

# Initial sequence of the chimpanzee genome and comparison with the human genome

The Chimpanzee Sequencing and Analysis Consortium\*

**Here we present a draft genome sequence of the common chimpanzee (*Pan troglodytes*). Through comparison with the human genome, we have generated a largely complete catalogue of the genetic differences that have accumulated since the human and chimpanzee species diverged from our common ancestor, constituting approximately thirty-five million single-nucleotide changes, five million insertion/deletion events, and various chromosomal rearrangements. We use this catalogue to explore the magnitude and regional variation of mutational forces shaping these two genomes, and the strength of positive and negative selection acting on their genes. In particular, we find that the patterns of evolution in human and chimpanzee protein-coding genes are highly correlated and dominated by the fixation of neutral and slightly deleterious alleles. We also use the chimpanzee genome as an outgroup to investigate human population genetics and identify signatures of selective sweeps in recent human evolution.**

More than a century ago Darwin<sup>1</sup> and Huxley<sup>2</sup> posited that humans share recent common ancestors with the African great apes. Modern molecular studies have spectacularly confirmed this prediction and have refined the relationships, showing that the common chimpanzee (*Pan troglodytes*) and bonobo (*Pan paniscus* or pygmy chimpanzee) are our closest living evolutionary relatives<sup>3</sup>. Chimpanzees are thus especially suited to teach us about ourselves, both in terms of their similarities and differences with human. For example, Goodall's pioneering studies on the common chimpanzee revealed startling behavioural similarities such as tool use and group aggression<sup>4,5</sup>. By contrast, other features are obviously specific to humans, including habitual bipedality, a greatly enlarged brain and complex language<sup>5</sup>. Important similarities and differences have also been noted for the incidence and severity of several major human diseases<sup>6</sup>.

Genome comparisons of human and chimpanzee can help to reveal the molecular basis for these traits as well as the evolutionary forces that have moulded our species, including underlying mutational processes and selective constraints. Early studies sought to draw inferences from sets of a few dozen genes<sup>7–9</sup>, whereas recent studies have examined larger data sets such as protein-coding exons<sup>10</sup>, random genomic sequences<sup>11,12</sup> and an entire chimpanzee chromosome<sup>13</sup>.

Here we report a draft sequence of the genome of the common chimpanzee, and undertake comparative analyses with the human genome. This comparison differs fundamentally from recent comparative genomic studies of mouse, rat, chicken and fish<sup>14–17</sup>. Because these species have diverged substantially from the human lineage, the focus in such studies is on accurate alignment of the genomes and recognition of regions of unusually high evolutionary conservation to pinpoint functional elements. Because the chimpanzee lies at such a short evolutionary distance with respect to human, nearly all of the bases are identical by descent and sequences can be readily aligned except in recently derived, large repetitive regions. The focus thus turns to differences rather than similarities. An observed difference at a site nearly always represents a single event, not multiple indepen-

dent changes over time. Most of the differences reflect random genetic drift, and thus they hold extensive information about mutational processes and negative selection that can be readily mined with current analytical techniques. Hidden among the differences is a minority of functionally important changes that underlie the phenotypic differences between the two species. Our ability to distinguish such sites is currently quite limited, but the catalogue of human–chimpanzee differences opens this issue to systematic investigation for the first time. We would also hope that, in elaborating the few differences that separate the two species, we will increase pressure to save chimpanzees and other great apes in the wild.

Our results confirm many earlier observations, but notably challenge some previous claims based on more limited data. The genome-wide data also allow some questions to be addressed for the first time. (Here and throughout, we refer to chimpanzee–human comparison as representing hominids and mouse–rat comparison as representing murids—of course, each pair covers only a subset of the clade.) The main findings include:

- Single-nucleotide substitutions occur at a mean rate of 1.23% between copies of the human and chimpanzee genome, with 1.06% or less corresponding to fixed divergence between the species.
- Regional variation in nucleotide substitution rates is conserved between the hominid and murid genomes, but rates in subtelomeric regions are disproportionately elevated in the hominids.
- Substitutions at CpG dinucleotides, which constitute one-quarter of all observed substitutions, occur at more similar rates in male and female germ lines than non-CpG substitutions.
- Insertion and deletion (indel) events are fewer in number than single-nucleotide substitutions, but result in ~1.5% of the euchromatic sequence in each species being lineage-specific.
- There are notable differences in the rate of transposable element insertions: short interspersed elements (SINEs) have been threefold more active in humans, whereas chimpanzees have acquired two new families of retroviral elements.

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- Orthologous proteins in human and chimpanzee are extremely similar, with ~29% being identical and the typical orthologue differing by only two amino acids, one per lineage.
- The normalized rates of amino-acid-altering substitutions in the hominid lineages are elevated relative to the murid lineages, but close to that seen for common human polymorphisms, implying that positive selection during hominid evolution accounts for a smaller fraction of protein divergence than suggested in some previous reports.
- The substitution rate at silent sites in exons is lower than the rate at nearby intronic sites, consistent with weak purifying selection on silent sites in mammals.
- Analysis of the pattern of human diversity relative to hominid divergence identifies several loci as potential candidates for strong selective sweeps in recent human history.

In this paper, we begin with information about the generation, assembly and evaluation of the draft genome sequence. We then explore overall genome evolution, with the aim of understanding mutational processes at work in the human genome. We next focus on the evolution of protein-coding genes, with the aim of characterizing the nature of selection. Finally, we briefly discuss initial insights into human population genetics.

In recognition of its strong community support, we will refer to chimpanzee chromosomes using the orthologous numbering nomenclature proposed by ref. 18, which renumbers the chromosomes of the great apes from the International System for Human Cytogenetic Nomenclature (ISCN; 1978) standard to directly correspond to their human orthologues, using the terms 2A and 2B for the two ape chromosomes corresponding to human chromosome 2.

### Genome sequencing and assembly

We sequenced the genome of a single male chimpanzee (Clint; Yerkes pedigree number C0471; Supplementary Table S1), a captive-born descendant of chimpanzees from the West Africa subspecies *Pan troglodytes verus*, using a whole-genome shotgun (WGS) approach<sup>19,20</sup>. The data were assembled using both the PCAP and ARACHNE programs<sup>21,22</sup> (see Supplementary Information 'Genome sequencing and assembly' and Supplementary Tables S2–S6). The former was a *de novo* assembly, whereas the latter made limited use of human genome sequence (NCBI build 34)<sup>23,24</sup> to facilitate and confirm contig linking. The ARACHNE assembly has slightly greater continuity (Table 1) and was used for analysis in this paper. The draft genome assembly—generated from ~3.6-fold sequence redundancy of the autosomes and ~1.8-fold redundancy of both sex chromosomes—covers ~94% of the chimpanzee genome with >98% of the sequence in high-quality bases. A total of 50% of the sequence (N50) is contained in contigs of length greater than 15.7 kilobases (kb) and supercontigs of length greater than 8.6 megabases (Mb). The assembly represents a consensus of two haplotypes, with one allele from each heterozygous position arbitrarily represented in the sequence.

**Assessment of quality and coverage.** The chimpanzee genome assembly was subjected to rigorous quality assessment, based on comparison to finished chimpanzee bacterial artificial chromosomes (BACs) and to the human genome (see Supplementary Information

'Genome sequencing and assembly' and Supplementary Tables S7–S16).

Nucleotide-level accuracy is high by several measures. About 98% of the chimpanzee genome sequence has quality scores<sup>25</sup> of at least 40 (Q40), corresponding to an error rate of  $\leq 10^{-4}$ . Comparison of the WGS sequence to 1.3 Mb of finished BACs from the sequenced individual is consistent with this estimate, giving a high-quality discrepancy rate of  $3 \times 10^{-4}$  substitutions and  $2 \times 10^{-4}$  indels, which is no more than expected given the heterozygosity rate (see below), as 50% of the polymorphic alleles in the WGS sequence will differ from the single-haplotype BACs. Comparison of protein-coding regions aligned between the WGS sequence, the recently published sequence of chimpanzee chromosome 21 (ref. 13; formerly chromosome 22 (ref. 18)) and the human genome also revealed no excess of substitutions in the WGS sequence (see Supplementary Information 'Genome sequencing and assembly'). Thus, by restricting our analysis to high-quality bases, the nucleotide-level accuracy of the WGS assembly is essentially equal to that of 'finished' sequence.

Structural accuracy is also high based on comparison with finished BACs from the primary donor and other chimpanzees, although the relatively low level of sequence redundancy limits local contiguity. On the basis of comparisons with the primary donor, some small supercontigs (most <5 kb) have not been positioned within large supercontigs (~1 event per 100 kb); these are not strictly errors but nonetheless affect the utility of the assembly. There are also small, undetected overlaps (all <1 kb) between consecutive contigs (~1.2 events per 100 kb) and occasional local misordering of small contigs (~0.2 events per 100 kb). No misoriented contigs were found. Comparison with the finished chromosome 21 sequence yielded similar discrepancy rates (see Supplementary Information 'Genome sequencing and assembly').

The most problematic regions are those containing recent segmental duplications. Analysis of BAC clones from duplicated ( $n = 75$ ) and unique ( $n = 28$ ) regions showed that the former tend to be fragmented into more contigs (1.6-fold) and more supercontigs (3.2-fold). Discrepancies in contig order are also more frequent in duplicated than unique regions (~0.4 versus ~0.1 events per 100 kb). The rate is twofold higher in duplicated regions with the highest sequence identity (>98%). If we restrict the analysis to older duplications ( $\leq 98\%$  identity) we find fewer assembly problems: 72% of those that can be mapped to the human genome are shared as duplications in both species. These results are consistent with the described limitations of current WGS assembly for regions of segmental duplication<sup>26</sup>. Detailed analysis of these rapidly changing regions of the genome is being performed with more directed approaches<sup>27</sup>.

**Chimpanzee polymorphisms.** The draft sequence of the chimpanzee genome also facilitates genome-wide studies of genetic diversity among chimpanzees, extending recent work<sup>28–31</sup>. We sequenced and analysed sequence reads from the primary donor, four other West African and three central African chimpanzees (*Pan troglodytes troglodytes*) to discover polymorphic positions within and between these individuals (Supplementary Table S17).

A total of 1.66 million high-quality single-nucleotide polymorphisms (SNPs) were identified, of which 1.01 million are heterozygous within the primary donor, Clint. Heterozygosity rates were estimated to be  $9.5 \times 10^{-4}$  for Clint,  $8.0 \times 10^{-4}$  among West African chimpanzees and  $17.6 \times 10^{-4}$  among central African chimpanzees, with the variation between West and central African chimpanzees being  $19.0 \times 10^{-4}$ . The diversity in West African chimpanzees is similar to that seen for human populations<sup>32</sup>, whereas the level for central African chimpanzees is roughly twice as high.

The observed heterozygosity in Clint is broadly consistent with West African origin, although there are a small number of regions of distinctly higher heterozygosity. These may reflect a small amount of central African ancestry, but more likely reflect undetected regions of segmental duplications present only in chimpanzees.

**Table 1 | Chimpanzee assembly statistics**

Assembler	PCAP	ARACHNE
Major contigs*	400,289	361,782
Contig length (kb; N50)†	13.3	15.7
Supercontigs	67,734	37,846
Supercontig length (Mb; N50)	2.3	8.6
Sequence redundancy: all bases (Q20)	$5.0 \times (3.6 \times)$	$4.3 \times (3.6 \times)$
Physical redundancy	20.7	19.8
Consensus bases (Gb)	2.7	2.7

\*Contigs >1 kb.

