Predicting Human Minisatellite Polymorphism

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We seek to define sequence-based predictive criteria to identify polymorphic and hypermutable minisatellites in the human genome. Polymorphism of a representative pool of minisatellites, selected from human chromosomes 2I and 22, was experimentally measured by PCR typing in a population of unrelated individuals. Two predictive approaches were tested. One uses simple repeat characteristics (e.g., unit length, copy number, nucleotide bias) and a more complex measure, termed HistoryR, based on the presence of variant motifs in the tandem array. We find that HistoryR and percentage of GC are strongly correlated with polymorphism and, as predictive criteria, reduce by half the number of repeats to type while enriching the proportion with heterozygosity ≥0.5, from a background level of 43% to 59%. The second approach uses length differences between minisatellites in the two releases of the human genome sequence (from the public consortium and Celera). As a predictor, this similarly enriches the number of polymorphic minisatellites, but fails to identify an unexpectedly large number of these. Finally, typing of the highly polymorphic minisatellites in large families identified one new hypermutable minisatellite, located in a predicted coding sequence. This may represent the first coding human hypermutable minisatellite.

[Supplemental material is available online at www.genome.org.]

Tandem repeats represent a significant fraction of vertebrate genomes and have been classified as satellites, minisatellites, and microsatellites according to the length of the repeated unit and the overall length of the array. Minisatellites are usually defined as the tandem repeats of a short (10- to 100-bp) motif spanning several hundred to several thousand base pairs and are associated with interesting features of genome biology (for review, see Vergnaud and Denoeud 2000).

Minisatellites frequently exhibit length polymorphism, which results from variation in the number of internal copies, making them valuable genomic markers. They provided the first highly polymorphic, multiallelic markers for linkage studies (Bell et al. 1982; Nakamura et al. 1987) and were used in the early stages of human genome mapping (NIH/CEPH Collaborative Mapping Group, 1992). Chromosomal distribution of minisatellites in the human genome is highly skewed toward telomeres and ancestrally telomeric regions (Amarger et al. 1998). Highly polymorphic minisatellites are thus a good tool for detection of microdeletions in the ends of chromosomes, associated with human pathologies such as mental retardation (Giraudeau et al. 2001). Polymorphic minisatellites are also found in bacterial genomes (Le Fleche et al. 2001), in which they have proven to be a powerful tool for bacterial strain identification.

Although the abundance of polymorphic minisatellites suggests that they are fast-evolving sequences, most of them are, in fact, quite stable. New alleles that display changes in the number of tandem copies have been observed at only a few loci, called hypermutable minisatellites. Changes at these loci in the germline can be observed in the next generation, and in humans, one locus, D2S90 (CEB1), has been found to

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Article and publication are at http://www.genome.org/cgi/doi/10.1101/ gr.574403. Article published online before print in April 2003. change in as many as 13% of the gametes (Vergnaud et al. 1991; Vergnaud and Denoeud, 2000). Hypermutable minisatellites may provide a potent source of information on the mechanism of minisatellite instability. In humans, this instability apparently arises at least in part through gene conversion events, during or shortly after meiosis, many of which involve interallelic transfers of information (Buard and Vergnaud 1994; Jeffreys et al. 1994; May et al. 1996; Buard et al. 1998). Similar intraallelic and interallelic recombination events are found in MS32 and CEB1 minisatellite sequences, when they are placed close to a meiotic hotspot in Saccharomyces cerevisiae (Appelgren et al. 1997, 1999; Debrauwère et al. 1999). Most likely, these events result from the gene conversion repair of double-strand breaks, as recent evidence indicates that meiotic recombination in mammals and yeast is initiated by the Spo11p endonuclease (Bergerat et al. 1997; Keeney et al. 1997; Baudat et al. 2000; Romanienko and Camerini-Otero 2000), which is also essential to the meiotic instability of the minisatellites introduced in yeast (Debrauwère et al. 1999). In agreement with these observations, it has been proposed that the meiotic hypermutability of some minisatellite structures is the byproduct of the coincidence of an ordinary minisatellite with a double-strand break hotspot (Vergnaud and Denoeud 2000).

Interestingly, hypermutable minisatellites might additionally provide biomarkers for low-dose exposure of the human germline to ionizing radiation (Dubrova et al. 1993, 1997; Dubrova and Plumb 2002). Unfortunately, <10 human hypermutable loci have been characterized so far, using approaches developed >10 years ago, whereas the population studies conducted to evaluate the effect of low-dose irradiation would greatly benefit from the availability of a larger panel of probes.

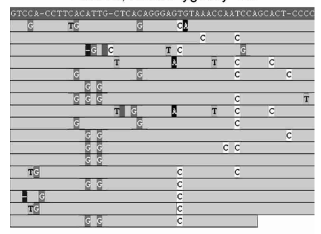
Given the multifaceted utility of minisatellites, determining which are polymorphic/hypermutable would seem a valuable task. Efficient tandem repeat detection software en-

ables the identification of tandem repeats across entire genomes (Benson 1999; Vergnaud and Denoeud 2000), so that testing for polymorphism is all that is required. But although the polymorphism of the few dozen minisatellites usually present in a small genome can be systematically assayed at a reasonable cost (Le Fleche et al. 2001), this is not a realistic option for the human genome. There, the number of minisatellite loci is estimated in the thousands (based on the sequence of chromosome 22: Vergnaud and Denoeud 2000). the proportion of highly polymorphic minisatellites among these is not known, and previous efforts to identify hypermutable loci among minisatellites have produced only very low yields (~1% to 3% of those examined). Furthermore, sequence analysis of a few hypermutable loci has not yet revealed specific features that might facilitate their identification (Murray et al. 1999). Of need are predictive criteria that can be applied before the expensive and labor-intensive step of polymorphism typing.

Earlier attempts at polymorphism prediction for tandem repeats focused on microsatellites. Fondon III et al. (1998) identified polymorphic loci by selecting microsatellites in which the individual copies were at least 90% identical to a core pattern, but that study did not include a control group to test whether selection yielded higher polymorphism values than the background rate. Wren et al. (2000) improved polymorphic microsatellite identification by requiring perfect homogeneity of the repetitive unit. Such results are in accordance with the mutation process of microsatellites (replication slippage): They are stabilized by variant repeats (Weber 1990), the presence of which facilitates detection of slippedstrand DNA by the mismatch repair system (Strand et al. 1993; Heale and Petes 1995). In the case of minisatellites, in which internal conservation is not the rule at currently known hypermutable loci (Murray et al. 1999; Vergnaud and Denoeud 2000), such a high conservation requirement imposes too great a restriction on the set of potentially useful repeats and, as we report below, would preclude finding both highly polymorphic and hypermutable repeats.

The purpose of this report is to define inexpensive strategies to accelerate the search for highly polymorphic minisatellites. The goal has been the development of sequencebased predictive criteria for polymorphism. Results are based on the study of a representative pool of minisatellites selected from human chromosomes 21 and 22. Polymorphism for these loci was experimentally measured by typing in a population of unrelated individuals. This was followed by typing the most polymorphic loci across a number of large families to test for hypermutability. Two predictive approaches were tested. The most straightforward takes advantage of the availability of two different releases of the human genome sequence: one from the public genome sequencing project and the other from the private Celera project. The second approach uses sequence-based characteristics of the repeats including such simple measures as unit length, copy number, degree of conservation, percentage of GC (%GC)— and a more complex measure based on the internal organization of variant motifs in the tandem array. A repeat that contains several distinct sets of nearly identical mutations exhibits prima facie evidence of multiple rounds of expansion and may be more likely to exist as multiple alleles than a repeat that contains mostly unique mutations (Fig. 1). This later measure is analyzed by using history reconstruction (Benson and Dong 1999), a type of parsimony analysis that infers how the present day sequence could have evolved from a single

CEB252: 16.8 x 50 bp HistoryR=0.76, %matches=88% 4 alleles, Heterozygosity=0.6



CEB233: 17.3 x 43 bp HistoryR=0.14, %matches=82%, 1 allele, heterozygosity=0

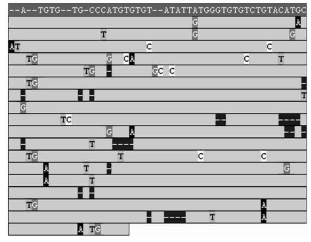


Figure 1 Multiple alignments of tandem repeats CEB252 and CEB233. In each alignment, the upper darker line is a consensus pattern for the basic unit, shown for reference, and the lighter lines are the individual copies, ordered from *top* to *bottom* as they occur in the repeat. Only differences with the consensus are shown. Heterozygosity for CEB252 is 0.6. Note several redundant patterns of mutation resulting in a high HistoryR score. Heterozygosity for CEB233 is zero. No clear organization of mutations resulting in a low HistoryR score.

ancestral copy while undergoing a minimum number of point mutations interspersed with duplications.

