivity went from 13% to 83% with a significant improvement in precision, reinforcing the
need for bigger and more diverse reference sets to analyze metagenomics data.

Table 7 compares the assembly level classification between centrifuge, ganon and krakenuniq. There is an overall decrease in precision and sensitivity from RefSeq-OLD to RefSeq-CG. Precision is greater using RefSeq-ALL but sensitivity is still greater with RefSeq-OLD. However, RefSeq-CG has more than 6 times the number of assemblies of RefSeq-OLD, while RefSeq-ALL has almost 50 times more assemblies (Table 1). As reported before [11], higher diversity in the references does



Figure 4: Cumulative-based precision, sensitivity and F1-Score for the real reads. Colors represent different reference sets: blue = RefSeq-OLD, orange = RefSeq-CG-top-3, red = RefALL-top-3

Reference	Method	$\mathbf{S}\mathbf{e}\mathbf{n}\mathbf{s}\mathbf{i}\mathbf{t}\mathbf{i}\mathbf{v}\mathbf{i}\mathbf{t}\mathbf{y}$	Precision	F1-Score
RefSeq-OLD	centrifuge	0.51%	2.24%	0.84%
	clark	0.49%	3.21%	0.86%
	diamond	0.00%	0.00%	0.00%
	ganon	0.45%	6.68%	0.85%
	kraken	0.50%	3.13%	0.86%
RefSeq-CG-top-3	centrifuge	2.41%	7.03%	3.59%
	clark	2.34%	9.57%	3.76%
	diamond	1.74%	11.23%	3.02%
	ganon	1.89%	19.14%	3.44%
	kraken	2.39%	9.61%	3.83%
RefSeq-ALL-top-3	diamond	12.38%	55.84%	20.27%
-	ganon	25.03%	38.97%	$\mathbf{30.48\%}$

Table 6: Rank-based precision, sensitivity and F1-Score values for the real reads at **species level.** The use of a larger reference set with RefSeq-ALL-top-3 significantly improves results. Only ganon and diamond indexed the RefSeq-ALL-top-3 in less than 24 hours. Highlighted values for the best results with this dataset at species level. Results for all taxonomic levels are in the Supplementary Material 1 - Figure 8 and Supplementary Material 2

not always translate to an improved accuracy in the classification. This was also noticed when 382 using the complete NCBI-nt database to analyze the same dataset (Supplementary Material 1 -

383

Figure 13). 384 In the specific case of methods evaluated here, small differences between very similar assemblies 385 are difficult to be identified due to the resolution of each method. This means that they, in general, 386 can correctly classify sequences to target assemblies given a certain similarity threshold. However, 387 they are unable to select the correct assembly, thus providing the lowest common ancestor at a lower 388 resolution. This can be seen in Supplementary Material 1 - Figure 12, where the overall sensitivity 389 and precision of all tools executing in assembly mode did not affect the taxonomic metrics and are 390 comparable to the same tools running in taxonomic mode. Even though the assembly step does 391 not provide accurate enough results, centrifuge and ganon are the only tools that can provide a 392

list of all matches/candidates that can be further analyzed with high resolution methods [32]. 393

In most scenarios evaluated, ganon consistently provides greater precision classifying reads to 394 their ground truth targets within the same reference set, while keeping sensitivity values high, with 395 little variation to the other methods. High precision translates to fewer reads with a wrong clas-396



Figure 5: AMBER completeness/sensitivity (green) and purity/precision (blue) values for real reads. Results for diamond (left) and ganon (right) using RefSeq-ALL-top-3 set of references. Strain level in AMBER plots are equivalent to species+ in our evaluations.

Reference	Method	Sensitivity	Precision	F1-Score
RefSeq-OLD	centrifuge ganon krakenuniq	22.78% 22.32% 22.68%	$64.54\%\ 77.95\%\ 69.66\%$	33.68% 34.70% 34.22%
RefSeq-CG	centrifuge ganon krakenuniq	$\frac{11.82\%}{11.52\%}\\11.67\%$	$30.77\%\ 37.25\%\ 32.45\%$	17.08% 17.60% 17.17%
RefSeq-ALL	ganon	21.56%	87.89%	34.62%

Table 7: Rank-based precision, sensitivity and F1-Score values for the simulated reads at assembly level. Only ganon indexed the RefSeq-ALL in less than 24 hours. Highlighted values for the best results with this dataset at assembly level. Results for all taxonomic levels are in the Supplementary Material 2.

sification. Sensitivity is strongly improved in more diverse reference sets, especially with RefSeqALL-top-3. Looking at rank-by-rank performance, ganon improved F1-Score in every taxonomic
rank (Supplementary Material 1 - Figures 7 and 8), with F1-Score up to 46% higher than diamond
with the same reference at species level (Table 5).