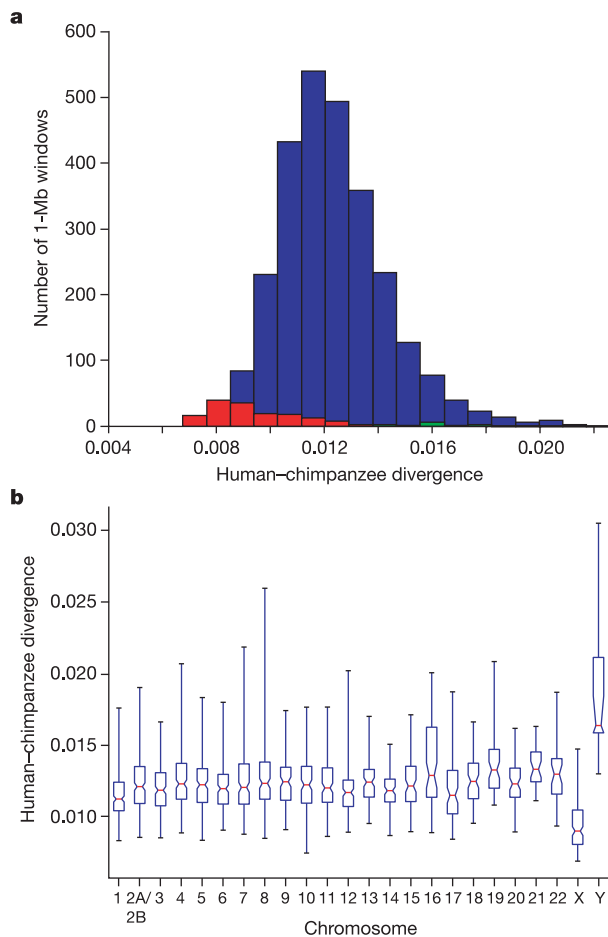
†N50 length is the size  $x$  such that 50% of the assembly is in units of length at least  $x$ .

## Genome evolution

We set out to study the mutational events that have shaped the human and chimpanzee genomes since their last common ancestor. We explored changes at the level of single nucleotides, small insertions and deletions, interspersed repeats and chromosomal rearrangements. The analysis is nearly definitive for the smallest changes, but is more limited for larger changes, particularly lineage-specific segmental duplications, owing to the draft nature of the genome sequence.

**Nucleotide divergence.** Best reciprocal nucleotide-level alignments of the chimpanzee and human genomes cover ~2.4 gigabases (Gb) of high-quality sequence, including 89 Mb from chromosome X and 7.5 Mb from chromosome Y.

**Genome-wide rates.** We calculate the genome-wide nucleotide divergence between human and chimpanzee to be 1.23%, confirming recent results from more limited studies<sup>12,33,34</sup>. The differences between one copy of the human genome and one copy of the chimpanzee genome include both the sites of fixed divergence between the species and some polymorphic sites within each species. By correcting for the estimated coalescence times in the human and chimpanzee populations (see Supplementary Information ‘Genome evolution’), we estimate that polymorphism accounts for 14–22% of the observed divergence rate and thus that the fixed divergence is ~1.06% or less.



**Figure 1 | Human-chimpanzee divergence in 1-Mb segments across the genome.** **a**, Distribution of divergence of the autosomes (blue), the X chromosome (red) and the Y chromosome (green). **b**, Distribution of variation by chromosome, shown as a box plot. The edges of the box correspond to quartiles; the notches to the standard error of the median; and the vertical bars to the range. The X and Y chromosomes are clear outliers, but there is also high local variation within each of the autosomes.

Nucleotide divergence rates are not constant across the genome, as has been seen in comparisons of the human and murid genomes<sup>16,17,24,35,36</sup>. The average divergence in 1-Mb segments fluctuates with a standard deviation of 0.25% (coefficient of variation = 0.20), which is much greater than the 0.02% expected assuming a uniform divergence rate (Fig. 1a; see also Supplementary Fig. S1).

Regional variation in divergence could reflect local variation in either mutation rate or other evolutionary forces. Among the latter, one important force is genetic drift, which can cause substantial differences in divergence time across loci when comparing closely related species, as the divergence time for orthologues is the sum of two terms:  $t_1$ , the time since speciation, and  $t_2$ , the coalescence time for orthologues within the common ancestral population<sup>37</sup>. Whereas  $t_1$  is constant across loci (~6–7 million years<sup>38</sup>),  $t_2$  is a random variable that fluctuates across loci (with a mean that depends on population size and here may be on the order of 1–2 million years<sup>39</sup>). However, because of historical recombination, the characteristic scale of such fluctuations will be on the order of tens of kilobases, which is too small to account for the variation observed for 1-Mb regions<sup>40</sup> (see Supplementary Information ‘Genome evolution’). Other potential evolutionary forces are positive or negative selection. Although it is more difficult to quantify the expected contributions of selection in the ancestral population<sup>41–43</sup>, it is clear that the effects would have to be very strong to explain the large-scale variation observed across mammalian genomes<sup>16,44</sup>. There is tentative evidence from in-depth analysis of divergence and diversity that natural selection is not the major contributor to the large-scale patterns of genetic variability in humans<sup>45–47</sup>. For these reasons, we suggest that the large-scale variation in the human–chimpanzee divergence rate primarily reflects regional variation in mutation rate.

**Chromosomal variation in divergence rate.** Variation in divergence rate is evident even at the level of whole chromosomes (Fig. 1b). The most striking outliers are the sex chromosomes, with a mean divergence of 1.9% for chromosome Y and 0.94% for chromosome X. The likely explanation is a higher mutation rate in the male compared with female germ line<sup>48</sup>. Indeed, the ratio of the male/female mutation rates (denoted  $\alpha$ ) can be estimated by comparing the divergence rates among the sex chromosomes and the autosomes and correcting for ancestral polymorphism as a function of population size of the most recent common ancestor (MRCA; see Supplementary Information ‘Genome evolution’). Estimates for  $\alpha$  range from 3 to 6, depending on the chromosomes compared and the assumed ancestral population size (Supplementary Table S18). This is significantly higher than recent estimates of  $\alpha$  for the murids (~1.9) (ref. 17) and resolves a recent controversy based on smaller data sets<sup>12,24,49,50</sup>.

The higher mutation rate in the male germ line is generally attributed to the 5–6-fold higher number of cell divisions undergone by male germ cells<sup>48</sup>. We reasoned that this would affect mutations resulting from DNA replication errors (the rate should scale with the number of cell divisions) but not mutations resulting from DNA damage such as deamination of methyl CpG to TpG (the rate should scale with time). Accordingly, we calculated  $\alpha$  separately for CpG sites, obtaining a value of ~2 from the comparison of rates between autosomes and chromosome X. This intermediate value is a composite of the rates of CpG loss and gain, and is consistent with roughly equal rates of CpG to TpG transitions in the male and female germ line<sup>51,52</sup>.

Significant variation in divergence rates is also seen among autosomes (Fig. 1b;  $P < 3 \times 10^{-15}$ , Kruskal–Wallis test over 1-Mb windows), confirming earlier observations based on low-coverage WGS sampling<sup>12</sup>. Additional factors thus influence the rate of divergence between chimpanzee and human chromosomes. These factors are likely to act at length scales significantly shorter than a chromosome, because the standard deviation across autosomes (0.21%) is comparable to the standard deviation seen in 1-Mb windows across the genome (0.13–0.35%). We therefore sought to

understand local factors that contribute to variation in divergence rate.

**Contribution of CpG dinucleotides.** Sites containing CpG dinucleotides in either species show a substantially elevated divergence rate of 15.2% per base; they account for 25.2% of all substitutions while constituting only 2.1% of all aligned bases. The divergence at CpG sites represents both the loss of ancestral CpGs and the creation of new CpGs. The former process is known to occur at a rapid rate per base due to frequent methylation of cytosines in a CpG context and their frequent deamination<sup>53,54</sup>, whereas the latter process probably proceeds at a rate more typical of other nucleotide substitutions. Assuming that loss and creation of CpG sites are close to equilibrium, the mutation rate for bases in a CpG dinucleotide must be 10–12-fold higher than for other bases (see Supplementary Information ‘Genome evolution’ and ref. 51).

Because of the high rate of CpG substitutions, regional divergence rates would be expected to correlate with regional CpG density. CpG density indeed varies across 1-Mb windows (mean = 2.1%, coefficient of variation = 0.44 compared with 0.0093 expected under a Poisson distribution), but only explains 4% of the divergence rate variance. In fact, regional CpG and non-CpG divergence is highly correlated ( $r = 0.88$ ; Supplementary Fig. S2), suggesting that higher-order effects modulate the rates of two very different mutation processes (see also ref. 47).

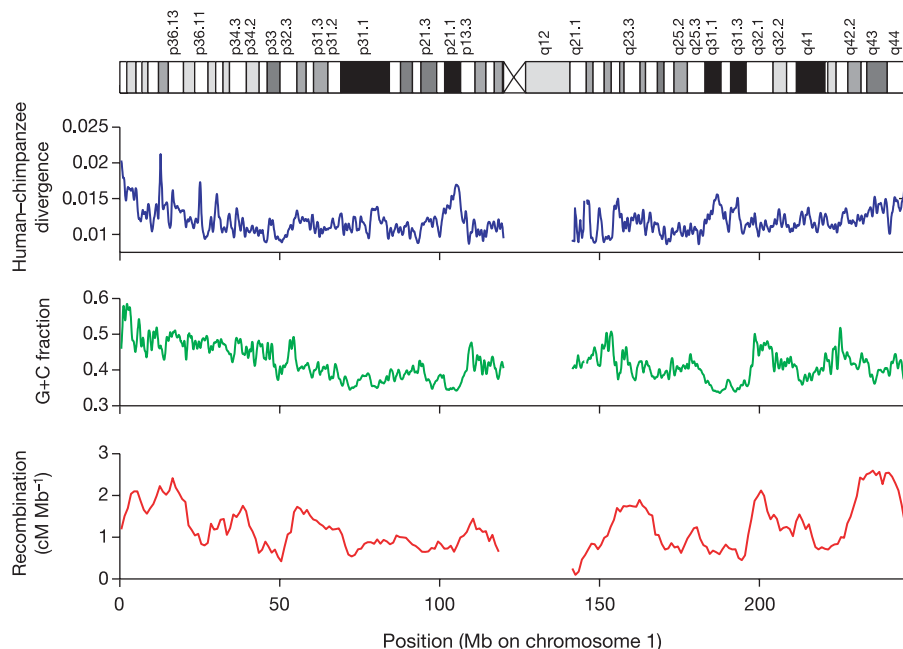
**Increased divergence in distal regions.** The most striking regional pattern is a consistent increase in divergence towards the ends of most chromosomes (Fig. 2). The terminal 10 Mb of chromosomes (including distal regions and proximal regions of acrocentric chromosomes) averages 15% higher divergence than the rest of the genome (Mann–Whitney  $U$ -test;  $P < 10^{-30}$ ), with a sharp increase towards the telomeres. The phenomenon correlates better with physical distance than relative position along the chromosomes and may partially explain why smaller chromosomes tend to have higher divergence (Supplementary Fig. S3; see also ref. 15). These observations suggest that large-scale chromosomal structure, directly or indirectly, influences regional divergence patterns. The cause of this effect is unclear, but these regions (~15% of the genome) are

notable in having high local recombination rate, high gene density and high G + C content.