RESULTS

Characterization of Chromosome 21 and 22 Minisatellites

Human chromosomes 21 and 22 contain ~15,000 tandem repeats each (as detected by tandem repeats finder [TRF] in the

publicly available sequences that exclude heterochromatin; Benson 1999). For this study, the empirical definition of minisatellites follows the suggestion made in Vergnaud and Denoeud (2000), which is more stringent than the usual definition of minisatellites mentioned in the introduction: (1) unit length \geq 17 bp, (2) copy number \geq 10, (3) total length \geq 350 bp, (4) percent matches ≥70%, and (5) GC bias (i.e., strand asymmetry for G and C; see Methods section) ≥0.35. This definition includes repeats clearly classified as minisatellites, not microsatellites, allows minisatellites shorter than the ≥800 bp usually identified by Southern blotting (Vergnaud 1989; Amarger et al. 1998) and removes repeats with highly diverged copies. On chromosomes 21 and 22, 127 tandem repeats fulfill these criteria. Table 1 indicates their position on the chromosomes. As described before for minisatellites derived by classical approaches (Amarger et al. 1998), they are mainly located toward chromosome ends (both chromosomes are acrocentric). Analysis shows no statistically significant differences between the minisatellites from chromosome 21 and 22 for any of the characteristics listed in Supplementary Table 1. The two chromosomes will subsequently be considered together.

PCR Typing Results

Polymorphism results, that is, number of alleles observed and heterozygosity, are given in Table 1, as well as dbSNP accession numbers for the polymorphic minisatellites that were submitted to the SNP database (http://www.ncbi.nlm.nih. gov/SNP/index.html). Supplementary data about polymorphism is also available at http://minisatellites.u-psud.fr. For the minisatellites that were typed first ("training set"), the study was made on a population of 76 unrelated individuals. Results were comparable to those obtained with a subset of 28 unrelated individuals from the set of 76. Subsequent PCR typings (minisatellites from the "test set") were performed only on the 28 individuals, except for the most polymorphic loci that were typed in all 76 individuals in order to evaluate their polymorphism more accurately.

Among the 127 minisatellites, 118 were successfully amplified (55 on chromosome 21 and 63 on chromosome 22) by using the selected primer pair (Table 1). Not surprisingly, long minisatellites (>2 kb) are the most difficult to amplify: Only five among eight were successfully amplified under the conditions used. Figure 2A shows the image of the gel obtained for minisatellite CEB285 on 32 individuals (including 28 unrelated individuals): Six different alleles can be assigned. About 75% of the minisatellites successfully amplified are polymorphic (i.e., two alleles or more), and 42% have a heterozygosity value \geq 0.5.

Polymorphism Prediction: Sequence Characteristics and History Reconstruction

Training Set

Twenty-five out of 60 and 32 out of 67 minisatellites were picked randomly, from chromosomes 21 and 22 respectively, to be typed first: They form the training set. PCR amplification was successful on 51 out of 57. A comparison of the sequence and polymorphism characteristics between the training set and the remaining minisatellites showed that the two sets have comparable distributions except for percentage of matches, purine/pyrimidine bias, and GC bias. To determine if some sequence characteristics are associated with high

polymorphism, correlations between sequence characteristics and allele number or heterozygosity were calculated for the training set. The greatest correlations were obtained for HistoryR (a measure derived from the tandem repeats history reconstruction algorithm [Benson and Dong 1999]; see Methods section) and %GC (Fig. 3). Weaker correlations were also found for average entropy (strongly correlated with HistoryR), and unit length (data not shown). Based on these observations, we chose to test three predictive criteria: criterion 1, minisatellites with HistoryR \geq 0.54; criterion 2, minisatellites with %GC \geq 48%; and criterion 3, minisatellites with HistoryR \geq 0.54 and %GC \geq 48%.

Test Set

Of the remaining 70 minisatellites, 67 were successfully amplified and used as a test set in order to confirm the predictive criteria deduced from the training set. For each of the three criteria, the test set was partitioned into two groups: a positive group fitting the predictive criterion and a negative group. Figure 4A illustrates the results: All three criteria are predictive, that is, heterozygosity and allele number are significantly higher in the positive group compared with the negative group. The best polymorphism prediction was obtained with criterion 3 (HistoryR and %GC combined). It produces an enrichment of repeats having heterozygosity ≥0.5 from 43% (29 of 67) in the test set to 59% (19 of 32) in the positive group and a diminishment of monomorphic repeats from 25% (17 of 67) in the test set to 6% (two of 32) in the positive group. Criterion 3 thus reduces by half (67 to 32) the number of minisatellites to type while eliminating most monomorphic minisatellites and keeping most polymorphic ones (Fig. 4A). One among five highly polymorphic minisatellites (heterozygosity ≥0.85%) would have been missed using criterion 3.

Polymorphism Prediction: Direct Sequence Comparison

The experimental polymorphism values measured here indicate that greatly enhanced efficiency of polymorphic loci identification is possible if the sequences of two independent alleles for each locus are available. The reasoning is that two random samples of a moderately or highly polymorphic locus will, with high probability, yield different alleles, whereas for a monomorphic or only slightly polymorphic locus, the alleles will likely be identical. Thus, selection based on observed allele difference in the two samples should enhance the proportion of loci obtained that are polymorphic. The applicability of this approach was directly tested by comparing sequences from the Human International Genome Sequencing Consortium (HGP) and Celera genomics. We establish selection criterion 4 to be different reported lengths in these two sequences. For the 127 minisatellites previously identified in the HGP sequence, repeat sizes in the sequence provided by Celera (Venter et al. 2001) were obtained by BLAST with the PCR primers. Three tandem repeats were not found in the Celera sequence, including two that were typed (CEB230, CEB256) and one long repeat (CEB215; length expected from HGP = 2834 bp) that could not be typed. Of the remainder, 51% (29 of 57) have a different length in the two sequences for chromosome 21 and 22% (15 of 67) for chromosome 22. From the measured heterozygosity values, we would expect 37% (43 of 116) to have different lengths between the two sequences, essentially the same as found. None of these

900

580

1090 920

(pb)

720 890 760

(continued)

