Analysis of Primate Genomic Variation Reveals a Repeat-Driven Expansion of the Human Genome

Ge Liu,¹ NISC Comparative Sequencing Program,² Shaying Zhao,³ Jeffrey A. Bailey,¹ S. Cenk Sahinalp,¹ Can Alkan,¹ Eray Tuzun,¹ Eric D. Green,² and Evan E. Eichler^{1,4}

¹Department of Genetics, Center for Human Genetics and Center for Computational Genomics, Case Western Reserve University School of Medicine and University Hospitals of Cleveland, Cleveland, Ohio 44106, USA; ²Genome Technology Branch and NIH Intramural Sequencing Center, National Human Genome Research Institute, Bethesda, Maryland 20892, USA; ³The Institute for Genome Research, Rockville, Maryland 20850, USA

We performed a detailed analysis of both single-nucleotide and large insertion/deletion events based on large-scale comparison of 10.6 Mb of genomic sequence from lemur, baboon, and chimpanzee to human. Using a human genomic reference, optimal global alignments were constructed from large (>50-kb) genomic sequence clones. These alignments were examined for the pattern, frequency, and nature of mutational events. Whereas rates of single-nucleotide substitution remain relatively constant ($1-2 \times 10^{-9}$ substitutions/site/year), rates of retrotransposition vary radically among different primate lineages. These differences have lead to a 15%–20% expansion of human genome size over the last 50 million years of primate evolution, 90% of it due to new retroposon insertions. Orthologous comparisons with the chimpanzee suggest that the human genome continues to significantly expand due to shifts in retrotransposition activity. Assuming that the primate genome sequence we have sampled is representative, we estimate that human euchromatin has expanded 30 Mb and 550 Mb compared to the primate genomes of chimpanzee and lemur, respectively.

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

Initial studies of primate genome variation were based largely on indirect evidence obtained from DNA hybridization kinetics (Sibley and Ahlquist 1984; Powell and Caccone 1990; Sibley et al. 1990). Molecular studies have been limited mainly by the lack of large-scale DNA sequence data (Bailey et al. 1991; Smith 1992; Horai et al. 1995; Kaessmann et al. 1999; Bohossian et al. 2000; Chen and Li 2001). In the past, such large-scale comparisons were dependent upon PCR crossamplification among diverse primate taxa and, therefore, were biased to either conserved regions or limited to closely related species. With the anticipated completion of the human genome sequence (International Human Genome Sequencing Consortium 2001; Venter et al. 2001) and the development of primate BAC library resources (Eichler and De-Jong 2002), it is now possible to initiate large-scale genomic comparisons (Thomas et al. 2002) in an unbiased fashion to assess the nature and pattern of primate genomic variation. Direct comparison of high-quality finished sequence from BAC clones of orthologous loci will not only elucidate mechanisms of genome evolution, but also shed light into the historical events that have shaped our species.

A variety of mutational forces are thought to have molded the human genome. These include both small-scale (single-base pair changes, microsatellite slippage, insertion/ deletions) as well as large-scale events (retrotransposition, genomic rearrangements, segmental duplication). To date, most

⁴Corresponding author.

E-MAIL eee@po.cwru.edu, FAX (216) 368-3432.

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evolution studies have focused on either single-base pair changes or microsatellite evolution (Chen et al. 2001; Ebersberger et al. 2002; Smith et al. 2002; Webster et al. 2002). Estimating rates of retrotransposition has been difficult in part due to the paucity of such de novo events over short stretches of DNA sequence as well as biases in repeat classification and genomic insertion sites (Chiaromonte et al. 2001; Batzer and Deininger 2002). Over the last 60 million years of evolution, the human genome has been bombarded by a variety of repeat elements through successive waves of retrotransposition (Smit 1999; Deininger and Batzer 2002). Among these, L1 (long interspersed repeat element 1) and Alu (a short interspersed repeat) elements are most prevalent (Moran et al. 1999; Batzer and Deininger 2002). Combined, they account for an estimated 26%-27% of the human genome (International Human Genome Sequencing Consortium 2001; Venter et al. 2001).

In this study, we analyzed genomic sequence from three species (chimpanzee, baboon, and lemur) and compared it to the human genome. These three species are estimated to have diverged from human at three very different time points, approximately 5.5, 25, and 55 million years ago (Goodman 1999). This analysis therefore provides a snapshot of genomic change representative of the evolutionary depth of the primate order. Most of the sequence was generated by the NIH Intramural Sequencing Center (http://www.nisc.nih.gov/) and represents orthologous regions to human chromosome 7. As part of this study, we generated large-scale alignments (ranging in length from 50–150 kb), providing a baseline for our analysis of genomic variation. The objective was to assess

RESULTS AND DISCUSSION

Alignment Validation

One of the most significant challenges to large-scale genomic analyses is the generation of biologically meaningful global alignments (Chen et al. 2001). A total of 10.6 Mb of aligned sequence between human and nonhuman primates were analyzed which included human orthologous comparisons with 51 chimpanzee, 42 baboon, and 9 lemur BAC clones or subclones. For each species, we chose a subset of gap opening and gap extension penalties which minimized the frequency of both single-nucleotide and insertion/deletion events. An assessment of both types of variation simultaneously, we reasoned, should provide the most biologically meaningful optimal global alignment (Methods). In order to validate the reliability of our alignment parameters, a number of tests were performed. First, we analyzed the nature of the sequence underlying insertion/deletions within each alignment (Methods). A variety of biological events are known to create insertion/deletions, including lineage-specific amplification of tandem repeats, homology-mediated genomic deletions, and retrotransposition events. Alignment parameters were favored where such large-scale insertion/deletions were effectively treated as a single event. All individual alignments and patterns of single-nucleotide variation were manually inspected and are available online (http://eichlerlab.cwru.edu/ primategenome/).

As a second test, we compared overall estimates of sequence divergence (Table 1) with previous reports in the literature (Li 1997; Chen and Li 2001; Chen et al. 2001; Ebersberger et al. 2002; Fujiyama et al. 2002). These studies are particularly relevant for human-chimpanzee alignments where similar sequence comparison studies using different alignment parameters have been performed. Although our results for human and chimpanzee divergence (K = 1.14%) are comparable to those of previous studies (1.18%-1.24%; Chen and Li 2001; Chen et al. 2001; Fujiyama et al. 2002; Smith et al. 2002), our estimate is lower. In our study, we excluded regions that harbored large, low-copy repeat sequences, as the orthologous relationship of these could not be unambiguously determined. Such segmental duplicated regions may significantly inflate estimates of divergence due to nonorthologous sequence relationships (Chen et al. 2001; Bailey et al. 2002) or gene conversion (Hurles 2001). Because comparable sets of data do not exist for other nonhuman primates, we generated 1000 randomly selected end sequences from existing BAC libraries (Eichler and DeJong 2002) for each species. Comparing high-quality alignments of BAC end sequence with these optimal global alignments, we observed similar estimates of sequence divergence between humanlemur (20%-21%), human-baboon (5%-6%), and humanchimpanzee (1%-1.2%). The variation distribution pattern of these short alignments (400-500 bp; data not shown) was remarkably similar to the distribution observed for nonoverlapping 500-bp windows generated from chromosome 7 optimal global alignments (Fig. 1, Suppl. Fig. S4).