**Correlation with chromosome banding.** Another interesting pattern is that divergence increases with the intensity of Giemsa staining in cytogenetically defined chromosome bands, with the regions corresponding to Giemsa dark bands (G bands) showing 10% higher divergence than the genome-wide average (Mann–Whitney  $U$ -test;  $P < 10^{-14}$ ) (see Fig. 2). In contrast to terminal regions, these regions (17% of the genome) tend to be gene poor, (G + C)-poor and low in recombination<sup>55,56</sup>. The elevated divergence seen in two such different types of regions suggests that multiple mechanisms are at work, and that no single known factor, such as G + C content or recombination rate, is an adequate predictor of regional variation in the mammalian genome by itself (Fig. 3). Elucidation of the relative contributions of these and other mechanisms will be important for formulating accurate models for population genetics, natural selection, divergence times and the evolution of genome-wide sequence composition<sup>57</sup>.

**Correlation with regional variation in the murid genome.** Given that sequence divergence shows regional variation in both hominids (human–chimpanzee) and murids (mouse–rat), we asked whether the regional rates are positively correlated between orthologous regions. Such a correlation would suggest that the divergence rate is driven, in part, by factors that have been conserved over the ~75 million years since rodents, humans and apes shared a common ancestor. Comparative analysis of the human and murid genomes has suggested such a correlation<sup>58–60</sup>, but the chimpanzee sequence provides a direct opportunity to compare independent evolutionary processes between two mammalian clades.

We compared the local divergence rates in hominids and murids across major orthologous segments in the respective genomes (Fig. 4). For orthologous segments that are non-distal in both hominids and murids, there is a strong correlation between the divergence rates ( $r = 0.5$ ,  $P < 10^{-11}$ ). In contrast, orthologous segments that are centred within 10 Mb of a hominid telomere have disproportionately high divergence rates and G + C content relative to the murids (Mann–Whitney  $U$ -test;  $P < 10^{-11}$  and



**Figure 2 | Regional variation in divergence rates.** Human–chimpanzee divergence (blue), G + C content (green) and human recombination rates<sup>173</sup> (red) in sliding 1-Mb windows for human and chimpanzee chromosome 1. Divergence and G + C content are noticeably elevated near the 1p telomere,

a trend that holds for most subtelomeric regions (see text). Internally on the chromosome, regions of low G + C content and high divergence often correspond to the dark G bands.



$P < 10^{-4}$ ), implying that the elevation in these regions is, at least partially, lineage specific. The same general effect is observed (albeit less pronounced) if CpG dinucleotides are excluded (Supplementary Fig. S4). Increased divergence and G + C content might be explained by 'biased gene conversion'<sup>61</sup> due to the high hominid recombination rates in these distal regions. Segments that are distal in murids do not show elevated divergence rates, which is consistent with this model, because the recombination rates of distal regions are not as elevated in mouse and rat<sup>62</sup>.

Taken together, these observations suggest that sequence divergence rate is influenced by both conserved factors (stable across mammalian evolution) and lineage-specific factors (such as proximity to the telomere or recombination rate, which may change with chromosomal rearrangements).

**Insertions and deletions.** We next studied the indel events that have occurred in the human and chimpanzee lineages by aligning the genome sequences to identify length differences. We will refer below to all events as insertions relative to the other genome, although they may represent insertions or deletions relative to the genome of the common ancestor.

The observable insertions fall into two classes: (1) 'completely covered' insertions, occurring within continuous sequence in both species; and (2) 'incompletely covered' insertions, occurring within sequence containing one or more gaps in the chimpanzee, but revealed by a clear discrepancy between the species in sequence length. Different methods are needed for reliable identification of modest-sized insertions (1 base to 15 kb) and large insertions (>15 kb), with the latter only being reliably identifiable in the human genome (see Supplementary Information 'Genome evolution').

The analysis of modest-sized insertions reveals ~32 Mb of human-specific sequence and ~35 Mb of chimpanzee-specific sequence, contained in ~5 million events in each species (Supplementary Information 'Genome evolution' and Supplementary Fig. S5). Nearly all of the human insertions are completely covered, whereas only half of the chimpanzee insertions are completely covered. Analysis of the completely covered insertions shows that the vast majority are small (45% of events cover only 1 base pair (bp), 96% are <20 bp and 98.6% are <80 bp), but that the largest few contain most of the

sequence (with the ~70,000 indels larger than 80 bp comprising 73% of the affected base pairs) (Fig. 5). The latter indels >80 bp fall into three categories: (1) about one-quarter are newly inserted transposable elements; (2) more than one-third are due to microsatellite and satellite sequences; (3) and the remainder are assumed to be mostly deletions in the other genome.

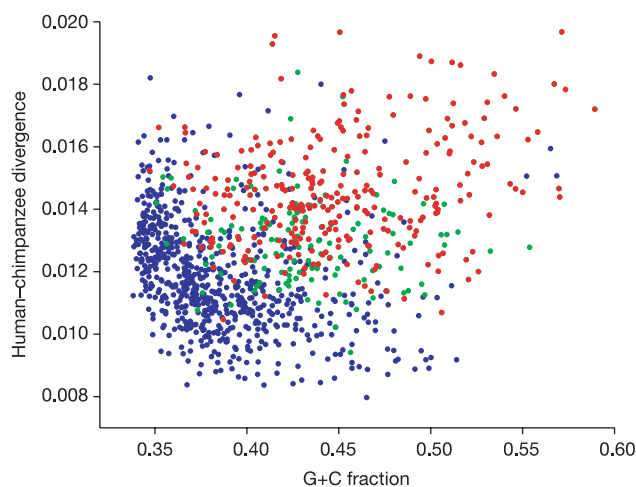
The analysis of larger insertions (>15 kb) identified 163 human regions containing 8.3 Mb of human-specific sequence in total (Fig. 6). These cases include 34 regions that involve exons from known genes, which are discussed in a subsequent section. Although we have no direct measure of large insertions in the chimpanzee genome, it appears likely that the situation is similar.

On the basis of this analysis, we estimate that the human and chimpanzee genomes each contain 40–45 Mb of species-specific euchromatic sequence, and the indel differences between the genomes thus total ~90 Mb. This difference corresponds to ~3% of both genomes and dwarfs the 1.23% difference resulting from nucleotide substitutions; this confirms and extends several recent studies<sup>63–67</sup>. Of course, the number of indel events is far fewer than the number of substitution events (~5 million compared with ~35 million, respectively).

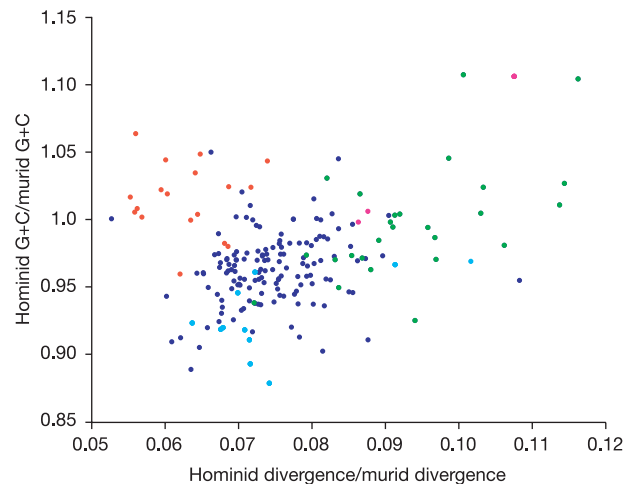
**Transposable element insertions.** We next used the catalogue of lineage-specific transposable element copies to compare the activity of transposons in the human and chimpanzee lineages (Table 2).

**Endogenous retroviruses.** Endogenous retroviruses (ERVs) have become all but extinct in the human lineage, with only a single retrovirus (human endogenous retrovirus K (HERV-K)) still active<sup>24</sup>. HERV-K was found to be active in both lineages, with at least 73 human-specific insertions (7 full length and 66 solo long terminal repeats (LTRs)) and at least 45 chimpanzee-specific insertions (1 full length and 44 solo LTRs). A few other ERV classes persisted in the human genome beyond the human–chimpanzee split, leaving ~9 human-specific insertions (all solo LTRs, including five HERV9 elements) before dying out.

Against this background, it was surprising to find that the chimpanzee genome has two active retroviral elements (PtERV1 and PtERV2) that are unlike any older elements in either genome;



**Figure 3 | Divergence rates versus G + C content for 1-Mb segments across the autosomes.** Conditional on recombination rate, the relationship between divergence and G + C content varies. In regions with recombination rates less than  $0.8 \text{ cM Mb}^{-1}$  (blue), there is an inverse relationship, where high divergence regions tend to be (G + C)-poor and low divergence regions tend to be (G + C)-rich. In regions with recombination rates greater than  $2.0 \text{ cM Mb}^{-1}$ , whether within 10 Mb (red) or proximal (green) of chromosome ends, both divergence and G + C content are uniformly high.



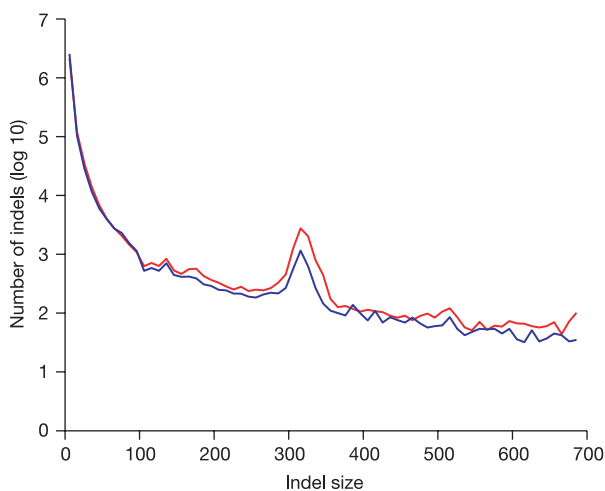
**Figure 4 | Disproportionately elevated divergence and G + C content near hominid telomeres.** Scatter plot of the ratio of human–chimpanzee divergence over mouse–rat divergence versus the ratio of human G + C content over mouse G + C content across 199 syntenic blocks for which more than 1 Mb of sequence could be aligned between all four species. Blocks for which the centre is within 10 Mb of a telomere in hominids only (green) or in hominids and murids (magenta), but not in murids only (light green), show a significant trend towards higher ratios than internal blocks (dark blue). Blocks on the X chromosome (red) tend to show a lower divergence ratio than autosomal blocks, consistent with a smaller difference between autosomal and X divergence in murids than in hominids (lower  $\alpha$ ).

these must have been introduced by infection of the chimpanzee germ line. The smaller family (PtERV2) has only a few dozen copies, which nonetheless represent multiple (~5–8) invasions, because the sequence differences among reconstructed subfamilies are too great (~8%) to have arisen by mutation since divergence from human. It is closely related to a baboon endogenous retrovirus (BaEV, 88% ORF2 product identity) and a feline endogenous virus (ECE-1, 86% ORF2 product identity). The larger family (PtERV1) is more homogeneous and has over 200 copies. Whereas older ERVs, like HERV-K, are primarily represented by solo LTRs resulting from LTR–LTR recombination, more than half of the PtERV1 copies are still full length, probably reflecting the young age of the elements. PtERV1-like elements are present in the rhesus monkey, olive baboon and African great apes but not in human, orang-utan or gibbon, suggesting separate germline invasions in these species<sup>68</sup>.