1980-2550 Observed size range 1600-4500 1500-1850 1100-2900 830-3100 900-1450 660-1420 910-960 650-850 810-1200 900-1060 650-1200 600-670 490-550 610-640 820-870 640-690 600-640 410-770 640-860 850-900 610-710 830-880 500-690 1510 840-870 590-670 1050 1660 ALLELE SIZE INFORMATION sednence Celera 1017 1044 2038 7305 000 (pp) 720 917 1026 636 819 357 813 642 904 498 280 1511 541 069 657 539 040 1241 483 290 635 765 682 491 831 763 644 901 647 891 Length predicted by Human Genome Project sednence (pb) 2227 1454 1045 720 543 917 1086 830 636 498 1192 813 642 899 009 580 1241 1508 838 912 3109 1654 650 1033 1040 2553 635 765 645 821 737 589 643 901 891 761 Heterozygosity POLYMORPHISM (28ind) 0.88* 0.39 0.83* 0.19 0.79* 0.14 0.36 0.36 0.64 0.04 0.47 0.29 0.56 69.0 0.57 99.0 0.37 0.27 0.17 0.04 0.23 0.57 0.7 0 0 9.0 0 0 0.51 0.07 0 0 0 0 0 0 alleles (28ind) Table 1. List of the 118 Minisatellites That Were Typed: PCR Conditions, Polymorphism Results, and Allele Size Information 14* 10* 2 2 * က 2 2 9 2 0 8 4 / 7 N emperature Annealing O.89 O.89 57°C 64°C S,89 0.89 C 0.89 C 59°C 63°C O.89 0.89 O.89 S8°C O.89 S89 O.89 0.89 O.89 57°C 08°C 59°C 0°89 O.89 0.89 0.89 0.89 0.89 08°C 54°C C8°C 68°C C8°C 68°C 57°C C89 55°C 64°C CCTGAAGAGACAAGGAGAAAGGATGAC GCTTATGTATGAGGCAGCAAATAGGATCAG AAGACATAAGTGCTCCAGGTGTTAACAGGA STTGAGGAGGACTGAGCCTTCAGAAGTTAG TAAACTAGATGAAAGAGTTGCTGCGCCTGA GACATCCAAAAGGCCAATGAGTATATGAAA GGAATGTTTAGCAGCAACCGAATATACCAG AAAGAAAAAGAAAAGCAGGGCTCATACC TATCCAAGTGGGCTCTAGATCCAGTGATAC AGGTTCAGTATTTGAAACAGCCAGATGGTA GTAATAATTCATCCACCCAGATTTGTATGC ATCTTAAATGGGCAGGTTAATGGATTGATG CTTGCAGTATGAGGTCCTCTGTATCTGGTC AATAGCGAAGAGGATGTTCCCAAAAAT GTTTGTCTCCTCTTGCCCTCTGGAAAGTAA CGTGCTGTGGACGTATTAACTTACTGGAAA TACATTTTCTGACTTCTGTGGTCTTCATGG TCTATCTTATTTGACCTCTGCCCGTAGTCC **ATTICTCTGGGTATTTTTCATCTGGAAGCA** ACTCATCTCCTGGTGGAGAGACACTCAC ACCCTAGATGACCTAGTGGGACCTACAC AGCATAACACATTCTGTTTTGGGCATTA CAGTATTCTCACCACTGCACTCAGCAAGT AGAAAACAGGAGACTCACCACGATCAACT **ATCGCCAAGTCATAAAGGTGTACTGTGGT** CTACTTGGGATGCTGAGGCAGGATTATG ATACCAGAGTCCAGCCAAGTTAGCCGTT1 CTGTACCCGGTTAGATCCACACCCTATG AGAGTGTGTGCACACTGTCAGTGAA TTTTGTTGCTGGAACCCTGACAATTTA AGTGCATGGACTGCAGATATTGGGACT AACACACAGAGGCGCAGCAGAGAC ACTGCACTCCAGCCTCTCCCATCCCTA VAACAAAGAGTCCAGGAGGCCCTGAGAGA AATCTGCTCTTTGCCTCCATTCCTCAT **ATGAAGCTGACGCGGAAGATGGTTCT** GAGAGGCTGCCGTCCCCCGGATTGCT CCCAGAATCCCATTCCTGCCCAATG CTTCTTGGGAACATCCATGGCTCAG PCR CONDITIONS AGGTGTACCAGTACAGCAGCTTTGGACCTA GAGACAGTATCAGACACACCAGACAAAAGC CTGAACGAATTAAGTGATGTACCCAGAAGC GATGTAGTTGCATTGCTTGAGTGCATTAAC ACATAACAATCAAAGCAGAGCCTCACTGAC GCTTTCAGTGGCTCTGGAGGTTTTAGTAAG **TTGGAATCCAGAGGAAGCAAGTTACACCAAT** SGGTTCTGGTCTGCAGTTTTTATCTGAGTT **ICTGTGGATAAACGTGAATATGCCCGAAAT IGCAAAATCTCTCCCTCTCTGTTGATAAAA** SATACAGATACGGATGATTCTTGCTCTTGG TGCATTITCTTCTAGGGGGGGTATGACAAGT **AATTGTTGAGGAGACTTCATATTGCTTTCC** AAAGAGCCAAGTGAAGTCCCTTCTCTGAA STTCAGGAGGCTCTGTCTCTGGGTAGAAG STGTGAACATTTTGTAGCCATGTTGTGTTA SCTCAGTTCTCTCTCTATTCGACTTGGTC AGAGCACACCCACCACTAAAATCACTG **ITTTAAATTCTGATTTTCTTGCGAAGGTGA** GCACTTTTGTCCCCATGTGTCATTCTAAC 3GAAAGTGACAGCTCCCTGTTTGAATTAT CAGCTTGGGCAATGGAGTGAGACTGTCT SAGTAGCCACAGGACAGAACTGAGAAGC **ACTCCTTGAGTCTTGGAGGGACTGACAC** CTGGAGGCCAGAAGTTCAAAATCAAGTT AGAGGTGATGAGCACAGGTGGTGAGAG STEGAACCCTGACAATTTTCAAGTGAGG SAACCTCCACCTCCCAGAAAAAAAA CAGGTGCCCCAAGGAGGGGGAGGAG GCACCAACCAGATAGGCCACTGAGAT SCCTAGGGGGGGGGCCTAAGACCA STGCTGTAACCCAGGCTCACAAACCT TCATTCCTGTGAAGCACAGCGTTT Physical position (kb) 19963 21556 23006 23639 28940 29314 30145 30440 30516 30670 30771 30829 31147 31420 31572 31675 31833 32177 32246 32690 33109 33318 24912 26682 31619 32529 33317 33383 33481 10501 31252 33286 4445 7748 3948 220 IDENTIFIER INFORMATION dbSNP ss# 6313648 6313631 6313632 6313646 6313649 6313629 6313638 6313639 6313640 6313643 6313644 6313645 6313628 6313630 6313633 6313635 6313636 6313637 6313641 6313642 6313647 6313634 CEB234 CEB238 CEB245 CEB258 CEB263 CEB265 **CEB266** CEB269 CEB235 **CEB236 CEB272** CEB274 CEB275 **CEB276** CEB239 CEB278 CEB240 CEB241 CEB280 **CEB244** CEB282 CFB256 CEB255 CEB260 CEB261 **CEB264 CEB268** CEB270 **CEB273 CEB237** CEB279 **CEB242** CEB281 CEB243 CEB271 **CEB277** 21 21 21 7 21 21 21 21 21 21 7 21 5 21 7 21 21 21 2 21 21 21 21 21 21 5 21 2 21 21