Single-Nucleotide Variation

Based on estimated divergence times from the human lineage (Goodman 1999), we calculated the substitution rate for each species comparison (Table 1). Estimates of overall single-base

pair substitution rate ranged from 1.0×10^{-9} mutations/site/ year for human–chimpanzee comparisons to 2.1 imes 10⁻⁹ mutations/site/year for human-lemur comparisons (Table 1). It has been suggested that the rate of substitution has slowed by as much as 50% among hominoids (humans and apes) after their separation from the Old World monkey lineage (Goodman et al. 1971; Koop et al. 1986; Li and Tanimura 1987). Indeed, a noticeably higher substitution rate was calculated based on human-baboon sequence alignments compared to estimates from human-chimpanzee alignments (Table 1). This effect becomes more dramatic when CpG dinucleotide sites are excluded. Human-lemur sequence comparisons indicated that the most dramatic change in the rate of substitution occurred early in primate evolution (25-55 million years ago), possibly owing to generation-time differences among prosimian and simian lineages (Ruvolo 1997). Although significant differences in the mean genetic distance were observed between human and nonhuman primates, the variance of these estimates was not constant. When we analyzed nonoverlapping 3-kb blocks of aligned genomic sequence, a considerable increase in genomic variance in sequence identity was observed as a function of evolutionary divergence (Fig. 1).

We performed a substitution relative rate test for all instances where three or more homologous sequences were available. We constructed 19 multiple alignments for human, chimpanzee, and baboon (2.5 Mb) and five multiple alignments for all four species (0.51 Mb). Relative rate tests were performed using Tajima's test (Kumar et al. 2001). Seventeen of the 19 rate tests supported the molecular clock hypothesis for human, chimpanzee, and baboon alignments. Similarly, when lemur was used as an outgroup, human and chimpanzee were found to have nearly identical substitution rates. In contrast, when using lemur as an outgroup, both human and chimpanzee had slower rates of substitution compared to baboon. Therefore, a local molecular clock seems to hold well between human and chimpanzee (Zuckerkandl and Pauling 1965). It is worth noting that the data used in these studies were limited to four species, and rate calculations may be confounded by incorrect estimates of species divergence times. However, even if more distant divergence times are used, the data clearly indicate that substitution rate has at least doubled among prosimians compared to the haplorhime species.

Retrotransposition

Retrotransposition typically creates large sequence insertions ranging in size from a few 100 bp to ~10 kb in length. Three major classes of retroelements have shaped the primate genome in recent evolutionary history: L1 (LINE), Alu (SINE), and LTR (long terminal repeat elements of endogenous retroviruses; International Human Genome Sequencing Consortium 2001). We examined all insertion/deletion (indel) events in excess of 100 bp within baboon and chimpanzee/human alignments in order to identify new insertions that had occurred over the last 25 million years of evolution. An indel was classified as a retrotransposition event if at least 80% of the indel contained one predominant repeat. We considered the known interspersed repeat phylogeny based on the established repeat subclasses as reported (Smit 1999). All insertions were considered, including the ancient repeat subclasses that passed our test. Further, in the case of L1 and Alu repeats, insertion sequences were examined for the presence of target-

Table 1. Primate Sir	ndle-Nu	cleotide Variati	ion versus Sequ	uence Class							
	# loci	Alignment length (bp)	Aligned bases (bp)	Matches (bp)	Mismatches (bp)	Transitions (s)	Transversions (v)	s/v	ldentity (%)	Kimura Distance (%)	$R^a \times 10^{-9}$
Human–Chimpanzee Overall	51	4968069	4853708	4798947	54761	36914	17847	2.07	98.87 ± 0.00	1.14 ± 0.00 ^b	1.034 ± 0.004
Overall-CG	51	4968069	4764283	4723249	41034	15414	25620	1.66	99.14 ± 0.00	0.87 ± 0.00	0.788 ± 0.004
Exon	24	69051	68957	68543	414	296	118	2.51	99.40 ± 0.03	0.60 ± 0.03	0.548 ± 0.027
Unique noncoding	51	2749584	2720023	2692593	27430	8913	18517	2.08	98.99 ± 0.01	1.02 ± 0.01	0.924 ± 0.006
Repetitive	51	2201336	2098297	2070750	26547	8393	18154	2.16	98.73 ± 0.01	1.28 ± 0.01	1.162 ± 0.007
Alu	51	446212	419379	412882	6497	4577	1920	2.38	98.45 ± 0.02	1.57 ± 0.02	1.425 ± 0.018
Alu-CG	51	446212	399048	395013	4035	1567	2468	1.57	98.99 ± 0.02	1.02 ± 0.02	0.926 ± 0.015
[]	51	837035	767774	758213	9561	6322	3239	1.95	98.75 ± 0.01	1.26 ± 0.01	1.143 ± 0.012
Human–Baboon											
Overall	42	4984965	4456507	4204745	251762	167380	84382	1.98	94.35 ± 0.01	$5.90 \pm 0.01^{\rm b}$	1.181 ± 0.002
Overall-CG	42	4984965	4351198	4140103	211095	75237	135858	1.81	95.15 ± 0.01	5.03 ± 0.01	1.007 ± 0.002
Exon	24	48578	48098	46627	1471	1042	429	2.43	96.94 ± 0.08	3.13 ± 0.08	0.627 ± 0.016
Unique noncoding	42	3148255	3022715	2862848	159867	106002	53865	1.97	94.71 ± 0.01	5.51 ± 0.01	1.102 ± 0.003
Repetitive	42	1973097	1555295	1456755	98540	66258	32282	2.05	93.66 ± 0.02	6.66 ± 0.02	1.332 ± 0.004
Alu	42	404917	292641	267869	24772	17219	7553	2.28	91.54 ± 0.05	9.07 ± 0.06	1.814 ± 0.012
Alu-CG	42	404917	275514	256962	18552	6506	12046	1.85	93.27 ± 0.05	7.10 ± 0.05	1.419 ± 0.011
[]	42	778575	538863	507237	31626	20465	11161	1.83	94.13 ± 0.03	6.14 ± 0.04	1.228 ± 0.007
Human-Lemur											
Overall	6	623139	423139	341061	82078	47053	35025	1.34	80.60 ± 0.06	22.73 ± 0.08^{b}	2.066 ± 0.008
Overall-CG	6	623139	406011	332321	73690	31863	41827	1.31	81.85 ± 0.06	21.01 ± 0.08	1.910 ± 0.007
Unique	6	370337	313512	255977	57535	33526	24009	1.40	81.65 ± 0.07	21.31 ± 0.09	1.938 ± 0.009
Repetitive	6	244728	103787	80333	23454	12909	10545	1.22	77.40 ± 0.13	27.25 ± 0.19	2.477 ± 0.017
Orthologous sequence: distance (window size 2 and 47,015 \pm 34,144 database). Repetitive s human-chimpanzee co thereiore not partitione CpG dinucleotides in e ^a Substitution rate calcu blf suboptimal alignmei	kb, slid bp for equence mparisc d. Due t ach spec ach spec rations é	(lobally aligned le 100 bp). These human-chimpau es were detecte sons, Alu and L1 to the enrichmet cies (Overall-Cpi assume diverger included in the	with ALIGN (Me with ALIGN (Me nzee, human-b ausing Repeat d using Repeat to of CpG dinuc G).	sthods). A suk anot included ii aboon, and h tMasker (vers t separately. I leotides with human linea verall genetic	optimal alignm, n the analysis. Th numan-lenur cc ion 3.0). Uniqu Relatively few L1 Alu repeats, we o ge from chimpa ge from chimpa	ent was defined ent was defined mean and sta mparisons. Exc e noncoding ru e noncoding trepea and Alu repea considered subs considered subs rnzee, baboon a ses to 1.14 ± 0	as any alignment as any alignment ndard deviation of sequence was 1 egions excluded t its were orthologo titutions without 0 nd lemur of 5.5, $(0.0\%, 6.05 \pm 0.0)$	which exalignme estricted ooth exo betwe pG dinu 1% and 5	cceeded two stan intlengths are 106 only to well-ann nic and repetitivu ic and repetitivu cleotides (Alu-CC) 5 Mya (Goodman 25.69 ± 0.07%, I	dard deviations of 1 dard deviations of 1 5, 107 \pm 41,659, 9 otated human ger regions. For hur mur genomic aligi nas well as the over 1999). respectively (Metho	the mean genetic $5,171 \pm 38,751$, less (NCBI RefSeq nan-baboon and mments and were all content minus ods).