**Higher Alu activity in humans.** SINE (Alu) elements have been threefold more active in humans than chimpanzee (~7,000 compared with ~2,300 lineage-specific copies in the aligned portion), refining the rather broad range (2–7-fold) estimated in smaller studies<sup>13,67,69</sup>. Most chimpanzee-specific elements belong to a subfamily (AluYc1) that is very similar to the source gene in the common ancestor. By contrast, most human-specific Alu elements belong to two new subfamilies (AluYa5 and AluYb8) that have evolved since the chimpanzee–human divergence and differ substantially from the ancestral source gene<sup>69</sup>. It seems likely that the resurgence of Alu elements in humans is due to these potent new source genes. However, based on an examination of available finished sequence, the baboon shows a 1.6-fold higher Alu activity relative to human new insertions, suggesting that there may also have been a general decline in activity in the chimpanzee<sup>67</sup>.

Some of the human-specific Alu elements are highly diverged (92 with >5% divergence), which would seem to suggest that they are much older than the human–chimpanzee split. Possible explanations include: gene conversion by nearby older elements; processed pseudogenes arising from a spurious transcription of an older element; precise excision from the chimpanzee genome; or high local mutation rate. In any case, the presence of such anomalies suggests that caution is warranted in the use of single-repeat elements as homoplasy-free phylogenetic markers.

**New Alu elements target (A + T)-rich DNA in human and chimpanzee genomes.** Older SINE elements are preferentially found in gene-rich,



**Figure 5 | Length distribution of small indel events, as determined using bounded sequence gaps.** Sequences present in chimpanzee but not in human (blue) or present in human but not in chimpanzee (red) are shown. The prominent spike around 300 nucleotides corresponds to SINE insertion events. Most of the indels are smaller than 20 bp, but larger indels account for the bulk of lineage-specific sequence in the two genomes.

(G + C)-rich regions, whereas younger SINE elements are found in gene-poor, (A + T)-rich regions where long interspersed element (LINE)-1 (L1) copies also accumulate<sup>24,70</sup>. The latter distribution is consistent with the fact that Alu retrotransposition is mediated by L1 (ref. 71). Murid genomes revealed no change in SINE distribution with age<sup>17</sup>.

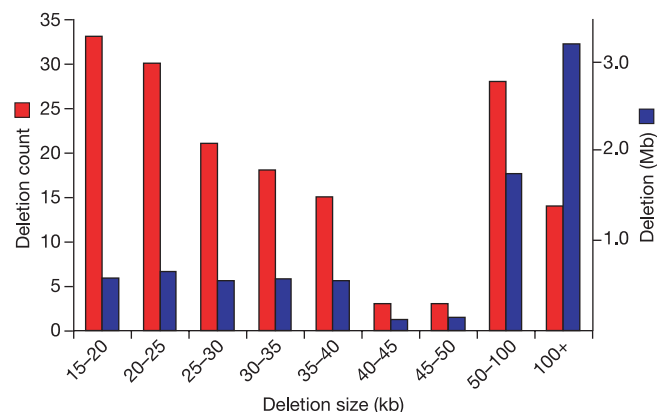
The human pattern might reflect either preferential retention of SINEs in (G + C)-rich regions, due to selection or mutation bias, or a recent change in Alu insertion preferences. With the availability of the chimpanzee genome, it is possible to classify the youngest Alu copies more accurately and thus begin to distinguish these possibilities.

Analysis shows that lineage-specific SINEs in both human and chimpanzee are biased towards (A + T)-rich regions, as opposed to even the most recent copies in the MRCA (Fig. 7). This indicates that SINEs are indeed preferentially retained in (G + C)-rich DNA, but comparison with a more distant primate is required to formally rule out the possibility that the insertion bias of SINEs did not change just before speciation.

**Equal activity of L1 in both species.** The human and chimpanzee genomes both show ~2,000 lineage-specific L1 elements, contrary to previous estimates based on small samples that L1 activity is 2–3-fold higher in chimpanzee<sup>72</sup>.

Transcription from L1 source genes can sometimes continue into 3' flanking regions, which can then be co-transposed<sup>73,74</sup>. Human–chimpanzee comparison revealed that ~15% of the species-specific insertions appear to have carried with them at least 50 bp of flanking sequence (followed by a poly(A) tail and a target site duplication). In principle, incomplete reverse transcription could result in insertions of the flanking sequence only (without any L1 sequence), mobilizing gene elements such as exons, but we found no evidence of this.

**Retrotransposed gene copies.** The L1 machinery also mediates retrotransposition of host messenger RNAs, resulting in many intronless (processed) pseudogenes in the human genome<sup>75–77</sup>. We identified 163 lineage-specific retrotransposed gene copies in human and 246 in chimpanzee (Supplementary Table S19). Correcting for incomplete sequence coverage of the chimpanzee genome, we estimate that there are ~200 and ~300 processed gene copies in human and chimpanzee, respectively. Processed genes thus appear to have arisen at a rate of ~50 per million years since the divergence of human and chimpanzee; this is lower than the estimated rate for early primate evolution<sup>75</sup>, perhaps reflecting the overall decrease in L1 activity. As expected<sup>78</sup>, ribosomal protein genes constitute the largest class in both species. The second largest class in chimpanzee corresponds to zinc finger C2H2 genes, which are not a major class in the human genome.



**Figure 6 | Length distribution of large indel events (>15 kb), as determined using paired-end sequences from chimpanzee mapped against the human genome.** Both the total number of candidate human insertions/chimpanzee deletions (blue) and the number of bases altered (red) are shown.

*The retrotransposon SVA and distribution of CpG islands by transposable elements.* The third most active element since speciation has been SVA, which created about 1,000 copies in each lineage. SVA is a composite element (~1.5–2.5 kb) consisting of two Alu fragments, a tandem repeat and a region apparently derived from the 3' end of a HERV-K transcript; it is probably mobilized by L1 (refs 79, 80). This element is of particular interest because each copy carries a sequence that satisfies the definition of a CpG island<sup>81</sup> and contains potential transcription factor binding sites; the dispersion of 1,000 SVA copies could therefore be a source of regulatory differences between chimpanzee and human (Supplementary Table S20). At least three human genes contain SVA insertions near their promoters (Supplementary Table S21), one of which has been found to be differentially expressed between the two species<sup>82,83</sup>, but additional investigations will be required to determine whether the SVA insertion directly caused this difference.

*Homologous recombination between interspersed repeats.* Human–chimpanzee comparison also makes it possible to study homologous recombination between nearby repeat elements as a source of genomic deletions. We found 612 deletions (totalling 2 Mb) in the human genome that appear to have resulted from recombination between two nearby Alu elements present in the common ancestor; there are 914 such events in the chimpanzee genome. (The events are not biased to (A + T)-rich DNA and thus would not explain the preferential loss of Alu elements in such regions discussed above.) Similarly, we found 26 and 48 instances involving adjacent L1 copies and 8 and 22 instances involving retroviral LTRs in human and chimpanzee, respectively. None of the repeat-mediated deletions removed an orthologous exon of a known human gene in chimpanzee.

The genome comparison allows one to estimate the dependency of homologous recombination on divergence and distance. Homologous recombination seems to occur between quite (>25%) diverged copies (Fig. 8), whereas the number of recombination events ( $n$ ) varies inversely with the distance ( $d$ , in bases) between the copies (as  $n \approx 6 \times 10^6 d^{-1.7}$ ;  $r^2 = 0.9$ ).

**Large-scale rearrangements.** Finally, we examined the chimpanzee genome sequence for information about large-scale genomic alterations. Cytogenetic studies have shown that human and chimpanzee chromosomes differ by one chromosomal fusion, at least nine pericentric inversions, and in the content of constitutive heterochromatin<sup>84</sup>. Human chromosome 2 resulted from a fusion of two ancestral chromosomes that remained separate in the chimpanzee lineage (chromosomes 2A and 2B in the revised nomenclature<sup>18</sup>, formerly chimpanzee chromosomes 12 and 13); the precise fusion point has been mapped and its duplication structure described in detail<sup>85,86</sup>. In accord with this, alignment of the human and chimpanzee genome sequences shows a break in continuity at this point.

We searched the chimpanzee genome sequence for the precise locations of the 18 breakpoints corresponding to the 9 pericentric inversions (Supplementary Table S22). By mapping paired-end sequences from chimpanzee large insert clones to the human genome, we were able to identify 13 of the breakpoints within the

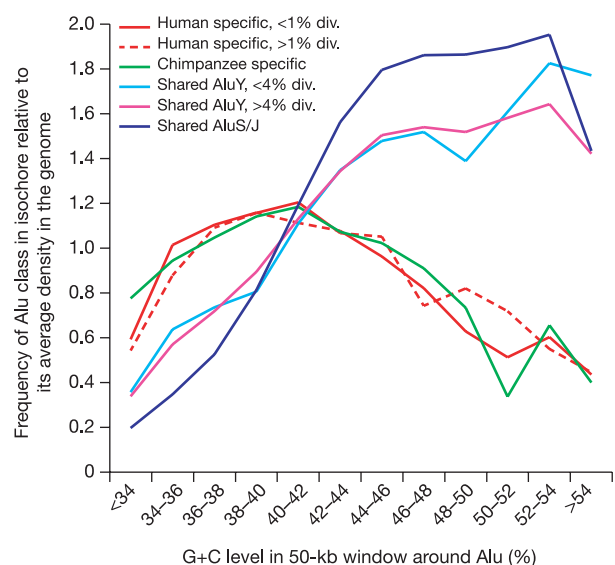
assembly from discordant end alignments. The positions of five breakpoints (on chromosomes 4, 5 and 12) were tested by fluorescence *in situ* hybridization (FISH) analysis and all were confirmed. Also, the positions of three previously mapped inversion breakpoints (on chromosomes 15 and 18) matched closely those found in the assembly<sup>87,88</sup>. The paired-end analysis works well in regions of unique sequence, which constitute the bulk of the genome, but is less effective in regions of recent duplication owing to ambiguities in mapping of the paired-end sequences. Beyond the known inversions, we also found suggestive evidence of many additional smaller inversions, as well as older segmental duplications (<98% identity; Supplementary Fig. S6). However, both smaller inversions and more recent segmental duplications will require further investigations.

## Gene evolution

We next sought to use the chimpanzee sequence to study the role of natural selection in the evolution of human protein-coding genes. Genome-wide comparisons can shed light on many central issues, including: the magnitude of positive and negative selection; the variation in selection across different lineages, chromosomes, gene families and individual genes; and the complete loss of genes within a lineage.