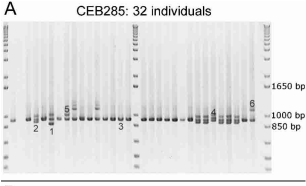
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020-1110 1740-1960 820-910 1300-1340 1300-1350 770-1300 540-1850 580-1100 980-1330 870-1100 630-680 940-1500 1300-1600 1020-4700 1860-2200 1080-1300 1020-2500 1420-1670 1650-2090 880-1090 560-850 900-1700 630-800 710-770 600-1350 830-1800 740-880 430-570 690-820 670-920 550-600 700-720 320-900 680-740 500-520 5000 1200 650 670 530 999 019 1045 092 1514 033 1092 290 868 993 96/ 545 430 3451 374 548 993 539 809 2301 758 539 713 270 394 775 487 858 555 528 352 822 671 738 292 723 758 418 739 713 533 200 0 772 1045 656 1066 1510 1305 2144 647 1737 1016 092 833 563 866 1322 758 649 713 570 822 983 1401 858 555 624 650 805 822 670 269 723 998 820 1790 899 737 713 533 0.39 0.61 0.35 99.0 0.87 0.93 0.55 0.49 0.75 0.53 0.23 0.78 0.79 0.55 99.0 0.51 0.55 0.64 0.07 0.52 0.56 0.58 0.33 0.13 0.71 0 0.3 0.64 0.5 0 0.2 0.5 C 9.0 0 0 0.24 0 0 0.04 0 0.5 9 19 2 2 9 9 S 7 *6 2 2 3 4 4 က 3 2 c 9 c. 3 0 3 2 00 ω 3 3 2 N 3 N S 89 C89°C S89 O.09 C89°C 62°C O.89 C89°C O.89 O.89 0.89 64°C 0°09 J.89 0.89 0.89 59°C O.89 64°C 0.89 C 0.89 08°C 59°C S8°C O.89 C8°C 0°C 68°C 0.89 0°C S89°C 68°C S8°C 2°89 S89 C0.09 58°C S8°C 0°C 38°C 08°C GTTTAGAGAGGAATGGACGCAGAGTGTC CAGCAGCTAATTCTAACGTTTAAGGTCATCC GTAAACCCAACATCGAGAAAACAAGGATGG AAGCGAAGAGAGTGTGTGTGACAGTGACTA GAAAATGTAATCAAGGGACAGGAAGGAAAC GGAGACTAACAGTGGCTACGGGATGTGTTA AAAGTGAGGACAGTAACACAAGGCTTATCG SAAGAGAACTTTCAGATCAGAGCACGAGTT AATCCTAGGTGCATTTCCGTAAACGGTGGT AAGACTAGTTTCTTTGCAGAGCACCAGAAG AATCATCTCTGGGGAGGTGCCGTTTACATA GACTTATGAATCAATCAAGGGCTTGTGCTG TAATCTTAGTTCACCCAGACATGCCCTAAG **ICCCATTTCCTACTTTAGAAACTTTGCGATT** GTTGTCTCGTTGAAAGGAAAAGGACTGTGT AGAGACGAATAAACCAGTGAGGGTAAAAGG TGTAAAAGGACATTAGCAAAACCACACGATT AATTACTTATATTCCGGCCCATCTGTGTGT CAGACTTAAATTCTCCCTTAAATTCACAAA AAACTTAAACGAACCGTGTTAATATGTCCA CTGACACTCCATCAGTAGGATGGACACTG CCCAGGGTTATAGGACAATTTTGAACCTG **ICAGTAATTGTTTGCTTCTCCTCTCCTTCA** CTCCTTACGAGTGGATGAGGCTCGTTTAT AACTCCAGGAAACTGGCAAGTCAGTCAG **ATCATTTTAGATCCATGACTCCCCTTGGA** GCCCTGCCCTAGATGAAGTAGAATC ACAGGATCTCATGAACCTGAGTCAGTGG AAGGAGAGGCCAGACAATGTGGAC CAAGGAATCCTGACTGGTAGTGGTGCT GAGGATGACCATGGGTATAATGCACAA GAGAATCCATTCCTTGCCTTTTCAACCT **AAGGGCCCCCAAGTCACTGGAATACAT** CTGAAGAGTTGTGTGCAAGGATGCTCT CTGAGCAGGGACAGAGGCTCTCATCT AGAGGCTCTCCCAGAGCACCTACAAT GAATCCCATGAGGCAGCTGAGAGAG AGCTGGAAGAAAGCAGCAGGGTCAC GGAGGCCCCTCACAGGGGAGTAGATA CACCCGGCTGTGTGTAACCTCT GACACAGCCTCCAGGTGACCCCACT CAAGACCCAGCTGTCCTCCAGGAAT **GCTCCTGCCACCATGTCCATCTAAT** CTTCTGCATCCTCTGCACCCAGCAT AATTAGAAGAGCAGTGAAACCACAGAGTCG SCTGTACATAGTGAAGGTGGGTTCTATTGC GATACTTCTCCAGCAGGGGAAACAAGAAGT TTTGGTGCAAGGTACAGATATCTCCTTATG ATAAACTAGGTTTTCTTGGAGCAGCAGGAG CTAGCCTTACCCATCCCAACTAGCCTTACC **ACCAATGTAATCACAGGGTCCTTAGAGAGG** ATTAAGATACAGACAAAAGCAGGGATGCTG **AAGTAAGGACTGAAAGGTCAGCATTCTTGC** SAAGCAAGAACAGAGGGATTTAGGATTCA AACAGAAGCTCCTGCTGATGAATAATTTCC ACTTCCTCCACTCCTGGACATCGTAGTCTC ACTCAGGCAGTTAGGGGTACACATCCTAT CTACTTCCCCTGCTTAGGAGCTTAGCCATC AGCTTTTTCTACCACAGATACCCTCACCTG GGACATGCTTGAGGGAATTTTTACTTTGTC CAACTCAGTCTCATTCCCACTCTGAGATTT SGGAACAACTTATCACCAGAGCCTAATA ATTGCACTGTGGTTATCTGGATGTTGTTTT TCTTCTCATTACAAAAGAAGCATGTTCAAA CCCAAGGTCACACAGGATGTTATATTTCTT GGTGACAAAGTGCCAGTCAGTGATTGTAT SCCATGTATGTGTGCATTTCTCTATCCATT GTTTCCCAATGCAAGTGTTTTGGTTTATTT TTTTGCTTTCCGTAAGATAACAATTTTCC AGATGGACAGGAGAGCCAAGGCCTAAGT ACTAAAGCAGTACTGGCTCCCTCCCCTCT ACACTTACCTCCATAGTGTGTGGCTGTGT ACAGCTCAAAGTGCGAGACAGGAACAC AGCCCAGAGGTTCAAGGCTACGATTAG ACCTGCCTGATTCCTACATTCCTACCAC CTTTGAGGTGAGTGGTGACTCCTGCTC **ATCCTAACTTTCAGGGGCTTTGGTCCT** ATCCCTGGTTCTGAAATCCTCAGCTTC GTCACCTTCGTGTTTTCTGCCATCAG STCAGGCTGCCTCTACACGTGAAATC SATGCTGACTCACGTGGCTTCTCTGT GAGAGGTCAGCTATCAGGCCCATCC GGGACCCTGCATTTCCGTTTCAGGT AGAACTCTTCTGGTTTCCCGCTGCT 34830 34932 34950 35145 33920 34474 34992 10395 12766 13055 16333 19972 22447 23529 25144 26130 33832 34282 34327 34384 34390 34454 34683 34741 34854 35084 35165 11741 15802 16228 16658 18295 20986 21044 21327 21541 22157 33711 1528 4095 1571 3522 6313650 6313652 6313654 6313655 6313656 6313657 6313659 6313661 6313662 6313663 6313665 6313666 6313667 6313668 6313669 6313651 6313653 6313658 6313660 6313664 6313670 6313675 6313672 6313673 6313674 6313676 6313677 6313671 CEB283 **CEB294 CEB246 CEB248 CEB249** CEB250 **CEB292** CEB254 CEB213 **CEB293 CEB225** CEB216 CEB228 **CEB284 CEB247 CEB285 CEB286 CEB287 CEB288 CEB289 CEB290** CEB251 **CEB252** CEB253 CEB224 CEB226 CEB295 **CEB222 CEB296** CEB227 **CEB297 CEB298 CEB299 CEB300** CEB301 CEB214 **CEB302** CEB303 CFB304 CEB229 CEB291 CEB201 21 21 21 21 21 21 21 21 21 21 21 21 21 22 22 22 22 22 22 21 21 21 21 22 22 22 22 22 22 22 22 22 22 22 22 22 22 22 22