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sequences and their problematic

annotation, more detailed analysis

of this apparent LTR increase is war-

ranted. Genomic comparison with

lemur sequence demonstrates the

most dramatic difference in new

retroelement insertions. Compared to orthologous human genome se-

quence, significant decreases in the

amount of retroelement sequence are observed overall (P < 0.0001,

 χ^2 = 183.17) for most classes of ret-

roposons (Table 2). The most pro-

nounced effect once again is found

among Alu elements. In our analy-

sis of 623 kb of aligned orthologous

sequence, we identified a total of 96 Alu elements in lemur sequences,

compared to 519 Alu elements in

human sequences overall. The ma-

jority of these events appeared to be specific to each lineage (Table 2).

Similar decreases were obtained

based on baboon-lemur genomic

comparisons, indicating that a ma-

jor burst in retrotransposition activ-

ity occurred 25-50 million years



Figure 1 Single-nucleotide variation. A scatter plot of genetic distances (changes/bp) determined from nonoverlapping 3-kb sliding windows for human–chimpanzee (51 loci, 5.0 Mb, 9684 windows), human–baboon (42 loci, 5.0 Mb, 8893 windows), and human–lemur (9 loci, 0.62 Mb, 841 windows) sequence alignments. These were plotted against human divergence times of 5.5, 25, and 55 Mya for chimpanzee, baboon, and lemur alignments, respectively. Suboptimal alignments were excluded. The means and their standard deviations are shown.

site duplications and a polyadenylation tail at the site of integration (Methods). The determination of new insertion elements was based exclusively on the analysis of pairwise sequence alignments. Because the vast majority of retroelement events are irreversible genetic character states (Perna et al. 1992; Deininger and Batzer 1999), unlike other insertion/ deletion events, the directionality of the event could be unambiguously assigned to a specific lineage (Table 2).

Analysis of the chimpanzee data shows a general decline in the level of L1, SINE, and LTR activity compared to the human. A significant decrease (P < 0.05, $\chi^2 = 5.90$) was observed in the number of new elements in chimpanzee (n = 16)compared to human (n = 34). To test whether this difference in the frequency of new retrotransposition events could be observed in an independent data set, we assessed the occurrence of "young" Alu elements in a random sample of 148,102 chimpanzee and 743,245 human BAC-end sequence clones (Zhao et al. 2000; Fujiyama et al. 2002). Lineage-specific Alu elements were identified based on new Alu insertions within human-chimpanzee orthologous genomic sequences (5.0 Mb) that had been identified in the present study. A similar analysis was performed with consensus sequences from Alu subfamilies (Ya5, Yb8, etc). After normalizing for sequence content, we observed a significant decrease (P < 0.001, $\chi^2 = 25.01$) in the number of Alu elements within chimpanzee BAC end sequences compared to human (Table 3).

In contrast to the chimpanzee, the baboon showed a highly significant increase (P = 0.0003; $\chi^2 = 13.05$) in the number of retroelement insertions compared to human orthologous genomic sequence. This overall increase was almost exclusively due to the 1.6-fold increase in the number of Alu insertions observed in the baboon lineage (96 Alu insertions in human compared to 153 insertions in baboon; Table 2). Interestingly, humans showed a significant increase in the number of retroviral LTR insertions (P = 0.0126, $\chi^2 = 6.23$) compared to baboon. Due to the hypermutability of retroviral

ago, consistent with a previous analysis based on Alu subfamily diversity (Shen et al. 1991).

These data predict extreme variability in rates of fixation and/or retrotransposition in different primate lineages. Within the human lineage, the rates of Alu and L1 insertions have remained relatively constant over the last 25 million years. Assuming that our 5.0-Mb subsample is representative of the human genome, we estimate the fixation of 990 and 960 new insertions of L1 elements per genome per million years (chimpanzee/human and baboon/human comparisons, respectively). Similarly for Alu elements, we calculate a remarkably constant rate of new insertion; between 2450 and 2580 new insertions per million years (based on chimpanzee/ human and baboon/human alignments, respectively). Changes in new insertion frequencies, therefore, appear to have occurred within nonhuman primate lineages as opposed to human (Table 2), although additional sequence data from New World and other prosimian lineages will be required before any general trends can be firmly established. Several factors have been proposed to account for lineage-specific changes in retrotransposition activity, including changes in insertion site availability, competence of active progenitor elements, and efficiency of reverse transcription (Deininger and Batzer 1999). The fact that the frequency of both L1 and Alu new insertions is decreased within the chimpanzee genome may point to a reduction in reverse transcriptase activity, since both elements are dependent on the same enzymatic machinery for propagation. In this regard, it is interesting that the average length of new L1 insertions appears to be much smaller in chimpanzee (880 bp) than in human (3500 bp)-a possible indicator of lowered processivity and also a source of a reduced amount of enzyme.