Table 8 compares the performance of the analyzed tools in terms of how many base pairs they 401 can classify per minute (Mbp/m), wall/elapsed time and memory usage. Kraken is the tool with 402 the fastest runtime on classification step and diamond with the slowest. Although comparisons 403 with diamond were made, it is important to notice that the tool works in a very different way using 404 protein data and performing alignments, thus explaining the huge difference in execution times. 405 Ganon can be configured to run in offset mode, thus skipping a certain number of k-mers and 406 speeding up classification. offset = 1 means that all k-mers are being evaluated while offset = 2407 means that every 2nd k-mer is being skipped. The trade-off between offset and precision/sensitivity 408 for ganon results can be seen in Supplementary Material 1 - Figure 11. Speed variation between 409 simulated and real reads is partly explained due to their classification rate: on average 70% of the 410 simulated reads are classified while only 20% of the real reads are classified. Memory consumption 411 is mainly based on the index size of each tool (Table 3), with little variation besides that. 412

413 4 Discussion

We presented ganon, a novel method to index big sets of genomic sequences and classify short reads against them in a taxonomic oriented scheme. Ganon's strengths are an ultra-fast indexing method for large sets of reference sequences that incorporates a novel application of Interleaved

		Simulated			Real		
Reference	Method	Mbp/m	Wall time	Memory	Mbp/m	Wall time	Memory
RefSeq-CG-top-3	centrifuge	298	$00:24:59~(\pm~51s)$	14	802	$00:09:19 \ (\pm 4s)$	14
	clark	1104	$00:06:44~(\pm~5s)$	108	1208	$00:06:11~(\pm 4s)$	107
	diamond	36	$03:27:00~(\pm 259s)$	15	33	$03:40:55~(\pm~170s)$	16
	ganon	406	$00:44:05~(\pm~54s)$	65	573	$00:14:19~(\pm~6s)$	65
	kraken	2113	00:03:46 (± 1s)	190	2734	00:02:57 (± 3s)	190
RefSeq-ALL-top-3	diamond	6	$18:23:09 \ (\pm \ 729)$	23	5	$21:23:00 \ (\pm \ 181s)$	23
	ganon	115	04:42:32 (± 216s)	261	163	00:49:11 (± 12s)	261

Table 8: Classification performance. Memory in GiB. Full set of simulated and real reads classified with 48 threads. Centrifuge, clark and diamond performance in Mbp/m calculated from wall time. Values are the average of 4 out 5 consecutive runs (excluding the slowest run), with standard deviation for the run time in parentheses. Computer specifications and parameters used are in the Supplementary Material 1 - Section 1.1 and 1.4

Bloom Filters and a precise classification with k-mer counting and filtering. Unlike DREAM-Yara, an alignment-based read mapper that uses the IBF as a pre-filter for the distributed Yara mapper, ganon uses the IBF as the main index structure to provide an alignment-free assignment of sequences. This is only possible by creating taxonomic constrained clusters with TaxSBP in any desired taxonomic level. Ganon additionally applies an LCA algorithm as a final step to have one classification per sequence. In addition it also provides updatability of indices, multi-hierarchy support for classification, assembly level support and taxonomic reports.

By indexing large sets of reference sequences and turn them into searchable indices, ganon allows 424 scientists to make most of their data. Short turnaround times for index building and updating 425 are crucial for many bioinformatics applications (e.g. outbreak investigation). In our evaluations, 426 building the complete RefSeq and classifying 49 million reads against it performed under 2 hours 427 with ganon, from raw reference sequences and reads to taxonomic reports, while diamond required 428 more than 22 hours to classify the same set. Other methods required at least 24 hours to build the 429 indices. Without a dedicated infrastructure for constant reconstruction of indices and databases, 430 tools evaluated in this work are unable to keep up with the fast growing rate of reference sequence 431 repositories. That results in either long time to start analysis or use of outdated reference sets. 432 Ganon facilitates database maintenance, allowing short increments on a daily basis being the only 433 realistic option to keep-up with the fast pace of data generation. In addition, ganon indices are 434 flexible and can be built for different taxonomic levels (e.g. genus), requiring less space and memory, 435 consequently improving classification speed. A trade-off between filter size, clustering and false 436 positive rate is also possible, simply by sacrificing precision over performance or disk usage over 437 classification speed. 438