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	CEB307	6313679	27382	CAGCTTCAGAATCTAAACCCCTGGTTCTAA	CTGAGCAGACCAAGGACAATAATAGAGAC	O.89	4	0.58	<u>528</u>	528	430-640
	CEB217		27398	стстесаваавсатсттсссттся	TACAGCTTCCATCTGCGTGGTGCTTTAGAC	O.89	-	0	1527	1527	1530
	CEB308	6313680	27654	AGTAGCCTCAGTAAATCGAGGAACTCTCCA	CTTAACTGACGCCTTGCTCTCTGTCGTAT	63°C	7*	0.78*	1191	1191	600-1200
22	CEB309		27773	CTCCAAAACCAAAAATCTCTATGACCCAAT	CAAAGTCACATGCTTTACCCTCAAACAAAG	58°C	-	0	513	513	513
22	CEB212	6313681	28490	CCCCAGCTGACCTTGATCACTAT	TATAGTTGGGTTTAGGCCCACCATCTGTTA	O.89	4	0.49	1201	1203	1100-1400
22	CEB202	6313682	29067	SAAC	ACTITICATCCCTCGCACCAGCTCAG	೦°09	19*	0.92*	206	606	630-1900
22	CEB230		30257	GTGTGTACGAGGCTGAGATCTAGGGATG	CGTGCCTCCACTGGTACTTGACACC	0°89	5	0.77	683	684	720-1200
	CEB231	6313683	30336	GAGTGTGCACTGAACCCATTCTTTTATCAG	GTTCTGGCTTCTGAGGGTAACTGGTTATG	O.89	3	0.45	1862	1844	1620-1850
	CEB310	6313684	30541	CCTTTTTATGGCTAAGTAGTATTCCATCGT	CGTTCAGGGAAAGAAAACAGAGATGACTT	0.89 C	12	0.85	866	866	900-1550
22	CEB311	6313685	30860	GTCTGGGTCGGTTGCAGTTGACAGTAG	CCAGCTGGATAAACTCTAAAGGTCTCAGGA	O.89	2	0.46	1115	1111	1110-1180
22	CEB232		31534	CTAAACCATTTGTCACCCTCTGGATTTGTT	CATAAGTGTGGAATTGTTGGTGGTTCTGAT	0,89	-	0	572	572	200-605
22	CEB312	6313686	31865	AGACTCTGCCAGGTGGAAATTTAAGATTGG	GCCTGATATGCCAGAGGAGATGCTTAG	O.89	2	0.04	615	615	610-640
22	CEB313		32217	ATGGTCAGCAAATAAACACCCCATGTATTT	GGCTGTTATCAGATTGTAGAGCAGAGCATC	64°C	,	0	811	811	810
22	CEB314	6313687	32267	AGAAGCTTGAAAGACAAGACTGGAGTGTCC	TCTGAGCTCTTCCCAGGTATTCCACATATT	63°C	4	0.56	1037	1036	630-1040
22	CEB203		32298	AACCACCTTCACATTTGAGAACTCGCTCTG	ACACCCAACCCATCATCCTGTCTAT	2,89	-	0	695	695	700
22	CEB315		32458	AAAGACCTCAGGGTGTAAGGACAGAAGAAA TAGGCCATCCTAAAAGGAAGGGACAGAG	TAGGCCATCCTAAAGGAAGGGACAGAG	2,89	2	0.72	1228	1245	1400-1600
22	CEB218		32693	ACCCACACAGCCTCCGTGAATTATTTAGTA	AACCTACAAGAGCACTTGGAAACCGAAG	S89	-	0	290	290	009
22	CEB316	6313688	32741	ACAAAGAGTGGACCCTGAATCACAATGAAT	AAAAATTTCCGCTGTTTAAGCTGCCTGGAC	೦.89	5	0.62	757	759	700-810
22	CEB219	6313689	32911	ACACTGAACATTGGGAAGGGCTTCTTCTAC	CCTGTGGCTTCTTCCTGTCTCAGGTAAC	2°89	က	0.51	559	559	200-700
22	CEB317		32915	ACCTTCAATATCCGTTCACCCTCTGTATCT	AAAAAGTGCTGGGAAAAGCCCTCATCT	೦.89	,	0	594	594	009
22	CEB204	6313690	32948	CAGTTCCTCAAACCCCAAGTGAAGATGA	CAGGATACTAGGGCCTAAGCAGTTTGGACA	℃ 9	5	99.0	875	298	850-1500
22	CEB318	6313691	32980	CACTATCACAAGAACAGCACGGGAAAACCT	CACATGTGAGGCTGCATGGGAAGAAC	O.89	က	0.43	777	777	730-860
22	CEB205	6313692	33057	GCGTTTAGAAAGACAGGTGCAAGAATTAGG	AGTGGTTGAGCGTCTCCTTCTGTCTCAAT	2°89	19*	0.93*	1318	1317	550-2500
22	CEB206	6313693	33318	AGAACACAGCAGCTGAAATGCCATACC	CCTAACAAAGAAGGTCAGAGTGAGCTGTG	2°89	2	0.44	1227	1226	820-1270
22	CEB233		33400	TGCTAGGAAATCCTTGTCTGCTGTGTATAA	CAGCACATGGCATTTGTTAGGATACACATA	O.89	-	0	856	856	850
22	CEB319		33414	ACTCTCCCTCCCATCCTCC	CTGCTGTTCTGCTCTCAGTTCATA	0.89°C	-	0	572	572	570
22	CEB320	6313694	33419	CTCCTCTCAGACCCTGTCACAGAACAAAC	GCTCATGAGAAAACAGTAGATCCATCTGAG	⊃.89	2	0.04	691	691	092-069
22	CEB321	6313695	33434	GAAGACCTCATCATGGGCAGTCC	ATCCTCTACTAGCCCTGATAGCACCACATC	S,€9	5	0.62	1272	1272	1240-1500
22	CEB322	6313696	33545	CCTCACTGCTCATTGGCATCAGACAT	GGTACTGCTTCTTGAGGACAGGGCTAACT	0.89	2	0.04	578	578	540-580
22	CEB220	6313697	33592	AAGACCCAAGATCTGAACCCTCAACTTCCT	GTGACTTGGCTTTCCATCTCTCTCTGT	O.89	3	0.28	785	785	700-930
22	CEB323		33618	ATGGAGAGTGGGGGCCCTTGAGTTAG	CTTTTCTTGCAACCTTAAAGGGCATCTG	၁့89	-	0	872	872	870
22	CEB207	6313698	33817	СТС	AGGAGTTTCTTAACTGTGACAGGGGATGCT	O.89	2	0.13	2617	2663	2600-3500
22	CEB324	6313699	33825	GTGGGGCAAGAGCATCTCCGTGAGT	CGCCCGGCAATAGGGGGGGGTCTTTA	2,89	19*	0.93*	985	244	400-3500
22	CEB325	6313700	33854	GCCCCTTCCTCTGGTTCCATACTG	CTCTGCCTTCAGAACCCCATACCTCT	58°C	4	0.53	926	928	930-1400
22	CEB221	6313701	33864	CAAAATAATTGGAGTAGGAGTGGGTGAAGC	AAGTGGTTTTGCACCCAAATCATTAGAAGA	J.89	က	0.51	802	989	750-830
22	CEB326	6313702	33965	AAAGCAGAAAGTGCATCTGGAAAACAAAG	TAAAGATCCTTGCATGTTTTCTGAGGGATG	0°89	က	0.25	1408	802	1250-1680
	CEB327		33983	CAGGAGGCGCGTGGGACTACACTT	GGCCGCCTTCCTCCACTCTCACCT	0.89 C	-	0	793	793	800
22	CEB223	6313703	34031	AATACCAGCACGATCGATTCTATCAGGACA CCCTGTGGAGACAGTGTTTTGTGATG	CCCTGTGGAGACAGTGTTTTGTGATG	O.89	2	0.5	1836	1864	1850-1950

Table 1. (Continued)



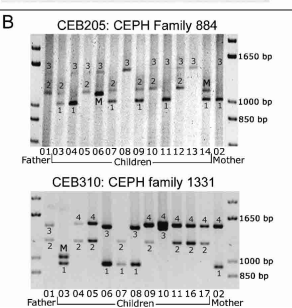


Figure 2 (*A*) Ethidium bromide–stained agarose gel showing PCR products for minisatellite CEB285. Six different alleles are scored among 32 individuals (in some cases, three bands are seen for one individual [the upper one is a PCR artifact as shown by segregation patterns in families]; this artifact occurs only in heterozygotes [data not shown], indicating a mechanism involving an interaction between the two alleles). (*B*) Image of the gels obtained for minisatellites CEB205 and CEB310 on CEPH families 884 and 1331, respectively. Two children inherit mutant alleles for CEB205, and one child inherits a mutant allele for CEB310. For CEB205, larger alleles are missed in the procedure used: The results were confirmed by Southern blot.

should be monomorphic, and ~75% (32 of 43) should have a heterozygosity value \geq 0.5.

Heterozygosity and allele number are significantly higher in the positive group for criterion 4 (over the entire set of typed repeats) compared with the negative group (Fig. 4B). Criterion 4 produces an enrichment of repeats having heterozygosity \geq 0.5 from 42% (49 of 118) in the whole set to 61% (25 of 41) in the positive group and a diminishment of monomorphic repeats from 25% (30 of 118) in the whole set to 12% (5 of 41) in the positive group. Criterion 4 thus reduces to nearly one third (116 to 41) the number of minisatellites to type while eliminating most monomorphic minisatellites and retaining 50% of the most polymorphic ones. By comparison, criterion 3, if applied to the entire set of typed repeats, (Fig. 4B) would reduce their number by roughly half (118 to 61), eliminating just two fewer monomorphs while retaining 69%

(34 of 49) of the most polymorphic repeats. Additionally, criterion 4 eliminates half (four of eight) of the highly polymorphic (heterozygosity \geq 0.85) minisatellites, whereas criterion 3 retains 75% (six of eight) of these.