Human Genome Expansion

During our analysis of orthologous genomic sequence, we noticed that the human genome sequence was consistently

Human-Chimpa	nzee		Chimpanzee					Human		
		Insertion	IS	Rate (/N	(VM)/dh		Insertion	SI	Rate (/N	/h/My)
Repeats	Events	Length (bp)	Mean length (bp)	Count	Base	Events	Length (bp)	Mean length (bp)	Count	Base
LINE/L1 SINE/Alu* Other Subtotal*	5 11 5 16	4404 3311 0 7715	881 301 482	0.18 0.41 0.59	162 122 0 69	9 23 34	32393 7036 2040 41469	3599 306 1020 1220	0.33 0.85 0.07 1.25	1195 259 75 369
Human-Baboon			Baboon					Human		
		Insertion	SI	Rate (/N	(My)		Insertion	SI	Rate (/N	(yM)
Repeats	Events	Length (bp)	Mean length (bp)	Count	Base	Events	Length (bp)	Mean length (bp)	Count	Base
LINE/L1 LTR*	26 2	48882 1407	1880 704	0.23 0.02	435 13	36 11	58670 31297	1630 2845	0.32 0.08	523 279
SINE/Alu* Other Subtotal*	153 1 182	45538 130 95957	298 130 527	1.36 0.01 1.62	406 1 855	96 2 145	29000 2836 121803	302 1418 840	0.86 0.02 1.29	258 25 1085
Human-Lemur			Lemur					Human		
		Insertion	SI	Rate (/N	(VM)		Insertion	SI	Rate (/N	/h/My)
	Events	Length (bp)	Mean length (bp)	Count	Base	Events	Length (bp)	Mean length (bp)	Count	Base
DNA LINE/L1*** LTR***	00 m m	4903 3223 2131	613 1074 426	0.19 0.07 0.12	119 78 52	5 53 16	1223 40635 8416	245 767 526	0.10 1.04 0.31	24 799 165
SINE/Alu*** Other Subtotal***	40 22 50	10659 281 16294	266 141 326	0.97 0.05 1.21	259 7 395	234 2 305	64991 750 114792	278 375 376	4.60 0.04 6.00	1278 15 2257
We examined all predominant rep including the and a polyadenylation alignment length $* P < 0.05; ***P <$	insertion d eat. We con cient repeat s swere 5.0, 0.001 by X	leletion events in ex subclasses that pass, subclasses that pass, e site of integration. 5.0, and 0.62 Mb ff ² test.	cess of 100 bp from glob nterspersed repeat phylog ed our test. Further, in the The rate calculation assum or human-chimpanzee, hu	al alignments. eny based on t e case of L1 an ied divergence iman-baboon,	An indel wa che establishe d Alu repeats times of the h and human-	s classified as d repeat subcli , insertion seq numan lineage lemur sequenc	a retrotransposition asses as reported pre uences were examin from chimpanzee, t ce alignment, respec	event if at least 80% of eviously (Smit 1999). All in hed for the presence of tan aboon, and lemur of 5.5, titvely.	the indel cont isertions were o get-site duplic 25, and 55 My	ained one considered ations and a. Pairwise

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Table 3.	Frequency of "Young"	Alu Elements	Within BAC
End Seque	ences		

	D	Database			
Lineage-specific Query Sequences	Human BES	Chimpanzee BES			
Human Alus Chimpanzee Alus	15.92* 19.49**	6.00 6.00			

Lineage-specific Alu retrotransposition events were identified by analysis of human-chimpanzee orthologous genomic sequence. Extracted representative events (query sequences) were searched against a database of BAC end sequences (BES) which included 743,245 human BES (354,136,231 bp; Zhao et al. 2000) and 148,102 chimpanzee BES (115,468,024 bp; Fujiyama et al. 2002). Only full-length Alu elements were considered. When guery sequence and BAC end-sequences were from the same species, a sequence similarity cutoff of 98.5% was used to account for sequencing errors within the single-pass BES database. When query sequence and BAC end sequences were from different species, sequence similarities greater or equal to 96.5% were counted (to account for sequencing error and species divergence). Human counts were further normalized by the size ratio of human and chimpanzee BAC end sequence library. *P < 0.05; **P < 0.01 by χ^2 test assuming equal distribution. In both cases, the human BES database shows a significant increase in the number of young Alu elements.

longer (Table 4) for each primate species comparison overall. The average human sequence expansion ranged from 0.6% for human–baboon comparisons to as much as 19% for human–lemur comparisons (Figs. 2,3). The expansion in human sequence compared to baboon is particularly striking, considering that an additional 16.5 kb of sequence has been introduced by the apparent increase in baboon Alu retroposon activity (Table 2). A permutation test of the difference was performed at the level of the alignment as well as at the level of individual insertion/deletion events (see Methods; Table 4)

for each species comparison. With the exception of the baboon, a significant increase (P < 0.05) in genome size was observed for the human genome in each case (Table 4). We divided the genome into two fractions, repetitive and unique DNA, to assess the source of this expansion. Most of the significant increase (80%–100%) could be assigned to an increase in retroposon content within the orthologous human sequence (Table 4).

Large-scale comparative sequencing of vertebrate genomes has shown that syntenic regions in other species are shorter and contain fewer repeats compared to human (Dehal et al. 2001; Aparicio et al. 2002; Mural et al. 2002). Our analysis extends this property of the human genome to at least two other primate species. One possible explanation for these differences is a change in the deletion rate of repetitive elements within different lineages (Dehal et al. 2001; Aparicio et al. 2002; Mural et al. 2002). The human lineage, for example, may retain more retroposon elements because its inherent mutational mechanisms are less efficient at deleting such events.

To determine whether an increase in the deletion rate in other primate lineages might account for this difference, we performed two tests. First, we analyzed all large insertion/ deletion events (>100 bp in length) for both baboon and chimpanzee comparisons. Three classes were distinguished: indels that were characteristic of a retrotransposition event (see above), those that were associated with a repetitive sequence at their junctions and were likely the result of a deletion event (Gilbert et al. 2002), and those that were not associated with repeats (nonrepeat-associated insertion/deletions, termed NRAIDs; Suppl. Table S5, Suppl. Fig. S7). No significant difference (P = 0.2–0.5, χ^2 = 0.2–1.71; Suppl. Table S5) was observed in the number of indels in the latter two categories. In contrast, estimates in the number of new insertion events that arose as a result of retrotransposition were significantly different for each species comparison.