Classification results presented here are on par with state-of-the-art methods with regards 439 to sensitivity, while improving precision rates in almost every scenario of our cumulative-based 440 evaluations. Results are consistent across all three evaluation methods (cumulative- and rank-441 based and amber) indicating the robustness of findings. We attribute this improvement to an 442 application of the k-mer counting lemma together with a progressive filtering step, which can 443 better separate false from true positives. The unique filtering step also allows for better selection 444 of false positives when taxonomic groups are underrepresented in the reference set. In addition, 445 instead of only reporting reads at a fixed LCA level, ganon provides every output for a read at a 446 taxonomic or assembly level. This is crucial for strain level analysis, where candidate organisms 447 are more insightful for further investigations than a conservative identification. 448

Even with ganon achieving improved results in classification, in general terms, the methods 449 tested here perform similarly when based on the same underlying set of reference sequences. The 450 difference in sensitivity when using a high quality set (RefSeq-ALL) compared to only complete 451 genomes (RefSeq-CG) or an outdated set (RefSeq-OLD) is very significant and tends to get bigger 452 with more sequences added to this repository. Thus the choice of the database is crucial and should 453 not be overlooked when analyzing metagenomics data. Even though centrifuge, clark, kraken and 454 krakenuniq could potentially perform well with more reference sequences, their indexing times are 455 highly prohibitive. 456

⁴⁵⁷ When using highly diverse reference sets or when aiming at high resolution classification (e.g. ⁴⁵⁸ assembly level), the evaluated methods shown decreased performance. However, in a scenario of

data exploration of an unknown environmental sample, the ability to classify reads against huge 459 sets of very diverse reference sequences (e.g. NCBI-nt) can be helpful. Therefore, in those scenarios 460 we recommend to perform analysis hierarchically, first classifying reads against high quality ref-461 erences and only using high diverse reference sets for unclassified sequences, adjusting error rates 462 accordingly. This approach can be easily done with ganon's implementation of multi-filter and 463 multi-hierarchy classification. This functionality tied to fast indexing of reference sets make ganon 464 a powerful tool for exploratory data analysis, enabling multiple combinations of indices and error 465 rates in an iterative manner. An example of this functionality can be found in Supplementary 466 Material 1 - Section 2.4, where we analyzed real data from TARA oceans [10], building several 467 indices and classifying reads against them in an exploratory-fashion. 468

Ganon's fast indexing performance is mainly due to the fact that k-mers are not being counted. 169 Instead, all of them are inserted into a space-efficient data structure (IBF) that also provides 470 quick look-up times. However, data generation is constantly increasing and in the long term this 471 approach will reach a limit. For that reason, a k-mer aware clustering combined with a minimizer 472 implementation could improve performance in the data structure as well as memory consumption. 473 These features are planned for future releases. Even though we based our analysis on large and 474 realistic datasets, time efficiency purely based on data can be misleading. Thus, the scalability of 475 the methods can only be deduced. As a future work we propose a comparison of time and space 476 complexities of each methodology and how they would perform in the long term, considering a 477 continuous and exponential data growth. 478

Ganon manages to index large sets of reference sequences while keeping them updated in very short time. In addition, classification results for ganon are as good as or better than the evaluated tools and it runs in competitive time. To the best of our knowledge, ganon is the only tool with update capabilities, which is performed in a fraction of the complete build time. This poses as an advantage to maintain up to date with the public repositories of genomic data and their frequent updates. To conclude, we believe that ganon can be a useful tool for metagenomics analysis in a time where reference sequence repositories are growing fast.

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6 Author contributions

V.C.P. designed, implemented and tested the tool and the experiments and wrote the manuscript.
T.H.D. created and implemented the IBF and reviewed the manuscript. E.S. improved and tested
the IBF and reviewed the manuscript. K.R. and B.Y.R. designed and supervised the project,
discussed the experiments and reviewed the manuscript. All authors read and approved the final
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