We note that for some highly polymorphic minisatellites, (CEB202, CEB205, CEB310, CEB291), predicted lengths are identical in the two sequences. In addition, the results for criterion 4 are not uniform for the two chromosomes, owing to the much greater agreement on predicted loci length in chromosome 22. We presume that this reflects the fact that the Celera sequence was assembled by using both public and Celera sequence reads (Venter et al. 2001). More surprisingly, for five minisatellites, which we found to be monomorphic, predicted lengths differ (CEB214, CEB255, CEB264, CEB247, CEB289). These findings raise unresolved questions about the accuracy of the HGP and Celera sequences with regard to minisatellites. Tandem arrays can present significant sequence assembly problems, in particular when the internal array contains regions of high homology and, potentially more seriously, when the repeat exhibits length polymorphism and data are drawn from more than one individual, as was done for the Celera sequence (Venter et al. 2001).

To examine this further, we compared the HGP and Celera predictions to the alleles we detected (in Table 1, predicted lengths are underlined and not shaded when they correspond to an observed allele). In 65% (75 of 116) of the repeats, HGP and Celera predict an identical allele length, which corresponds to an observed allele length with five exceptions (Table 2) and is the most common allele in 81% of these cases. In 35% of the repeats (41 of 116), HGP and Celera predict different length alleles (Table 2). The length predicted by the HGP sequence fits with an observed allele size in 36 cases (most common allele length in 20 of these), whereas the Celera prediction fits with an observed size in 10 cases (and was once the most common allele).

Among the tandem repeats that provide PCR products unmatched by the HGP sequence, six sufficiently informative ones (CEB230, CEB253, CEB295, CEB298, CEB315, CEB269), with at least three different alleles among the four parental chromosomes, were typed in large CEPH families to check their chromosomal origin. All map to the expected area of chromosome 21 or 22, indicating that the discrepancy between sequence data and PCR product size probably results from a sequencing error (or the sequencing of a very rare allele) and not from a PCR specificity problem.

 χ^2 tests were used to examine whether the similarities in prediction of the HGP and Celera findings could be explained by chance (see Methods). Differences identified by the tests had, in all cases, less than one one-thousandth probability of occurring by chance. Specifically, cases in which predictions disagreed and both allele sizes were detected were underrepresented (compared to expected frequency) in all tests, and cases in which only one or neither predicted size was detected were overrepresented in all but one test.

Identifying Hypermutable Loci

Hypermutable minisatellites are expected to belong to the class of highly polymorphic loci because they are, by definition, subject to frequent rearrangements that generate new alleles. For practical reasons linked to the size of available pedigrees, a minisatellite will usually be classified as hypermutable if its average mutation rate in the germline is >0.5%, that is, if an average of at least one or two mutant alleles is observed among 100 children.

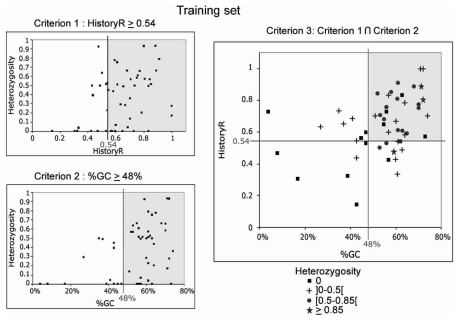


Figure 3 Criteria 1, 2, and 3 applied to the training set. For criteria 1 and 2, heterozygosity (28 individuals) versus HistoryR (criterion 1) or percentage of GC (%GC; criterion 2) are plotted. Correlations are significant at the 0.01 level. For criterion 3, HistoryR versus %GC is plotted, with different symbols representing the polymorphism. Lines represent the selected thresholds, and shaded areas contain the minisatellites selected by the criteria (criterion 1, HistoryR ≥0.54; criterion 2, %GC ≥48%; criterion 3, criteria 1 and 2 combined). Plots show that criteria select most of the polymorphic minisatellites and eliminate a majority of monomorphs or slightly polymorphic ones.

We typed the eight most polymorphic minisatellites (i.e., with heterozygosity ≥0.85) in the eight largest CEPH families (102 children) to search for mutant alleles. Comparing the results obtained by PCR and Southern blotting shows that even when some larger alleles are missing in the PCR products, the estimated heterozygosity rate (see Methods) is close to the heterozygosity rate obtained with Southern blots. This helps validate the simplified PCR-based polymorphism measurement. Among the eight minisatellites (CEB202, CEB205, CEB250, CEB310, CEB269, CEB291, CEB305, CEB324), two showed mutant alleles (CEB205 and CEB310; Fig. 2B). Both yielded two mutant alleles among 204 meioses, that is, 102 children (mutation rate, 0.12% to 3.5%; 95% confidence interval). For minisatellite CEB205, one mutation event occurred in the mother and the other in the father, whereas for CEB310, both mutations occurred in the father. The remaining six minisatellites yielded no mutant allele among 102 children (mutation rate, 0 to 1.79%; 95% confidence interval). They were not investigated further but can not be strictly excluded from being hypermutable. The two minisatellites that appeared hypermutable among 102 children were then typed in more families (32 other reference CEPH families). For CEB205, one new mutant allele was found among 352 meioses (mutation rate, 0.54%; 95% confidence interval, 0.11% to 1.57%), but no other mutant allele was detected for CEB310 among 476 additional meioses (mutation rate, 0.29%; 95% confidence interval, 0.04% to 1.06%). Based on these results, CEB205 appears to be hypermutable. It is a GC-rich minisatellite with a unit length of 33 bp repeated 10 to 70 times, located at 1.5 Mb from the end of the chromosome 22 sequence. It seems to be part of a predicted coding region (gene LOC129238; see http://www.ncbi.nlm.nih.gov/ LocusLink/LocRpt.cgi?l=129238, 31 July 2002 update).

DISCUSSION

This study, performed on the scale of entire human chromosomes, provides a first global evaluation of minisatellite polymorphism based on genome sequence data. The repeats studied here, chosen by using a detailed definition that is more stringent than the broad definition mentioned in the Introduction, are, in majority (75%), polymorphic in the population investigated, and 42% have a heterozygosity value ≥0.5. Minisatellites from chromosomes 21 and 22 are similar in physical distribution (higher frequency toward chromosome ends), sequence features, and polymorphism. Assuming that chromosomes 21 and 22 are representative of all human chromosomes and given that the two chromosomes represent ~2% of the genome, we speculate that the entire human genome contains ~6,000 minisatellites that match our definition, including 4,800 polymorphic and 2,500 very polymorphic ones. A few 10s of these might be expected to qualify as hypermutable loci. Be-

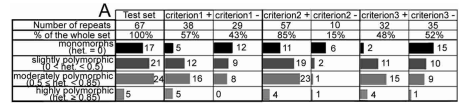
cause our definition precluded many other potentially polymorphic minisatellites, future research should seek to expand the category of minisatellites that are tested against our polymorphism prediction criteria.

Predicting Polymorphism

We showed that using the sequence properties %GC and HistoryR effectively improves polymorphic minisatellite selection. With them, we reduce the number of minisatellites for typing by about half while increasing the frequency of repeats with heterozygosity $\geq 0.5\%$ from the background rate of 43% to 59%. Internal conservation, used as a polymorphism predictor for microsatellites, is not applicable to minisatellites, presumably owing to the greater complexity of their mutation processes.

That %GC correlates with polymorphism is in agreement with earlier observations. Some of the first minisatellites to be characterized were detected via a shared 10- to 15-bp "core" sequence similar to the generalized recombination signal (χ) of *Escherichia coli* (GCTGTGG; Jeffreys et al. 1985). The majority of classical minisatellites (mostly polymorphic and/or hypermutable ones) are GC-rich, with a strong purine/pyrimidine strand asymmetry (Vergnaud and Denoeud 2000). In other genomes, though, (for instance bacterial genomes), %GC does not seem to be associated with minisatellites polymorphism (Le Fleche et al. 2001). Such a criterion may therefore not be universal, especially because GC content varies significantly across genomes.