As a second test, we compared the mean genetic distance

	All		Repea	ts	Unique	9
	Length (bp)	%	Length (bp)	%	Length (bp)	%
Human	5410556	100.00	2204532	40.75	3206024	59.25
Chimpanzee	5351536	98.91	2148580	39.71	3202956	59.20
Difference	59020	1.09**	55952	1.03**	3068	0.06
Human	5560707	100.00	2181276	39.23	3379431	60.77
Baboon	5527115	99.40	2143997	38.56	3383118	60.84
Difference	33592	0.60	37279	0.67ª	- 3687	0.07
Human	924753	100.00	374742	40.52	550011	59.48
Lemur	749135	81.01	216540	23.42	532595	57.59
Difference	175618	18.99**	158202	17.11**	17416	1.88
Baboon	790055	100.00	278145	35.21	511910	64.79
Lemur	675780	85.54	187084	23.68	488696	61.86
Difference	114275	14.46**	91061	11.53**	23214	2.94

For orthologous genomic comparisons, the length of aligned sequence and difference were considered for each species comparison. Repetitive and unique portions were identified using RepeatMasker (version 3.0) from human–chimpanzee (51 loci), human–baboon (42 loci), human–lemur (nine loci), and baboon–lemur (eight loci) comparisons. In the event that lemur common repeats were not efficiently masked, intraspecific sequence similarity searches (BLAST) were performed to identify potentially missing repeats. Relative percentages were calculated assuming the length of larger primate genome (human or baboon) as 100%. Significance of the difference in genome size was tested by a permutation test (10,000 replicates). *P < 0.05; **P < 0.01. a The difference in repeat composition is greater than the total due to an expansion of LTR content and deletion of 3687 bp of unique sequence. Table S8 shows a more detailed breakdown by repeat class for both human–lemur and baboon–lemur alignments.



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Figure 3 Primate genome size variation. Repetitive and unique portions of aligned orthologous sequences were identified by Repeat-Masker (version 3.0, slow option). Relative fractions were based on the larger primate genome. Significance of the difference in genome size was determined by a permutation test (10,000 replicates, see Methods). Asterisks over species bars indicate significant differences in overall lengths, and those between species bars indicate significant differences in either repetitive or unique lengths between two species. *, P < 0.05; **, P < 0.01.

for lineage-specific Alus within both the human and lemur lineages. If a higher deletion rate were responsible for the depletion of repeats within the lemur lineage, we would expect the mean genetic distance for Alu repeats within lemur to be lower as the longevity of Alu insertions would be reducedolder Alu elements would be more likely to be deleted or truncated within a background of increased deletion. A comparison of lineage-specific full-length Alus in lemur (K = 0.258 \pm 0.015, n = 30) and human (K = 0.284 \pm 0.014, n = 239) reveals comparable levels of Alu diversity. Similarly, analysis of older repetitive elements (L2, L3, MIR, and DNA transposons) that are believed to have inserted before the separation of the two lineages shows virtually no difference in either count or the relative proportion in the human and lemur genomes (Suppl. Table S9A). Combined, these data strongly suggest that recent (<50 Mya) changes in the rates of retrotransposition as opposed to deletion are responsible for the expansion of the hominoid genome.

Our comparison of human and baboon genomic sequence to lemur shows the most dramatic expansion (15%–19%) in genome size (Fig. 2). In a previous study based on a sampling of DNA content from 15 stepsirhines and 48 haplorhines species, it was reported that the genomes of prosimian species were significantly smaller (9% decrease; 7.1 ± 0.2 pg vs. 7.8 ± 0.2 pg; Pellicciari et al. 1982). It was unknown, however, whether such differences were attributable to centric chromatin, which is known to be cytogenetically reduced in size among prosimians (Martin 1990). Our analysis of lemur and human data indicates that the difference is in fact euchromatic in nature and that it is almost exclusively repeat-

derived (Fig. 3). All classes of younger retrotransposons (Alus, L1s, and LTR) contribute to this increase, whereas more ancient elements such as L2, L3, and DNA transposons do not contribute to this increase by differential deletion. Interestingly, although the number of Alu elements appears to be significantly increased, among the LTR and L1 elements, the average length of the insertion has increased whereas the number of such events has not. This effect is seen in both human and baboon compared to lemur. Assuming a divergence of human and baboon approximately 25 million years ago, the data would support a major increase in genome size due to an increase in retrotransposition fixation.

Summary

The analysis here provides a large-scale and unbiased assessment of primate genome variation. As such, it is expected that these data will serve as a valuable baseline for future studies of primate molecular evolution both at the level of singlenucleotide variation as well as that of retrotransposition. The human genome is particularly enriched in both number and length of retrotransposons. It has grown as a result of a major burst in Alu activity 25-55 million years ago and has subsequently continued to expand compared to more closely related primates, due to lineage-specific shifts in retroposon activity within the last 25 million years of evolution. Compared to every animal genome sequenced to date, the human genome is larger and harbors more repeats within its euchromatin. Since the rate of substitution is fundamentally lower and our generation time is longer compared to those species, such changes may have contributed to this repeat-enrichment. In this context, the finding that the human genome is significantly larger than that of the chimpanzee is unexpected. The mutation rate and generation time among the ancestors of these large-bodied hominoids are believed to have remained relatively constant (Ruvolo 1997; Webster et al. 2002), although the population history is believed to be radically different among these species. The repeat-associated reduction in the chimpanzee genome, however, is slight and must await further validation before being declared a general property. We cannot, for example, exclude the possibility that other rare and very large repetitive sequences (i.e., segmental duplications) may compensate for this difference. Nevertheless, it is interesting that similar expansions of smaller tandem microsatellites, such as dinucleotide and trinucelotide repeat sequences, have been reported (Cooper et al. 1998; Webster et al. 2002) in human compared to chimpanzee. Although the molecular basis for these differences is not well understood, combined the data support a repeat-driven expansion of our genome. Since such sequences have been shown to be potent mutagens at the structural as well as the genic level, it follows that their contribution to phenotypic change and evolution might be more significant than previously anticipated.