We began by identifying a set of 13,454 pairs of human and chimpanzee genes with unambiguous 1:1 orthology for which it was possible to generate high-quality sequence alignments covering virtually the entire coding region (Supplementary Information 'Gene evolution' and Table S23). The list contains a large fraction of the entire complement of human genes, although it underrepresents gene families that have undergone recent local expansion (such as olfactory receptors and immunoglobulins). To facilitate comparison with the murid lineage, we also compiled a set of 7,043 human, chimpanzee, mouse and rat genes with unambiguous 1:1:1:1 orthology and high-quality sequence alignments (Supplementary Table S24).

**Average rates of evolution.** To assess the rate of evolution for each gene, we estimated  $K_A$ , the number of coding base substitutions that result in amino acid change as a fraction of all such possible sites (the non-synonymous substitution rate). Because the background



**Figure 7 | Correlation of Alu age and distribution by G + C content.** Alu elements that inserted after human–chimpanzee divergence are densest in the (G + C)-poor regions of the genome (peaking at 36–40% G + C), whereas older copies, common to both genomes, crowd (G + C)-rich regions. The figure is similar to figure 23 of ref. 24, but the use of chimpanzee allows improved separation of young and old elements, leading to a sharper transition in the pattern.

**Table 2 | Transposable element activity in human and chimpanzee lineages**

Element	Chimpanzee*	Human*
Alu	2,340 (0.7 Mb)	7,082 (2.1 Mb)
LINE-1	1,979 (>5 Mb)	1,814 (5.0 Mb)
SVA	757 (>1 Mb)	970 (1.3 Mb)
ERV class 1	234 (>1 Mb)†	5 (8 kb)‡
ERV class 2	45 (55 kb)§	77 (130 kb)§
(Micro)satellite	7,054 (4.1 Mb)	11,101 (5.1 Mb)

\*Number of lineage-specific insertions (with total size of inserted sequences indicated in brackets) in the aligned parts of the genomes.

†P1ERV1 and P1ERV2.

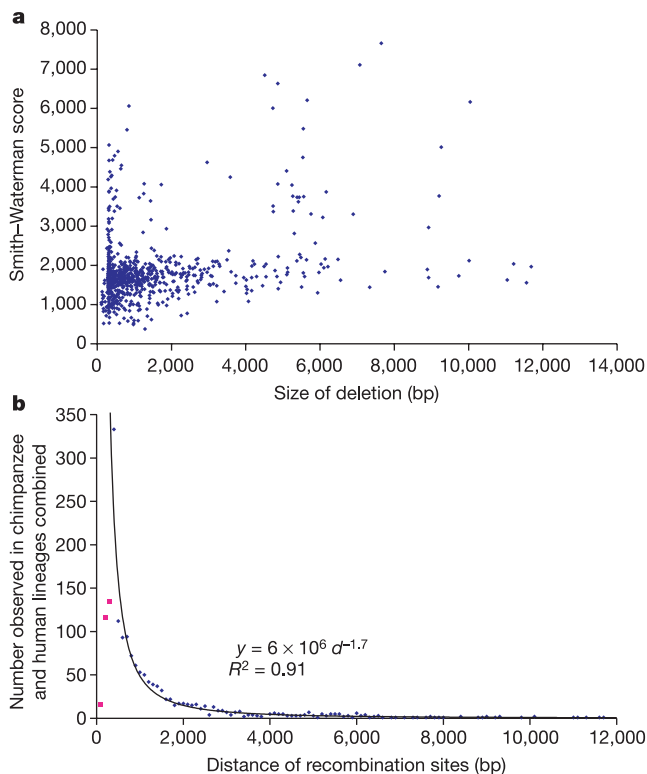
‡HERV9.

§Mostly HERV-K.

mutation rate varies across the genome, it is crucial to normalize  $K_A$  for comparisons between genes. A striking illustration of this variation is the fact that the mean  $K_A$  is 37% higher in the rapidly diverging distal 10 Mb of chromosomes than in the more proximal regions. Classically, the background rate is estimated by  $K_S$ , the synonymous substitution rate (coding base substitutions that, because of codon redundancy, do not result in amino acid change). Because a typical gene has only a few synonymous changes between humans and chimpanzees, and not infrequently is zero, we exploited the genome sequence to estimate the local intergenic/intronic substitution rate,  $K_I$ , where appropriate.  $K_A$  and  $K_S$  were also estimated for each lineage separately using mouse and rat as outgroups (Fig. 9).

The  $K_A/K_S$  ratio is a classical measure of the overall evolutionary constraint on a gene, where  $K_A/K_S \ll 1$  indicates that a substantial proportion of amino acid changes must have been eliminated by purifying selection. Under the assumption that synonymous substitutions are neutral,  $K_A/K_S > 1$  implies, but is not a necessary condition for, adaptive or positive selection. The  $K_A/K_I$  ratio has the same interpretation. The ratios will sometimes be denoted below by  $\omega$  with an appropriate subscript (for example,  $\omega_{\text{human}}$ ) to indicate the branch of the evolutionary tree under study.

**Evolutionary constraint on amino acid sites within the hominid lineage.** Overall, human and chimpanzee genes are extremely similar, with the encoded proteins identical in the two species in 29% of cases. The median number of non-synonymous and synonymous substitutions per gene are two and three, respectively. About 5% of the proteins show in-frame indels, but these tend to be small (median = 1 codon)



**Figure 8 | Dependency of homologous recombination between Alu elements on divergence and distance.** **a**, Whereas homologous recombination occurs between quite divergent (Smith–Waterman score <1,000), closely spaced copies, more distant recombination seems to favour a better match between the recombining repeats. **b**, The frequency of Alu–Alu-mediated recombination falls markedly as a function of distance between the recombining copies. The first three points (magenta) involve recombination between left or right arms of one Alu inserted into another. The high number of occurrences at a distance of 300–400 nucleotides is due to the preference of integration in the A-rich tail; exclusion of this point does not change the parameters of the equation.

and to occur in regions of repeated sequence. The close similarity of human and chimpanzee genes necessarily limits the ability to make strong inferences about individual genes, but there is abundant data to study important sets of genes.

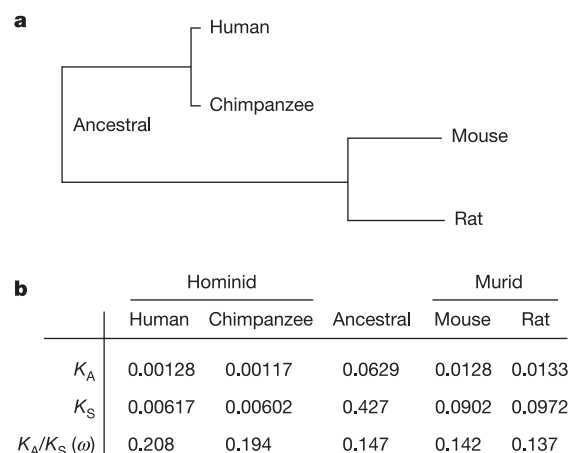
The  $K_A/K_S$  ratio for the human–chimpanzee lineage ( $\omega_{\text{hominid}}$ ) is 0.23. The value is much lower than some recent estimates based on limited sequence data (ranging as high as 0.63 (ref. 7)), but is consistent with an estimate (0.22) from random expressed-sequence-tag (EST) sequencing<sup>45</sup>. Similarly,  $K_A/K_I$  was also estimated as 0.23.

Under the assumption that synonymous mutations are selectively neutral, the results imply that 77% of amino acid alterations in hominid genes are sufficiently deleterious as to be eliminated by natural selection. Because synonymous mutations are not entirely neutral (see below), the actual proportion of amino acid alterations with deleterious consequences may be higher. Consistent with previous studies<sup>8</sup>, we find that  $K_A/K_S$  of human polymorphisms with frequencies up to 15% is significantly higher than that of human–chimpanzee differences and more common polymorphisms (Table 3), implying that at least 25% of the deleterious amino acid alterations may often attain readily detectable frequencies and thus contribute significantly to the human genetic load.

**Evolutionary constraint on synonymous sites within hominid lineage.** We next explored the evolutionary constraints on synonymous sites, specifically fourfold degenerate sites. Because such sites have no effect on the encoded protein, they are often considered to be selectively neutral in mammals.

We re-examined this assumption by comparing the divergence at fourfold degenerate sites with the divergence at nearby intronic sites. Although overall divergence rates are very similar at fourfold degenerate and intronic sites, direct comparison is misleading because the former have a higher frequency of the highly mutable CpG dinucleotides (9% compared with 2%). When CpG and non-CpG sites are considered separately, we find that both CpG sites and non-CpG sites show markedly lower divergence in exonic synonymous sites than in introns (~50% and ~30% lower, respectively). This result resolves recent conflicting reports based on limited data sets<sup>45,89</sup> by showing that such sites are indeed under constraint.

The constraint does not seem to result from selection on the usage of preferred codons, which has been detected in lower organisms<sup>90</sup> such as bacteria<sup>91</sup>, yeast<sup>92</sup> and flies<sup>93</sup>. In fact, divergence at fourfold



**Figure 9 | Human–chimpanzee–mouse–rat tree with branch-specific  $K_A/K_S$  ( $\omega$ ) values.** **a**, Evolutionary tree. The branch lengths are proportional to the absolute rates of amino acid divergence. **b**, Maximum-likelihood estimates of the rates of evolution in protein-coding genes for humans, chimpanzees, mice and rats. In the text,  $\omega_{\text{hominid}}$  is the  $K_A/K_S$  of the combined human and chimpanzee branches and  $\omega_{\text{murid}}$  of the combined mouse and rat branches. The slight difference between  $\omega_{\text{human}}$  and  $\omega_{\text{chimpanzee}}$  is not statistically significant; masking of some heterozygous bases in the chimpanzee sequence may contribute to the observed difference (see Supplementary Information ‘Gene evolution’).



degenerate sites increases slightly with codon usage bias (Kendall's  $\tau = 0.097$ ,  $P < 10^{-14}$ ). Alternatively, the observed constraint at synonymous sites might reflect 'background selection'—that is, the indirect effect of purifying selection at amino acid sites causing reduced diversity and thereby reduced divergence at closely linked sites<sup>42</sup>. Given the low rate of recombination in hominid genomes (a 1 kb region experiences only  $\sim 1$  crossover per 100,000 generations or 2 million years), such background selection should extend beyond exons to include nearby intronic sites<sup>94</sup>. However, when the divergence rate is plotted relative to exon–intron boundaries, we find that the rate jumps sharply within a short region of  $\sim 7$  bp at the boundary (Fig. 10). This pattern strongly suggests that the action of purifying selection at synonymous sites is direct rather than indirect, suggesting that other signals, for example those involved in splice site selection, may be embedded in the coding sequence and therefore constrain synonymous sites.

**Comparison with murids.** An accurate estimate of  $K_A/K_S$  makes it possible to study how evolutionary constraint varies across clades. It was predicted more than 30 years ago<sup>95</sup> that selection against deleterious mutations would depend on population size, with mutations being strongly selected only if they reduce fitness by  $s \gg 1/4N$  (where  $N$  is effective population size). This would predict that genes would be under stronger purifying selection in murids than hominids, owing to their presumed larger population size. Initial analyses (involving fewer than 50 genes<sup>96</sup>) suggested a strong effect, but the wide variation in estimates of  $K_A/K_S$  in hominids<sup>7,8,97</sup> and murids<sup>98</sup> has complicated this analysis<sup>45</sup>.