The HistoryR criterion is based on the hypothesis that tandem repeats expand through multiple rounds of duplication, with the new copies sharing the mutations that occur before duplication, whereas unique mutations accumulate once the repeat is no longer evolving. For example (Fig. 1),



D					
D	Whole set	criterion3 +	criterion3 -	criterion4 +	criterion4 -
Number of repeats	118	61	57	41	75
% of the whole set	100%	52%	48%	35%	65%
monomorphs (het. = 0)	30	7	23	5	25
slightly polymorphic (0 < het. < 0.5)	39	20	19	11	27
moderately polymorphic (0.5 ≤ het. < 0.85)	41	28	13	21	19
highly polymorphic (het. ≥ 0.85)	8	6	2	4	4

Figure 4 (A) Application of criteria 1 (HistoryR ≥0.54), 2 (%GC ≥48%), and 3 (HistoryR ≥0.54 and %GC ≥48%) to the test set. For each criterion, the distributions of minisatellites (from monomorphs to highly polymorphic) between positive (retained by the criterion) and negative (excluded by the criterion) sets are compared. All differences between sets + and - are statistically significant at the 0.01 level. (B) On the whole set, comparison of the results obtained with criterion 4 and criterion 3.

minisatellite CEB252 shows several redundant patterns of mutation, resulting in a high HistoryR score, whereas CEB233 shows no clear organization of mutations, resulting in a low HistorvR score.

This polymorphism criterion is likely to be applicable to any genome, even though the history reconstruction algorithm makes simplifying assumptions about the possible biological mechanisms involved in array expansion. These mechanisms, which include mutational events during mitotic replication and meiotic recombination, comprising both intraallelic and interallelic events, might occur independently or jointly. At present, there are no rules to predict which mechanism will occur preferentially at which locus (Maleki et al. 2002). Moreover, the individual mechanisms themselves are still poorly understood and, thus, impossible to model. Meiotic events, for instance, have been shown to result from the activity of nearby meiosis-specific double-strand break hot-spots. The nature of these sites, better known in yeast, is still unknown in the human genome (Debrauwère et al. 1999; Tamaki et al. 1999; Vergnaud and Denoeud 2000). In view of the current state of knowledge, it may be premature to hope for a perfect polymorphism predictor based on apparent array expansion.

Use of Two Human Sequences to Select for Polymorphic Loci Is Problematic

The availability of two versions of the human genome sequence provides an additional avenue to improve polymorphic minisatellite identification. However, in the repeats studied here, selection based on reported length differences discarded half the highly polymorphic minisatellites and, in particular, the hypermutable one from chromosome 22. In both chromosomes, the number of loci with different predicted lengths in the HGP and Celera sequences that were nonetheless both found was significantly underrepresented. This is apparently owing to the lack of independence resulting from sharing of data during assembly of the Celera sequence. In addition, in both chromosomes, the number of loci in which only one or no predicted allele was found is overrepresented, apparently owing to assembly errors. Because the Celera sequence—which when not in agreement with the

HGP data—usually provides copy numbers unobserved in any allele, it appears that the Celera sequence, at least with respect to minisatellites, is more prone to assembly error. As a result of the lack of independence/assembly errors in the Celera/HGP data, polymorphism prediction based on sequence comparison did not perform as well as anticipated.

One New Hypermutable Locus in a Coding Region

This study revealed one hypermutable minisatellite, CEB205, showing three mutant alleles among 278 children (mutation rate, 0.54%; 95% confidence interval, 0.11% to 1.57%). Interestingly, CEB205, with a 33-bp pattern, may be part of a coding region. The corresponding putative protein is 614 amino acids

long, half of which are derived from the tandem repeat (11 codon repetition) at the N terminus. Of the minisatellites studied here, 26 among 60 (43%) on chromosome 21, and 22 among 67 (33%) on chromosome 22 belong to genes (i.e., exons, introns, or UTRs), as determined by sequence similarity analyses in the human genome sequence (using BLAST and http://www.ncbi.nlm.nih.gov/genome/seq/, release of November 2001). None except CEB205 appear to contribute to the coding sequence itself. Although the proportion of tandem repeats that contribute to coding regions is important in bacterial genomes, it is relatively low in the human genome, and CEB205 might represent the first known, coding hypermutable minisatellite.

CEB310—which exhibited meiotic mutation events, but which we do not here classify as hypermutable—is unusual in that its sequence is 80% AT. It is reminiscent of the tandem repeats studied in Giraudeau et al. (1999), that is, minisatellites made up of degenerated microsatellite-like repeated units (in this case, [AC]m[AT]n). Although most hypermutable minisatellites known to date are GC-rich, some have been described as having a very high AT content, for instance, the one constituting the chromosomal fragile site FRA16B (Yu et al. 1997; Yamauchi et al. 2000). The highly polymorphic minisatellite MSY1, from human chromosome Y, is also very AT-rich (75% to 80%; Jobling et al. 1998).

Future research will expand the systematic exploration of human tandem repeat polymorphism by testing the %GC, HistoryR, and HGP/Celera criteria on other human chromosomes as the sequences are progressively finished and released (Deloukas et al. 2001).

METHODS

Constructing the Tandem Repeats Database

Tandem repeats were identified from chromosome 21 (Hattori et al. 2000) and chromosome 22 (Dunham et al. 1999) sequences by using the TRF software (Benson 1999) with the following options: alignment parameters of (2,3,5), minimum alignment score to report repeat of 50, maximum period size of 500. When the program reported redundant (overlapping) repeats, the redundancy was eliminated in the following way. For each group of overlapping repeats, two values were determined: Lmax, the maximal total length among the redundant alignments, and Mmax, the maximal percent matches among the redundant alignments with total length ≥80% of Lmax. Then, of all the alignments in the group with total length ≥80% of Lmax and percentage of matches ≥Mmax – 0.1, the one with smallest unit length was stored in the database. The nominal length of the stored repeat is the total length of the overlapping region, that is, from the first position of the first overlapping repeat to the last position of the last overlapping repeat. Twenty-two tandem repeats showed differences of >5% between the nominal length and the length of the stored repeat, (the difference exceeded 10% in 14 cases, and 30% in three cases: CEB311, 33%; CEB320, 46%; and CEB327, 50%). For these latter three, TRF cut the repeats into two parts, which were combined for further analysis. Variation between nominal and stored size of repeats does not affect allele size prediction, which is based on length of sequence between primers. The database, publicly available at http://minisatellites.u-psud.fr, can be queried according to a number of simple features (e.g., total length, unit length, copy number, %GC) and provides links to repeat alignments and flanking sequence data as described previously (Le Fleche et al. 2001).

PCR Typing of Minisatellites

DNA was provided by Centre d'Etudes du Polymorphisme Humain (CEPH; http://www.cephb.fr/). PCRs were performed in 15 µL reactions, using 50 ng of genomic DNA, Roche long template PCR buffer (1.75 mM MgCl₂, 50 mM Tris-HCl at pH 9.2 and 25°C, 16 mM [NH₄]₂[SO₄]), 0.033 U/μL Taq polymerase (Roche), 0.003 U/µL Pwo (Roche), 200 µM of each dNTP (Amersham-Pharmacia biotech), and 0.6 µM of each flanking primer (Table 1; primers were selected within the flanking sequences provided by TRF using Primer3 software: http:// www-genome.wi.mit.edu/cgi-bin/primer/primer3_www.cgi). PCRs were cycled for 5 min at 96°C, then for 15 sec at 96°C: for 20 sec at annealing temperature (Table 1; this temperature was optimized for each primer pair by using the temperature gradient provided by MJResearch PTC200), for 5 min at 68°C for 30 cycles, and for 10 min at 68°C, on Perkin Elmer 9600 thermocycler or MJResearch PTC200. Samples were run through a 13-cm-long 1% standard agarose (Qbiogen) gel in 0.5 × TBE buffer at 10 V/cm for 1.5 h and visualized by ethidium bromide staining using UV (1 \times TBE buffer is 89 mM Tris, 89 mM boric acid, 2 mM EDTA at pH 8).

Polymorphism Measures

A population of 96 CEPH individuals (from the 40 reference families) were typed for minisatellite polymorphism. This population includes 13 mother/father/child trios and altogether comprises 76 unrelated individuals. The 76 unrelated individuals form subpopulation 1. A subset of 28 unrelated individuals forms subpopulation 2. The exact list of the 96 individuals typed is provided in Supplementary Table 2.