METHODS

Orthologous Sequences

Large genomic sequences (>50 kb in length) from chimpanzee (RP43), baboon (RP41), and lemur (LB2) were retrieved from GenBank. To provide high-quality large-scale genomic alignments, we limited our analysis to genomic sequences that were completely finished or where the sequence contigs were ordered and oriented. Sequence accessions were considered

where there were fewer than three contigs and no internal ambiguous bases. Finished sequence was generated to the standards established for sequencing the human genome (see http://www.genome.wustl.edu/Overview/finrulesname. php?G16=1), which includes closure of all sequence gaps and achieving an estimated error rate of <1 in 10,000 bp (Felsenfeld et al. 1999). Among the working draft sequences, an analysis of the assembly quality revealed an ~10-11-fold redundancy of high-quality bases (Phred Q > = 20). To search for orthologous sequences, we extracted segments longer than 50 kb in length and masked the sequences for common repeat elements (Smit 1999, http://repeatmasker.genome.washington. edu/cgi-bin/RepeatMasker). Because duplicated regions of the genome complicate identification of orthologous segments and confound genetic distance estimates (Chen et al. 2001), we excluded any accession if it was located within a known duplicated region of the human genome (Bailey et al. 2002, http://humanparalogy.gene.cwru.edu/SDD/index.htm). A total of 102 nonhuman primate accessions met these criteria, corresponding to nine lemur, 42 baboon, and 51 chimpanzee genomic clones. Nonhuman primate genomic sequence was generated almost exclusively by the U.S. National Institutes of Health Intramural Sequencing Center (http:// www.nisc.nih.gov/open_page.html?staff.html). Ninety-nine of 102 of the sequences mapped to phylogenetic group chromosome 7 and were part of a targeted comparative sequencing effort to three gene-containing regions on chromosome 7 (Thomas et al. 2002). Five of the lemur genomic loci mapped to a gene-rich region near the CFTR locus on human 7q31. 2. The majority of nonhuman primate clones mapped primarily to two regions within 7p14.3 and 7q22.1 (positions 30,000,000-35,000,000 and 95,000,000-103,000,000 within build30, June 2002 assembly). A complete list of all accessions, their map location with respect to the human genome, and their sequence attributes are provided (http://eichlerlab. cwru.edu/primategenome/ and Suppl. Fig. S2, Suppl. Table S3). Orthologous human sequence was identified by sequence similarity searches (BLAST) of nonhuman primate sequence queried against a formatted version of the assembled human genome (http://genome.ucsc.edu/). Human genomic sequence underlying the assembly was obtained from GenBank accessions. Overlapping sequences within a species were excluded based on human genome assembly coordinates. Although only nine genomic regions are compared between human and lemur, each of these regions represents ~70 kb of orthologous sequence. Our genomic analysis (Fig. 1, Suppl. Fig. S4), therefore, involves the analysis of more than 841 nonoverlapping blocks of 3 kb of genomic sequence. Therefore, it is reasonable to believe that these datasets are sufficiently representative and robust enough to draw sound conclusions regarding rates and properties of primate genomic mutation. As a control for selection bias and rate variation among these genomic regions, we analyzed 1000 BAC-end sequences randomly selected from chimpanzee, baboon, and lemur BAC libraries. A comparison of these alignments to these large-scale genomic alignments showed comparable results (Results and Discussion).

Genomic Sequence Alignment

Orthologous sequence relationships between human and nonhuman primate genomic sequences were initially delineated using Miropeats (Parsons 1995, http://genome. wustl.edu/gsc/index.shtml), and the sequences were subsequently extracted using two_way_mirror (J. Bailey, unpubl.). We used the Myers-Miller algorithm (Myers and Miller 1988) to construct all optimal global alignments. One of the most significant challenges of large-scale genomic analyses is the generation of biologically meaningful global alignments (Chen et al. 2001). As sequence becomes increasingly divergent, the reliable treatment of insertion/deletions becomes

particularly problematic. Ineffective treatment of insertion/ deletions (indels) may lead to the formation of suboptimal global alignments providing erroneously higher estimates of sequence divergence. To establish the optimal parameters for global alignment, we initially analyzed a subset of large-scale sequence alignments between human, baboon, and lemur. Using the software ALIGN (Myers and Miller 1988), we tested a series of gap opening and extension penalties and their impact on the frequency of single-nucleotide and insertion/ deletion events (Suppl. Fig. S1). For each species we selected parameters that minimized sequence divergence and the number of indels. For equally parsimonious gap parameters, we selected parameters (-f 50 - g 1) where known "young" retrotransposition events were treated as a single insertion/ deletion event. All alignments were manually inspected for extreme fluctuations in genetic distance using align_slider_viewer (J. Bailey, unpubl.). A suboptimal alignment was defined as any alignment which exceeded two standard deviations of the mean genetic distance (window size 2 kb, slide 100 bp). These regions were considered separately in the analysis (Table 1). A total of six (16 kb), 23 (43 kb), and 17 (26 kb) such subalignments were classified as suboptimal for chimpanzee, baboon, and lemur comparisons to human, respectively. Altering gap parameters recovered approximately 50% of these suboptimal alignments for human-chimpanzee alignments but not for the other primate comparisons. Only a small fraction (<5%) of all aligned bases was classified as suboptimal. A total of 5.0, 5.0, and 0.62 Mb of genomic sequence was successfully aligned between human and chimpanzee, baboon, and lemur, respectively. We further constructed 19 multiple alignments for human, chimpanzee, and baboon (alignment length 2.5 Mb) and five multiple alignments for all four species (alignment length 0.51 Mb) using ClustalW. Tajima's relative rate tests were performed on these multiple alignments using MEGA. All alignments, including graphical assessments, are available online (http://eichlerlab. cwru.edu/primategenome/).

Genetic Distance Estimates

For all estimates of genetic distance (K; Table 1), we used Kimura's two-parameter method, which corrects for multiple events and transversion/transition mutational biases (Table 1; Kimura 1980). Insertion/deletion events were not factored into these calculations (Britten 2002). Repetitive, unique noncoding and exonic portions from the sequence alignments were extracted using MaM (Multiple Alignment Manipulator; Alkan et al. 2002, http://genomics.cwru.edu/MAM.html). Repeat coordinates were identified using the slow option of RepeatMasker v3.0. Five major classes of repeats were considered in this analysis (LINES, SINES, DNA Transposons, LTR, and simple repeats). To eliminate the possibility that more divergent or novel common repeats (particularly for the lemur) may not have been effectively masked by RepeatMasker, intraspecific sequence-similarity searches were performed. Exon definition was limited to well annoted human genes (NCBI RefSeq: http://www.ncbi.nlm.nih.gov/LocusLink/refseq. html). Among these, a total of 460 coding exons corresponding to 52 genes were analyzed. Sliding window analyses (Fig. 1) were performed using align_slider (J. Bailey, unpubl.). Rates of substitution were calculated using K/2T, where human divergence times of 5.5, 25, and 55 million years ago were used for chimpanzee, baboon, and lemur alignments, respectively (Kumar and Hedges 1998; Goodman 1999). All alignment attributes were maintained within a mySQL database, which facilitated cross-referencing with various properties of the genomic sequence. DNA sequences corresponding to recent retroelements (Alu and L1) were extracted from the aligned sequences. Multiple sequence alignments were generated (ClustalW) and within-group and between-group estimates of genetic distance were calculated (MEGA2; Kumar et al. 2001).