Using the large collection of 7,043 orthologous quartets, we calculated mean  $K_A/K_S$  values for the various branches of the four-species evolutionary tree (human, chimpanzee, mouse and rat; Fig. 9). The  $K_A/K_S$  ratio for hominids is 0.20. (This is slightly lower than the value of 0.23 obtained with all human–chimpanzee orthologues, probably reflecting slightly greater constraint on the class of proteins with clear orthologues across hominids and murids.)

The  $K_A/K_S$  ratio is markedly lower for murids than for hominids ( $\omega_{\text{murid}} \approx 0.13$  compared with  $\omega_{\text{hominid}} \approx 0.20$ ) (Fig. 9). This implies that there is an  $\sim 35\%$  excess of the amino-acid-changing mutations in the two hominids, relative to the two murids. Excess amino acid divergence may be explained by either increased adaptive evolution or relaxation of evolutionary constraints. As shown in the next section, the latter seems to be the principal explanation.

**Relaxed constraints in human evolution.** The  $K_A/K_S$  ratio can be used to make inferences about the role of positive selection in human evolution<sup>99,100</sup>. Because alleles under positive selection spread rapidly through a population, they will be found less frequently as common human polymorphisms than as human–chimpanzee differences<sup>8</sup>. Positive selection can thus be detected by comparing the  $K_A/K_S$  ratio for common human polymorphisms with the  $K_A/K_S$  ratio for

hominid divergence. These ratios have been estimated as  $\omega_{\text{polymorphism}} \approx 0.20$  based on an initial collection of common SNPs in human genes and  $\omega_{\text{divergence}} \approx 0.34$  based on comparison of human and Old World monkey genes<sup>8</sup>. Thus, the proportion of amino acid changes attributable to positive selection was inferred to be  $\sim 35\%$  (ref. 8). This would imply a huge quantitative role for positive selection in human evolution.

With the availability of extensive data for both human polymorphism and human–chimpanzee divergence, we repeated this analysis (using the same set of genes for both estimates). We find that  $\omega_{\text{polymorphism}} \approx 0.21$ – $0.23$  and  $\omega_{\text{divergence}} \approx 0.23$  are statistically indistinguishable (Table 3). Although some of the amino acid substitutions in human and chimpanzee evolution must surely reflect positive selection, the results indicate that the proportion of changes fixed by positive selection seems to be much lower than the previous estimate<sup>8</sup>. (Because the previous results involved comparison to Old World monkeys, it is possible that they reflect strong positive selection earlier in primate evolution; however, we suspect that they reflect the fact that relatively few genes were studied and that different genes were used to study polymorphism and divergence.)

Relaxed negative selection pressures thus primarily explain the excess amino acid divergence in hominid genes relative to murids. Moreover, because both  $\omega_{\text{human}}$  and  $\omega_{\text{chimpanzee}}$  are similarly elevated this explanation applies equally to both lineages.

We next sought to study variation in the evolutionary rate of genes within the hominid lineage by searching for unusually high or low levels of constraint for genes and sets of genes.

**Rapid evolution in individual genes.** We searched for individual genes that have accumulated amino acid substitutions faster than expected given the neutral substitution rate; we considered these genes as potentially being under strong positive selection. A total of 585 of the 13,454 human–chimpanzee orthologues (4.4%) have observed  $K_A/K_S > 1$  (see Supplementary Information 'Gene evolution'). However, given the low divergence, the  $K_A/K_S$  statistic has large variance. Simulations show that estimates of  $K_A/K_S > 1$  would be expected to occur simply by chance in at least 263 cases if purifying selection is allowed to act non-uniformly across genes (Supplementary Fig. S7).

Nonetheless, this set of 585 genes may be enriched for genes that are under positive selection. The most extreme outliers include glycoprotein C, which mediates one of the *Plasmodium falciparum* invasion pathways in human erythrocytes<sup>101</sup>; granulysin, which mediates antimicrobial activity against intracellular pathogens such as *Mycobacterium tuberculosis*<sup>102</sup>; as well as genes that have previously been shown to be undergoing adaptive evolution, such as the protamines and semenogelins involved in reproduction<sup>103</sup> and the Mas-related gene family involved in nociception<sup>104</sup>. With similar

**Table 3 | Comparison of  $K_A/K_S$  for divergence and human diversity**

Substitution type	$\Delta A$	$\Delta S$	$K_A/K_S$	Per cent excess*	Confidence interval†
Human–chimpanzee divergence	38,773	61,737	0.23	–	–
HapMap (European ancestry)‡					
Rare derived alleles (<15%)	1,614	1,540	0.39	67	[59, 75]
Common alleles	1,199	1,907	0.23	0	[–5, 6]
Frequent derived alleles (>85%)	209	356	0.22	–7	[–19, 7]
HapMap (African ancestry)‡					
Rare derived alleles (<5%)	849	842	0.36	61	[50, 72]
Common alleles	495	803	0.22	–2	[–10, 7]
Frequent derived alleles (>85%)	59	82	0.26	15	[–11, 48]
Affymetrix 120K (multi-ethnic)§					
Rare derived alleles (<15%)	74	82	0.33	44	[14, 80]
Common alleles	77	137	0.21	–11	[–28, 12]
Frequent derived alleles (>85%)	10	15	0.25	6	[–42, 95]

$\Delta A$ , Number of observed non-synonymous substitutions.  $\Delta S$ , Number of observed synonymous substitutions.

\* A negative value indicates excess of non-synonymous divergence over polymorphism.

† 95% confidence intervals assuming non-synonymous substitutions are Poisson distributed.

‡ Source: <http://www.hapmap.org> (Public Release no. 13).

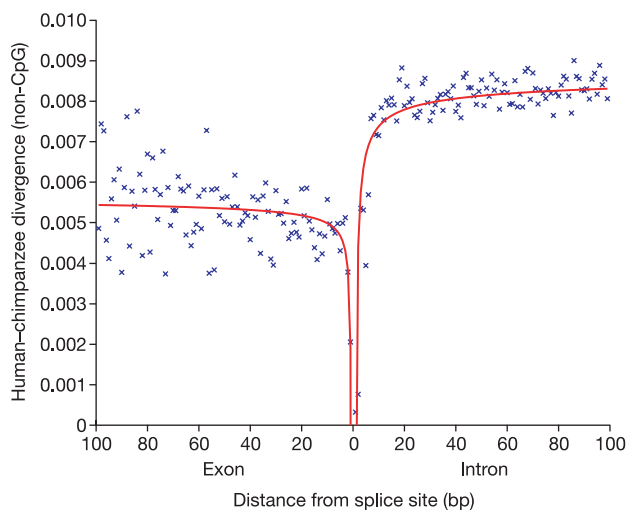
§ Source: <http://www.affymetrix.com>.

follow-up studies on candidates from this list, one may be able to draw conclusions about positive selection on other individual genes. In subsequent sections, we examine the rate of divergence for sets of related genes with the aim of detecting subtler signals of accelerated evolution.

**Variation in evolutionary rate across physically linked genes.** We explored how the rate of evolution varies regionally across the genome. Several studies of mammalian gene evolution have noted that the rate of amino acid substitution shows local clustering, with proteins encoded by nearby genes evolving at correlated rates<sup>16,105–107</sup>. *Variation across chromosomes.* On the basis of an analysis of ~100 genes<sup>108</sup>, it was recently reported that the normalized rate of protein evolution is greater on the nine chromosomes that underwent major structural rearrangement during human evolution (chromosomes 1, 2, 5, 9, 12, 15, 16, 17 and 18); it was suggested that such rearrangements led to reduced gene flow and accelerated adaptive evolution. A subsequent study of a collection of chimpanzee ESTs gave contradictory results<sup>109,110</sup>. With our larger data set, we re-examined this issue and found no evidence of accelerated evolution on chromosomes with major rearrangements, even if we considered each rearrangement separately (Supplementary Table S25).

Among all hominid chromosomes, the most extreme outlier is chromosome X with a mean  $K_A/K_I$  of 0.32. The higher mean seems to reflect a skewed distribution at both high and low values, with the median value (0.17) being more in line with other chromosomes (0.15). The excess of low values may reflect greater purifying selection at some genes, owing to the hemizyosity of chromosome X in males. The excess of high values may reflect increased adaptive selection also resulting from hemizyosity, if a considerable proportion of advantageous alleles are recessive<sup>111</sup>. Interestingly, the higher  $K_A/K_I$  value on the X chromosome versus autosomes is largely restricted to genes expressed in testis<sup>83</sup>.

*Variation in local gene clusters.* We next searched for genomic neighbourhoods with an unusually high density of rapidly evolving genes. Specifically, we calculated the median  $K_A/K_I$  for sliding windows of ten orthologues and identified extreme outliers ( $P < 0.001$  compared to random ordering of genes; see Supplementary Information 'Gene evolution'). A total of 16 such neighbourhoods were found, which greatly exceeds random expectation (Table 4). Repeating the analysis with larger windows (25, 50 and 100 orthologues) did not identify additional rapidly diverging regions.



**Figure 10 | Purifying selection on synonymous sites.** Mean divergence around exon boundaries at non-CpG, exonic, fourfold degenerate sites and intronic sites, relative to the closest mRNA splice junction. The divergence rate at exonic, fourfold degenerate sites is significantly lower than at nearby intronic sites (Mann–Whitney  $U$ -test;  $P < 10^{-27}$ ), suggesting that purifying selection limits the rate of synonymous codon substitutions.

In nearly all cases, the regions contain local clusters of phylogenetically and functionally related genes. The rapid diversification of gene families, postulated by ref. 112, can thus be readily discerned even at the relatively close distance of human–chimpanzee divergence. Most of the clusters are associated with functional categories such as host defence and chemosensation (see below). Examples include the epidermal differentiation complex encoding proteins that help form the cornified layer of the skin barrier (Supplementary Fig. S8), the WAP-domain cluster encoding secreted protease inhibitors with antibacterial activity, and the Siglec cluster encoding *CD33*-related genes. Rapid evolution in these clusters does not seem to be unique to either human or chimpanzee<sup>113,114</sup>.

**Variation in evolutionary rate across functionally related genes.** We next studied variation in the evolutionary rate of functional categories of genes, based on the Gene Ontology (GO) classification<sup>115</sup>.

*Rapidly and slowly evolving categories within the hominid lineage.* We started by searching for sets of functionally related genes with exceptionally high or low constraint in humans and chimpanzees. For each of the 809 categories with at least 20 genes,  $K_A/K_S$  was calculated by concatenating the gene sequences. The category-specific ratios were compared to the average across all orthologues to identify extreme outliers using a metric based on the binomial test (Supplementary Information 'Gene evolution' and Supplementary Tables S26–S29). The numbers of observed outliers below a specific threshold (test statistic  $< 0.001$ ) were then compared to the expected distribution of outliers given randomly permuted annotations.

A total of 98 categories showed elevated  $K_A/K_S$  ratios at the specified threshold (Table 5). Only 30 would be expected by chance, indicating that most (but not all) of these categories undergo significantly accelerated evolution relative to the genome-wide average ( $P < 10^{-4}$ ). The rapidly evolving categories within the hominid lineage are primarily related to immunity and host defence, reproduction, and olfaction, which are the same categories known to be undergoing rapid evolution within the broader mammalian lineage, as well as more distantly related species<sup>15,16,116</sup>. Hominids thus seem to be typical of mammals in this respect (but see below).