In this study, we examined only length polymorphism, not internal sequence variation. Two values, calculated on unrelated individuals, were used to quantify polymorphism: the number of alleles observed and the heterozygosity, calculated as $1 - \Sigma f^2$, where f are the allelic frequencies observed in the population of unrelated individuals. Heterozygosity represents the probability of having two different alleles. The simple PCR and ethidium bromide staining assay used here will usually detect only the smallest allele in individuals showing large length differences between alleles (as is often the case for highly polymorphic loci). The shorter allele often masks the longer one because it is easier to amplify. Such PCR artifacts are indicated with an asterisk in Table 1. They were detected because of the mother/father/child segregation controls and also because they do not satisfy the Hardy-Weinberg equilibrium, as tested with the HWE program, from the publicly available Linkage Utilities package (Ott 1999). For these loci, the heterozygosity value calculated from allelic frequencies was obtained by counting only one allele for individuals showing a single band (i.e., by assuming that the individual is heterozygous with one allele masked) instead of counting the same allele twice, as was done for loci in which homozygosity was not in question. The resulting heterozygosity value could be underestimated (if too many alleles are not seen), but it is sufficient to roughly evaluate the polymorphism.

Mutation Rate Estimation

Mutation rate of the most polymorphic (i.e., potentially hypermutable) minisatellites was evaluated by a combination of Southern blot hybridization and PCR typings, in recognition of the "masking" phenomenon described above. Typings were performed by using DNA from the eight largest CEPH families (F102, F884, F1331, F1332, F1347, F1362, F1413, F1416). Five µg of DNA were digested with AluI (CEB202, CEB250, CEB269, CEB291) or HinfI (CEB205, CEB324, CEB305; Boehringer Mannheim), electrophoresed through a 1% agarose gel and transferred to nylon membranes (Nytran+, Schleicher and Schuell) under vacuum (Pharmacia Biotech). Probes were obtained from PCR products and recovered from agarose using QIAquick gel extraction kit (Qiagen). Probes were labeled with a-[³²P]dCTP (Amersham Pharmacia Biotech) by the random priming procedure (Feinberg and Vogelstein 1984). Hybridization was conducted as described in Vergnaud (1989) in an hybridization oven at 65°C. After hybridization, the filters were washed in 1× SSC/0.1% SDS or 0.1× SSC/0.1% SDS at 65°C. Membranes were revealed by using a phosphoimager (Storm 860 Molecular Dynamics).

Sequence Characteristics of Repeats

The following sequence characteristics (calculated from the HGP sequence) were tested for correlation with either allele number or heterozygosity. Characteristics did not differ markedly when evaluated in the Celera sequence (in which differences with HGP typically involved deletion of adjacent copies reported in the HGP sequence):

1. Unit length: the length of the repetitive unit (consensus pattern).

Table 2. Success of the Public Human Genome Project (HGP) and Celera Sequences in Predicting Alleles That Were Actually Found to Occur									
116 tandem repeats (predicted by both sequences)									
Predicted length	s match: n = 75		Predicted length	ns differ: n = 41					
Allele found with	oredicted length:		Allele found with	predicted length:					
Yes	No	Yes, both predictions	HGP prediction only	Celera prediction only	Neither prediction				
70	5	10	26	0	5				

- 2. Copy number: the number of copies of the repetitive unit.
- 3. Total length: the length of the entire tandem array.
- Percent matches: the frequency at which a nucleotide at a
 position in one unit matches the corresponding nucleotide
 in the next unit (reading from left to right).
- 5. %GC: the percentage of nucleotides that are either G or C.
- 6. GC bias: strand asymmetry for G and C, |%G %C|/ (%G + %C).
- 7. Purine/Pyrimidine bias: strand asymmetry for purines and pyrimidines, |%Pur %Pyr|/(%Pur + %Pyr).
- 8. Average entropy: from the columns of a multiple alignment of the repeat copies, the average, over all columns, of the entropy calculated from nucleotide frequencies.
- 9. HistoryR: described below.

HistoryR is derived from the tandem repeats history reconstruction algorithm (Benson and Dong 1999), a greedy algorithm that chooses a series of least-cost contractions to convert a multicopy tandem array into a single putative ancestral copy. Greedy algorithms are not guaranteed to find the overall least-cost solution, but testing has shown this approach to work very well on simulated sequences. Input is a multiple alignment, M, of the individual copies in the repeat, with \mathbf{n} rows (number of copies) and \mathbf{k} columns (length of alignment). $M_{i,j}$ represents the **i**th row and **j**th column of M, and each $M_{i,j}$ contains one of the alphabet symbols (A,C,G,T,-). In a contraction, two or more consecutive, equallength subsequences (the contraction copies) are replaced by a single subsequence (the merged copy) of the same length (all subsequences selected have length equal to a multiple of \boldsymbol{k}). Each contraction reduces the number of rows in \boldsymbol{M} . If the contraction copies are identical, then one becomes the merged copy. Otherwise at every position at which the contraction copies differ, the merged copy contains the character that occurs most often, with ties being represented by an ambiguous character, that is, a set of all the most frequently occurring characters at that position. An ambiguous character created in one contraction may be converted to a single character in a subsequent contraction. This method is analogous to that used by Sankoff (1975; Sankoff and Rousseau 1975). The cost of a contraction is a ratio. The numerator is the cost of obtaining the contraction copies from the merged copy; that is, at each position of the merged copy, subtract the number of times the most frequent character occurs in the contraction copies from the total number of contraction copies, then sum all these differences. The denominator is the combined length of rows by which M is reduced, that is, the length of all contraction copies minus the length of the merged copy.

History reconstruction yields four numerical values: (1) Max, the maximum possible history cost; (2) Min, the minimum possible history cost; (3) BinaryActual, the calculated history cost when the number of contraction copies in every contraction is restricted to exactly two; and (4) ManyActual, the calculated cost when the number of contraction copies is unrestricted. Max and Min are sums of column values from the original alignment M. In the case of Max, the value of a single column is the number of characters that are not the most frequent character. Max is therefore the cost if the most frequent character is ancestral and if every character different from the ancestral character was produced by its own mutation. For Min, the value is one less than the number of distinct characters in a column, that is, at most four. Min is the history cost if every distinct character different from the ancestral character arose by a single mutation (with identical characters produced by duplication).

Combinations of the four numerical values were tested for polymorphism prediction in the training set and HistoryR, which produced the highest correlation with heterozygosity, was used for the remainder of the study. It is defined as

$$\textit{History R} = \begin{cases} (Max - \textit{BestActual})/(Max - Min) \text{ when } Max \neq Min \\ 1 \text{ otherwise} \end{cases}$$

where BestActual is the minimum of BinaryActual and ManyActual. Usually, this was BinaryActual. The HistoryR value can be thought of as the proportion of mutations that could be accounted for by duplication that actually are. When Max \neq Min, HistoryR \leq 1, with a higher ratio indicating more mutations accounted for by duplications (Fig. 1). When Max = Min, each mutation is unique, and we arbitrarily set the ratio to one. This occurred in only one repeat with a total of four mutations. The history reconstruction program is freely available for interactive use at http://tandem.biomath.mssm.edu/cgi-bin/history/history.exe.

Statistical Analysis

All statistical analysis was done with the SPSS program except for χ^2 tests which were done with StatXact 4. Correlations were determined by three methods: Pearson correlation, and nonparametric Kendall's τ_b and Spearman's ρ. Correlations are considered significant at the 0.01 level (two-tailed) of the test statistics. Group comparisons were determined by first conducting two tests of normality, Kolmogorov-Smirnov and Shapiro-Wilkinson, on the values within each group. Values were assumed to be normally distributed unless the test statistic fell within the 0.05 level of significance. If the values were normally distributed in the two groups, then a *t* test was used to compare the means, which were judged significantly different at the 0.01 level of the statistic (two-tailed). If the values were not normally distributed in either of the two groups, then a nonparametric Mann-Whitney test was used to compare the distributions, which were judged significantly different at the 0.01 level of the statistic (two-tailed).

 χ^2 tests were used to analyze HGP/Celera prediction data for chromosomes 21 and 22 separately. The data were divided into three categories: (1) identical predictions/allele size detected, (2) different predictions/both alleles sizes detected, and (3) one or neither predicted allele size detected. Two estimates for frequency of unobserved alleles were used (in order to calculate the probability of alleles being detected): 10% which corresponds to the largest frequency in the population for which the chance of not appearing in our sample of 28 individuals is \geq 0.05, and an arbitrary low estimate of 1%. The probability of identical predictions in the HGP and Celera sequences was obtained by summing the estimated heterozygosity values calculated separately for each locus based on observed frequencies in our sample (equivalent to using the average observed heterozygosity over all loci).

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