Insertion/Deletion Analysis

Insertion/deletion (indel) events within the pairwise alignments were initially separated by length into two groups (<100 bp and \geq 100 bp). This classification was based on the rationale that most retrotransposition events are greater than 100 bp in length, whereas the vast majority of the smaller events result from other mutational events (microsatellite variation, replication slippage, small local deletion events). More than 80% of all indels are equal to or less than 15 bp in length but contribute to less than 3.6% of the overall length differences within an alignment. This is in agreement with a recently published analysis (Britten 2002). A complete count of the total number of indels and their length distribution are available (Suppl. Table S5, Suppl. Fig. S6).

Large gaps (>100 bp) within a genomic pairwise alignment may occur as a result of a deletion in one species or an insertion in the other. Such events cannot, usually, be assigned. It is expected that, many such large events will be associated with a common repeat sequence due to homologybased deletion of repeat sequences (Gilbert et al. 2002) and retrotransposition-based insertion events. We therefore further subdivided indels (>100 bp) into one of two categories based on their association with a repeat sequence. We classified an indel as a retrotransposition if at least 80% of the indel contained one predominant repeat (LINE, SINE, LTR). We considered the known interspersed repeat phylogeny based on the established repeat subclasses as reported previously (Smit 1999). All insertions were considered including the ancient repeat subclasses that passed our test. Further, in the case of L1 and Alu repeats, insertion sequences were examined for the presence of target-site duplications and a polyadenylation tail at the site of integration. The vast majority of retroelements events are irreversible genetic character states (Perna et al. 1992; Deininger and Batzer 1999), and it is therefore highly unlikely that a deletion event would occur to precisely remove a retroelement during evolution. Unlike other insertion deletion events, then, the directionality of the event could be unambiguously assigned to a specific lineage (Table 2). Large indels in which one end or both ends were placed within a repetitive sequence were categorized separately (Suppl. Table S8).

Two basic statistical tests were performed during the analysis of indels. Differences in counts were assessed using the χ^2 test based on the assumption that alignment parameters would not show a species preference for insertions. Differences in genomic length (insertion/deletions) were examined using a permutation test of the difference for both orthologous loci and for individual indels (>100 bp). Briefly, for each alignment, the greater length was randomly assigned between the two species of interest. P-values were defined as the fraction of replicates out of 10,000 which surpassed or equaled the observed length differences. These permutations were also done on an indel by indel basis by effectively assigning any given insertion or deletion to a species randomly. The sum was then compared to the observed length differences to determine the P-value. Permutation tests, therefore, were performed at the level of the total alignment as well as at the level of the individual insertion/deletion events.

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REFERENCES

- Alkan, C., Tuzun, E., Eicher, E.E., Bailey, J.A., and Sahinalp, S.C. 2002. MaM: Multiple alignment manipulator. *Currents in computational molecular biology 2002*. pp. 3–4. Celera Genomics, Rockville, MD.
- Aparicio, S., Chapman, J., Stupka, E., Putnam, N., Chia, J.M., Dehal, P., Christoffels, A., Rash, S., Hoon, S., Smit, A., et al. 2002. Whole-genome shotgun assembly and analysis of the genome of *Fugu rubripes. Science* **297**: 1301–1310.
- Bailey, J.A., Gu, Z., Clark, R.A., Reinert, K., Samonte, R.V., Schwartz, S., Adams, M.D., Myers, E.W., Li, P.W., and Eichler, E.E. 2002. Recent segmental duplications in the human genome. *Science* 297: 1003–1007.
- Bailey, W.J., Fitch, D.H., Tagle, D.A., Czelusniak, J., Slightom, J.L., and Goodman, M. 1991. Molecular evolution of the ψ η -globin gene locus: Gibbon phylogeny and the hominoid slowdown. *Mol. Biol. Evol.* **8:** 155–184.
- Batzer, M.A. and Deininger, P.L. 2002. Alu repeats and human genomic diversity. *Nat. Rev. Genet.* **3:** 370–379.
- Bohossian, H.B., Skaletsky, H., and Page, D.C. 2000. Unexpectedly similar rates of nucleotide substitution found in male and female hominids. *Nature* **406**: 622–625.
- Britten, R.J. 2002. Divergence between samples of chimpanzee and human DNA sequences is 5%, counting indels. *Proc. Natl. Acad. Sci.* **99:** 13633–13635.
- Chen, F.C. and Li, W.H. 2001. Genomic divergences between humans and other hominoids and the effective population size of the common ancestor of humans and chimpanzees. *Am. J. Hum. Genet.* 68: 444–456.
- Chen, F.C., Vallender, E.J., Wang, H., Tzeng, C.S., and Li, W.H. 2001. Genomic divergence between human and chimpanzee estimated from large-scale alignments of genomic sequences. *J. Hered.* **92**: 481–489.
- Chiaromonte, F., Yang, S., Elnitski, L., Yap, V.B., Miller, W., and Hardison, R.C. 2001. Association between divergence and interspersed repeats in mammalian noncoding genomic DNA. *Proc. Natl. Acad. Sci.* 98: 14503–14508.

Cooper, G., Rubinsztein, D.C., and Amos, W. 1998. Ascertainment bias cannot entirely account for human microsatellites being longer than their chimpanzee homologues. *Hum. Mol. Genet.* 7: 1425–1429.

- Dehal, P., Predki, P., Olsen, A.S., Kobayashi, A., Folta, P., Lucas, S., Land, M., Terry, A., Ecale Zhou, C.L., Rash, S., et al. 2001. Human chromosome 19 and related regions in mouse: Conservative and lineage-specific evolution. *Science* 293: 104–111.
- Deininger, P.L. and Batzer, M.A. 1999. Alu repeats and human disease. *Mol. Genet. Metab.* **67**: 183–193.
- Deininger, P.L. and Batzer, M.A. 2002. Mammalian retroelements. Genome Res. 12: 1445–1465.
- Ebersberger, I., Metzler, D., Schwarz, C., and Paabo, S. 2002. Genomewide comparison of DNA sequences between humans and chimpanzees. *Am. J. Hum. Genet.* **70**: 1490–1497.
- Eichler, E.E. and DeJong, P.J. 2002. Biomedical applications and studies of molecular evolution: A proposal for a primate genomic library resource. *Genome Res.* **12**: 673–678.
- Felsenfelá, A., Peterson, J., Schloss, J., and Guyer, M. 1999. Assessing the quality of the DNA sequence from the Human Genome Project. *Genome Res.* 9: 1–4.
- Fujiyama, A., Watanabe, H., Toyoda, A., Taylor, T.D., Itoh, T., Tsai, S.F., Park, H.S., Yaspo, M.L., Lehrach, H., Chen, Z., et al. 2002. Construction and analysis of a human-chimpanzee comparative clone map. *Science* **295**: 131–134.
- Gilbert, N., Lutz-Prigge, S., and Moran, J.V. 2002. Genomic deletions created upon LINE-1 retrotransposition. *Cell* **110**: 315–325.
- Goodman, M. 1999. The genomic record of humankind's
- evolutionary roots. Am. J. Hum. Genet. **64:** 31–39. Goodman, M., Barnabas, J., Matsuda, G., and Moore, G.W. 1971.