A total of 251 categories showed significantly low  $K_A/K_S$  ratios (compared with ~32 expected by chance;  $P < 10^{-4}$ ). These include a wide range of processes including intracellular signalling, metabolism, neurogenesis and synaptic transmission, which are evidently under stronger-than-average purifying selection. More generally, genes expressed in the brain show significantly stronger average constraint than genes expressed in other tissues<sup>83</sup>.

*Differences between hominid and murid lineages.* Having found gene categories that show substantial variation in absolute evolutionary rate within hominids, we next examined variation in relative rates

**Table 4 | Rapidly diverging gene clusters in human and chimpanzee**

Location (human)	Cluster	Median $K_A/K_I$ *
1q21	Epidermal differentiation complex	1.46
6p22	Olfactory receptors and HLA-A	0.96
20p11	Cystatins	0.94
19q13	Pregnancy-specific glycoproteins	0.94
17q21	Hair keratins and keratin-associated proteins	0.93
19q13	CD33-related Siglecs	0.90
20q13	WAP domain protease inhibitors	0.90
22q11	Immunoglobulin- $\lambda$ /breakpoint critical region	0.85
12p13	Taste receptors, type 2	0.81
17q12	Chemokine (C-C motif) ligands	0.81
19q13	Leukocyte-associated immunoglobulin-like receptors	0.80
5q31	Protocadherin- $\beta$	0.77
1q32	Complement component 4-binding proteins	0.76
21q22	Keratin-associated proteins and uncharacterized ORFs	0.76
1q23	CD1 antigens	0.72
4q13	Chemokine (C-X-C motif) ligands	0.70

\*Maximum median  $K_A/K_I$  if the cluster stretched over more than one window of ten genes.

**Table 5 | GO categories with the highest divergence rates in hominids**

GO categories within 'biological process'	Number of orthologues	Amino acid divergence	$K_A/K_S$
GO:0007606 sensory perception of chemical stimulus	59	0.018	0.590
GO:0007608 perception of smell	41	0.018	0.521
GO:0006805 xenobiotic metabolism	40	0.013	0.432
GO:0006956 complement activation	22	0.013	0.428
GO:0042035 regulation of cytokine biosynthesis	20	0.011	0.402
GO:0007565 pregnancy	34	0.014	0.384
GO:0007338 fertilization	24	0.010	0.371
GO:0008632 apoptotic programme	36	0.010	0.358
GO:0007283 spermatogenesis	80	0.008	0.354
GO:0000075 cell cycle checkpoint	27	0.006	0.354

Listed are the ten categories in the taxonomy biological process with the highest  $K_A/K_S$  ratios, which are not significant solely due to significant subcategories.

between murids and hominids. The  $K_A/K_S$  of each of the GO categories are highly correlated between the hominid and murid orthologue pairs, suggesting that the selective pressures acting on particular functional categories have been largely proportional in recent hominid and recent murid evolution (Fig. 11). However, there are several categories with significantly accelerated non-synonymous divergence on each of the lineages, which might represent functions that have undergone lineage-specific positive selection or a lineage-specific relaxation of constraint (Supplementary Information 'Gene evolution' and Supplementary Tables S30–S39).

A total of 59 categories (compared with 11 expected at random,  $P < 0.0003$ ) show evidence of accelerated non-synonymous divergence in the murid lineage. These are dominated by functions and processes related to host defence, such as immune response and lymphocyte activation. Examples include genes encoding interleukins and various T-cell surface antigens (*Cd4*, *Cd8*, *Cd80*). Combined with the recent observation that genes involved in host defence have undergone gene family expansion in murids<sup>16,17</sup>, this suggests that the immune system has undergone extensive lineage-specific innovation in murids. Additional categories that also show relative acceleration in murids include chromatin-associated proteins and proteins involved in DNA repair. These categories may have similarly undergone stronger adaptive evolution in murids or, alternatively, they may contain fewer sites for mutations with slightly deleterious effects (with the result that the  $K_A/K_S$  ratios are less affected by the differences in population size<sup>96,117</sup>).

Another 58 categories (versus 14 expected at random,  $P < 0.0005$ ) show evidence of accelerated evolution in hominids, with the set dominated by genes encoding proteins involved in transport (for example, ion transport), synaptic transmission, spermatogenesis and perception of sound (Table 6). Notably, some outliers include genes with brain-related functions, compatible with a recent finding<sup>118</sup>. Potential positive selection on spermatogenesis genes in the hominids was also recently noted<sup>119</sup>. However, as above, it is possible that these categories could have more sites for slightly deleterious mutations and thus be more affected by population size differences. Sequence information from more species and from individuals

within species will be necessary to distinguish between the possible explanations.

*Differences between the human and chimpanzee lineage.* One of the most interesting questions is perhaps whether certain categories have undergone accelerated evolution in humans relative to chimpanzees, because such genes might underlie unique aspects of human evolution.

As was done for hominids and murids above, we compared non-synonymous divergence for each category to search for relative acceleration in either lineage (Fig. 12). Seven categories show signs of accelerated evolution on the human lineage relative to chimpanzee, but this is only slightly more than the four expected at random ( $P < 0.22$ ). Intriguingly, the single strongest outlier is 'transcription factor activity', with the 348 human genes studied having accumulated 47% more amino acid changes than their chimpanzee orthologues. Genes with accelerated divergence in human include homeotic, forkhead and other transcription factors that have key roles in early development. However, given the small number of changes involved, additional data will be required to confirm this trend. There was no excess of accelerated categories on the chimpanzee lineage.

We also compared human genes with and without disease associations, including mental retardation, for differences in mutation rate when compared to chimpanzee. Briefly, no significant differences were observed in either the background mutation rate or in the ratio of human-specific changes to chimpanzee-specific amino acid changes (see Supplementary Information 'Gene evolution' and Supplementary Tables S40 and S41).

We thus find minimal evidence of acceleration unique to either the human or chimpanzee lineage across broad functional categories. This is not simply due to general lack of power resulting from the small number of changes since the divergence of human and chimpanzee, because one can detect acceleration of categories in either hominid relative to either murid. For example, 29 accelerated categories versus 9 expected at random ( $P < 0.02$ ) can be detected on the human lineage, and 40 categories versus 11 expected at random ( $P < 0.007$ ) on the chimpanzee lineage, relative to mouse. But the

**Table 6 | GO categories with accelerated divergence rates in hominids relative to murids**

GO categories within 'biological process'	Number of orthologues	Amino acid divergence in hominids	Amino acid divergence in murids	$K_A/K_S$ in hominids	$K_A/K_S$ in murids
GO:0007283 spermatogenesis	43	0.0075	0.054	0.323	0.188
GO:0006869 lipid transport	22	0.0081	0.051	0.306	0.120
GO:0006865 amino acid transport	24	0.0058	0.033	0.218	0.084
GO:0015698 inorganic anion transport	29	0.0061	0.027	0.195	0.072
GO:0006486 protein amino acid glycosylation	50	0.0056	0.040	0.166	0.100
GO:0019932 second-messenger-mediated signalling	58	0.0049	0.036	0.159	0.083
GO:0007605 perception of sound	28	0.0052	0.033	0.158	0.085
GO:0016051 carbohydrate biosynthesis	27	0.0047	0.028	0.147	0.067
GO:0007268 synaptic transmission	93	0.0040	0.025	0.126	0.069
GO:0006813 potassium ion transport	65	0.0035	0.022	0.113	0.056

Listed are the ten categories in the taxonomy biological process with the strongest evidence for accelerated evolution in hominids relative to murids, which are not significant solely due to significant subcategories.

outliers are largely the same for both human and chimpanzee, indicating that the fraction of amino acid mutations that have contributed to human- and chimpanzee-specific patterns of evolution must be small relative to the fraction that have contributed to a common hominid and, to a large extent, mammalian pattern of evolution.

It was recently reported<sup>10</sup> that several functional categories are enriched for genes with evidence of positive selection in the human lineage or the chimpanzee lineage, and that these categories are largely different between the two lineages. These results and ours differ in ways that will require further investigation. With the potential exception of some developmental regulators, the categories that ref. 10 reported as showing the strongest enrichment of positive selection in one lineage (including cell adhesion, ion transport and perception of sound) are among those that we show as having accelerated divergence in both human and chimpanzee. This suggests that positive selection and relaxation of constraints may be correlated, or alternatively, that the results of ref. 10 may be enriched for false positives in categories that have experienced particularly strong relaxation of constraints in the hominids. Data from additional primates, as well as advances in analytical methods, will be necessary to distinguish between these alternatives. At present, strong evidence of positive selection unique to the human lineage is thus limited to a handful of genes<sup>120</sup>.

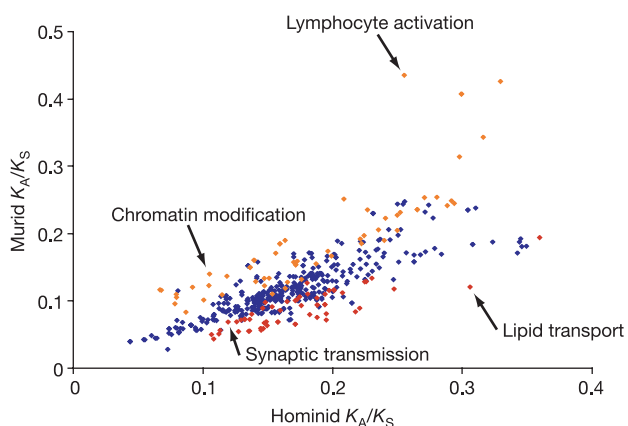
Our analysis above largely omitted genes belonging to large gene families, because gene family expansion makes it difficult to define 1:1:1 orthologues across hominids and murids. One of the largest such families, the olfactory receptors, is known to be undergoing rapid divergence in primates. Directed study of these genes in the draft assembly has suggested that more than 100 functional human olfactory receptors are likely to be under no evolutionary constraint<sup>121</sup>. Our analysis also omitted the majority of very recently duplicated genes owing to their lower coverage in the current chimpanzee assembly. However, recent human-specific duplications can be readily identified from the finished human genome sequence, and have previously been shown to be highly enriched for the same categories found to have high absolute rates of evolution in 1:1 orthologues here; that is, olfaction, immunity and reproduction<sup>23</sup>.