Molecular evolution in the descent of man. Nature 233: 604-613.

- Horai, S., Hayasaka, K., Kondo, R., Tsugane, K., and Takahata, N. 1995. Recent African origin of modern humans revealed by complete sequences of hominoid mitochondrial DNAs. Proc. Natl. Acad. Sci. 92: 532–536.
- Hurles, M.E. 2001. Gene conversion homogenizes the CMT1A paralogous repeats. BMC Genomics 2: 11-19.
- International Human Genome Sequencing Consortium. 2001. Initial sequencing and analysis of the human genome. Nature 409: 860-920
- Kaessmann, H., Wiebe, V., and Paabo, S. 1999. Extensive nuclear DNA sequence diversity among chimpanzees. Science **286:** 1159–1162.
- Kimura, M. 1980. A simple method for estimating evolutionary rates of base substitutions through comparative studies of nucleotide sequences. J. Mol. Evol. 16: 111-120.
- Koop, B.F., Miyamoto, M.M., Embury, J.E., Goodman, M., Czelusniak, J., and Slightom, J.L. 1986. Nucleotide sequence and evolution of the orangutan ϵ globin gene region and surrounding Alu repeats. J. Mol. Evol. 24: 94-102.
- Kumar, S. and Hedges, S.B. 1998. A molecular timescale for vertebrate evolution. Nature 392: 917-920.
- Kumar, S., Tamura, K., Jakobsen, I.B., and Nei, M. 2001. MEGA2: Molecular evolutionary genetics analysis software. Bioinformatics 17: 1244-1245
- Li, W. 1997. Molecular evolution. pp. 177-213. Sinauer Associates, Sunderland, MA.
- Li, W.H. and Tanimura, M. 1987. The molecular clock runs more slowly in man than in apes and monkeys. Nature 326: 93-96.
- Martin, R.D. 1990. Primate origins and evolution: A phylogenetic
- reconstruction. pp. 558–559. University Press, Princeton, NJ. Moran, J.V., DeBerardinis, R.J., and Kazazian, Jr., H.H. 1999. Exon shuffling by L1 retrotransposition. Science 283: 1530-1534.
- Mural, R.J., Adams, M.D., Myers, E.W., Smith, H.O., Miklos, G.L., Wides, R., Halpern, A., Li, P.W., Sutton, G.G., Nadeau, J., et al. 2002. A comparison of whole-genome shotgun-derived mouse chromosome 16 and the human genome. Science **296:** 1661-1671
- Myers, E.W. and Miller, W. 1988. Optimal alignments in linear space. Comput. Appl. Biosci. 4: 11-17.
- Parsons, J. 1995. Miropeats: Graphical DNA sequence comparisons. Comput. Appl. Biosci. 11: 615-619.

Pellicciari, C., Formenti, D., C.A. Redi, and M.R. M.G. 1982. DNA content variability in primates. J. Hum. Evol. 11: 131-141.

- Perna, N.T., Batzer, M.A., Deininger, P.L., and Stoneking, M. 1992. Alu insertion polymorphism: A new type of marker for human population studies. Hum. Biol. 64: 641-648.
- Powell, J.R. and Caccone, A. 1990. The TEACL method of DNA-DNA hybridization: Technical considerations. J. Mol. Evol. **30:** 267–272.
- Ruvolo, M. 1997. Molecular phylogeny of the hominoids: Inferences from multiple independent DNA sequence data sets. Mol. Biol. Evol. 14: 248-265.
- Shen, M.-R., Batzer, M., and Deininger, P. 1991. Evolution of the master Alu gene(s). J. Mol. Evol. 33: 311-320.
- Sibley, C.G. and Ahlquist, J.E. 1984. The phylogeny of the hominoid

primates, as indicated by DNA-DNA hybridization. J. Mol. Evol. **20:** 2–15.

- Sibley, C.G., Comstock, J.A., and Ahlquist, J.E. 1990. DNA hybridization evidence of hominoid phylogeny: A reanalysis of the data. J. Mol. Evol. 30: 202-236.
- Smit, A.F. 1999. Interspersed repeats and other mementos of transposable elements in mammalian genomes. Curr. Opin. Genet. Dev. 9: 657–663.
- Smith, J. 1992. Evolutionary biology. Byte-sized evolution. Nature 355: 772-773.
- Smith, N.G., Webster, M.T., and Ellegren, H. 2002. Deterministic mutation rate variation in the human genome. Genome Res. 12: 1350-1356.

Thomas, J.W., Prasad, A.B., Summers, T.J., Lee-Lin, S.Q., Maduro, V.V., Idol, J.R., Ryan, J.F., Thomas, P.J., McDowell, J.C., and Green, E.D. 2002. Parallel construction of orthologous sequence-ready clone contig maps in multiple species. Genome Res. 12: 1277-1285.

Venter, J.C., Adams, M.D., Myers, E.W., Li, P.W., Mural, R.J., Sutton, G.G., Smith, H.O., Yandell, M., Evans, C.A., Holt, R.A., et al. 2001. The sequence of the human genome. *Science* **291:** 1304–1351.

- Webster, M.T., Smith, N.G., and Ellegren, H. 2002. Microsatellite evolution inferred from human-chimpanzee genomic sequence alignments. Proc. Natl. Acad. Sci. 99: 8748–8753.
- Zhao, S., Malek, J., Mahairas, G., Fu, L., Nierman, W., Venter, J.C., and Adams, M.D. 2000. Human BAC ends quality assessment and sequence analyses. Genomics 63: 321–332.
- Zuckerkandl, E. and Pauling, L. 1965. Molecules as documents of evolutionary history. J. Theor. Biol. 8: 357-366.

WEB SITE REFERENCES

http://www.nisc.nih.gov/; NIH intramural sequencing center. http://eichlerlab.cwru.edu/primategenome/; supplemental material:

- pairwise and multiple alignments. http://www.genome.wustl.edu/Overview/finrulesname.php?G16=1;
- finishing standards for the human genome project. http://repeatmasker.genome.washington.edu/cgi-bin/RepeatMasker;
- Repeatmasker Web server. http://humanparalogy.gene.cwru.edu/SDD/index.htm; segmental
- duplication database at CWRU.
- http://www.nisc.nih.gov/open_page.html?staff.html; NIH intramural sequencing center staff page.
- http://genome.ucsc.edu/; UCSC genome bioinformatics site.
- http://genome.wustl.edu/gsc/index.shtml; Genome sequencing center at Washington University.
- http://genomics.cwru.edu/MAM.html; MaM download page at Center for Computational Genomics, CWRU
- http://www.ncbi.nlm.nih.gov/LocusLink/refseq.html; NCBI reference sequences.

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