**Gene disruptions in human and chimpanzee.** Whereas most genes have undergone only subtle substitutions in their amino acid sequence, a few dozen have suffered more marked changes. We found a total of 53 known or predicted human genes that are either deleted entirely (36) or partially (17) in chimpanzee (Supplementary

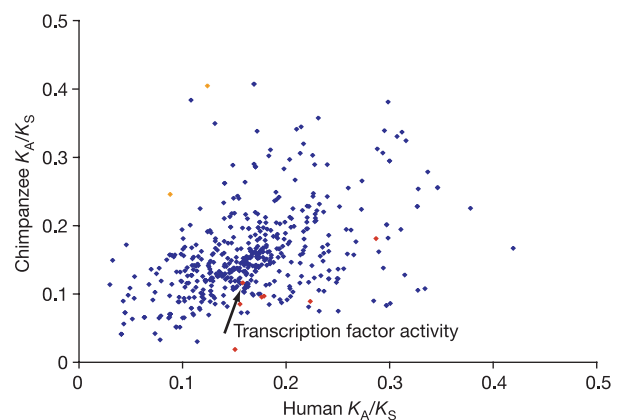
Table S42). We have so far tested and confirmed 15 of these cases by polymerase chain reaction (PCR) or Southern blotting. An additional eight genes have sustained large deletions (>15 kb) entirely within an intron. Some genes may have been missed in this count owing to limitations of the draft genome sequence. In addition, some genes may have suffered chain termination mutations or altered reading frames in chimpanzee, but accurate identification of these will require higher-quality sequence. The sensitivity of the reciprocal analysis of genes disrupted in human is currently limited by the small number of independently predicted gene models for the chimpanzee. Some of the gene disruptions may be related to interesting biological differences between the species, as discussed below. **Genetic basis for human- and chimpanzee-specific biology.** Given the substantial number of neutral mutations, only a small subset of the observed gene differences is likely to be responsible for the key phenotypic changes in morphology, physiology and behavioural complexity between humans and chimpanzees. Determining which differences are in this evolutionarily important subset and inferring their functional consequences will require additional types of evidence, including information from clinical observations and model systems<sup>122</sup>. We describe some novel examples of genetic changes for which plausible functional or physiological consequences can be suggested.

**Apoptosis.** Mouse and human are known to differ with respect to an important mediator of apoptosis, caspase-12 (refs 123–125). The protein triggers apoptosis in response to perturbed calcium homeostasis in mice, but humans seem to lack this activity owing to several mutations in the orthologous gene that together affect the protein produced by all known splice forms; the mutations include a premature stop codon and a disruption of the SHG box required for enzymatic activity of caspases. By contrast, the chimpanzee gene encodes an intact open reading frame and SHG box, indicating that the functional loss occurred in the human lineage. Intriguingly, loss-of-function mutations in mice confer increased resistance to amyloid-induced neuronal apoptosis without causing obvious developmental or behavioural defects<sup>126</sup>. The loss of function in humans may contribute to the human-specific pathology of Alzheimer's disease, which involves amyloid-induced neurotoxicity and deranged calcium homeostasis.

**Inflammatory response.** Human and chimpanzee show a notable difference with respect to important mediators of immune and inflammatory responses. Three genes (*IL1F7*, *IL1F8* and *ICEBERG*)



**Figure 11 | Hominid and murid  $K_A/K_S$  ( $\omega$ ) in GO categories with more than 20 analysed genes.** GO categories with putatively accelerated (test statistic <0.001; see Methods) non-synonymous divergence on the hominid lineages (red) and on the murid lineages (orange) are highlighted. Owing to the hierarchical nature of GO, the categories do not all represent independent data points. A non-redundant list of significant categories is provided in Table 8 and a complete list in Supplementary Table S30.



**Figure 12 | Human and chimpanzee  $K_A/K_S$  ( $\omega$ ) in GO categories with more than 20 analysed genes.** GO categories with putatively accelerated (test statistic <0.001; see Methods) non-synonymous divergence on the human lineage (red) and on the chimpanzee lineage (orange) are highlighted. The variance of these estimates is larger than that seen in the hominid–murid comparison owing to the small number of lineage-specific substitutions. Owing to the hierarchical nature of the GO ontology, the categories do not all represent independent data points. A complete list of categories is provided in Supplementary Table S30.



that act in a common pathway involving the caspase-1 gene all appear to be deleted in chimpanzee. *ICEBERG* is thought to repress caspase-1-mediated generation of pro-inflammatory *IL1* cytokines, and its absence in chimpanzee may point to species-specific modulation of the interferon- $\gamma$ - and lipopolysaccharide-induced inflammatory response<sup>127</sup>.

**Parasite resistance.** Similarly, we found that two members of the primate-specific *APOL* gene cluster (*APOL1* and *APOL4*) have been deleted from the chimpanzee genome. The *APOL1* protein is associated with the high-density lipoprotein fraction in serum and has recently been proposed to be the lytic factor responsible for resistance to certain subspecies of *Trypanosoma brucei*, the parasite that causes human sleeping sickness and the veterinary disease nagana<sup>128</sup>. The loss of the *APOL1* gene in chimpanzees could thus explain the observation that human, gorilla and baboon possess the trypanosome lytic factor, whereas the chimpanzee does not<sup>129</sup>.

**Sialic acid biology related proteins.** Sialic acids are cell-surface sugars that mediate many biological functions<sup>130</sup>. Of 54 genes involved in sialic acid biology, 47 were suitable for analysis. We confirmed and extended findings on several that have undergone human-specific changes, including disruptions, deletions and domain-specific functional changes<sup>113,131,132</sup>. Human- and chimpanzee-specific changes were also found in otherwise evolutionarily conserved sialyl motifs in four sialyl transferases (*ST6GAL1*, *ST6GALNAC3*, *ST6GALNAC4* and *ST8SIA2*), suggesting changes in donor and/or acceptor binding<sup>130</sup>. Lineage-specific changes were found in a complement factor H (*HFI1*) sialic acid binding domain associated with human disease<sup>133</sup>. Human *SIGLEC11* has undergone gene conversion with a nearby pseudogene, correlating with acquisition of human-specific brain expression and altered binding properties<sup>134</sup>.

**Human disease alleles.** We next sought to identify putative functional differences between the species by searching for instances in which a human disease-causing allele appears to be the wild-type allele in the chimpanzee. Starting from 12,164 catalogued disease variants in 1,384 human genes, we identified 16 cases in which the altered sequence in a disease allele matched the chimpanzee sequence, and had plausible support in the literature (Table 7; see also Supplementary Table S43). Upon re-sequencing in seven chimpanzees, 15 cases were confirmed homozygous in all individuals, whereas one (*PONI* I102V) appears to be a shared polymorphism (Supplementary Table S44).

Six cases represent *de novo* human mutations associated with simple mendelian disorders. Similar cases have also been found in comparisons of more distantly related mammals<sup>135</sup>, as well as

between insects<sup>136</sup>, and have been interpreted as a consequence of a relatively high rate of compensatory mutations. If compensatory mutations are more likely to be fixed by positive selection than by neutral drift<sup>136</sup>, then the variants identified here might point towards adaptive differences between humans and chimpanzees. For example, the ancestral Thr 29 allele of cationic trypsinogen (*PRSSI*) causes autosomal dominant pancreatitis in humans<sup>137</sup>, suggesting that the human-specific Asn 29 allele may represent a digestion-related molecular adaptation<sup>138</sup>.

The remaining ten cases represent common human polymorphisms that have been reported to be associated with complex traits, including coronary artery disease and diabetes mellitus. In all of these cases we confirmed that the disease-associated allele in humans is indeed the ancestral allele by showing that it is carried not only by chimpanzee but also by outgroups such as the macaque. These ancestral alleles may thus have become human-specific risk factors due to changes in human physiology or environment, and the polymorphisms may represent ongoing adaptations. For example, *PPARG* Pro 12 is the wild-type allele in chimpanzee but has been clearly associated with increased risk of type 2 diabetes in human<sup>139</sup>. It is tempting to speculate that this allele may represent an ancestral 'thrifty' genotype<sup>140</sup>.

The current results must be interpreted with caution, because few complex disease associations have been firmly established. The fact that the human disease allele is the wild-type allele in chimpanzee may actually indicate that some of the putative associations are spurious and not causal. However, this approach can be expected to become increasingly fruitful as the quality and completeness of the disease mutation databases improve.

### Human population genetics

The chimpanzee has a special role in informing studies of human population genetics, a field that is undergoing rapid expansion and acquiring new relevance to human medical genetics<sup>141</sup>. The chimpanzee sequence allows recognition of those human alleles that represent the ancestral state and the derived state. It also allows estimates of local mutation rates, which serve as an important baseline in searching for signs of natural selection.

**Ancestral and derived alleles.** Of ~7.2 million SNPs mapped to the human genome in the current public database, we could assign the alleles as ancestral or derived in 80% of the cases according to which allele agrees with the chimpanzee genome sequence<sup>142</sup> (see Supplementary Information 'Human population genetics'). For the remaining cases, no assignment could be made because of the following: the orthologous chimpanzee base differed from both human alleles (1.2%); was polymorphic in the chimpanzee sequences obtained (0.4%); or could not be reliably identified with the current draft sequence of the chimpanzee (18.8%), with many of these occurring in repeated or segmentally duplicated sequence. The first two cases arise presumably because a second mutation occurred in the chimpanzee lineage. It should be possible to resolve most of these cases by examining a close outgroup such as gorilla or orang-utan.

Mutations in the chimpanzee may also lead to the erroneous assignment of human alleles as derived alleles. This error rate can be estimated as the probability of a second mutation resulting in the chimpanzee sequence matching the derived allele (see Supplementary Information 'Human population genetics'). The estimated error rate for typical SNPs is 0.5%, owing to the low nucleotide substitution rate. The exceptions are those SNPs for which the human alleles are CpG and TpG and the chimpanzee sequence is TpG. For these, a non-negligible fraction may have arisen by two independent deamination events within an ancestral CpG dinucleotide, which are well-known mutational hotspots<sup>51</sup> (also see above). Human SNPs in a CpG context for which the orthologous chimpanzee sequence is TpG account for 12% of the total, and have an estimated error rate of 9.8%. Across all SNPs, the average error rate,  $\epsilon$ , is thus estimated to be ~1.6%.

We compared the distribution of allele frequencies for ancestral

**Table 7 | Candidate human disease variants found in chimpanzee**

Gene	Variant*	Disease association	Ancestral†	Frequency‡
<i>AIRE</i>	P252L <sup>159</sup>	Autoimmune syndrome	Unresolved	0
<i>MKKS</i>	R518H <sup>160</sup>	Bardet-Biedl syndrome	Wild type	0
<i>MLH1</i>	A441T <sup>161</sup>	Colorectal cancer	Wild type	0
<i>MYOC</i>	Q48H <sup>162</sup>	Glaucoma	Wild type	0
<i>OTC</i>	T125M <sup>163</sup>	Hyperammonaemia	Wild type	0
<i>PRSS1</i>	N29T <sup>137</sup>	Pancreatitis	Disease	0
<i>ABCA1</i>	I883M <sup>164</sup>	Coronary artery disease	Unresolved	0.136
<i>APOE</i>	C130R <sup>165</sup>	Coronary artery disease and Alzheimer's disease	Disease	0.15
<i>DIO2</i>	T92A <sup>166</sup>	Insulin resistance	Disease	0.35
<i>ENPP1</i>	K121Q <sup>167</sup>	Insulin resistance	Disease	0.17
<i>GSTP1</i>	I105V <sup>168</sup>	Oral cancer	Disease	0.348
<i>PON1S</i>	I102V <sup>169</sup>	Prostate cancer	Wild type	0.016
<i>PONI</i>	Q192R <sup>170</sup>	Coronary artery disease	Disease	0.3
<i>PPARG</i>	A12P <sup>139</sup>	Type 2 diabetes	Disease	0.85
<i>SLC2A2</i>	T110I <sup>171</sup>	Type 2 diabetes	Disease	0.12
<i>UCP1</i>	A64T <sup>172</sup>	Waist-to-hip ratio	Disease	0.12

\* This takes the following format: benign variant, codon number, disease/chimpanzee variant.

† Ancestral variant as inferred from closest available primate outgroups (Supplementary Information).

‡ Frequency of the disease allele in human study population.

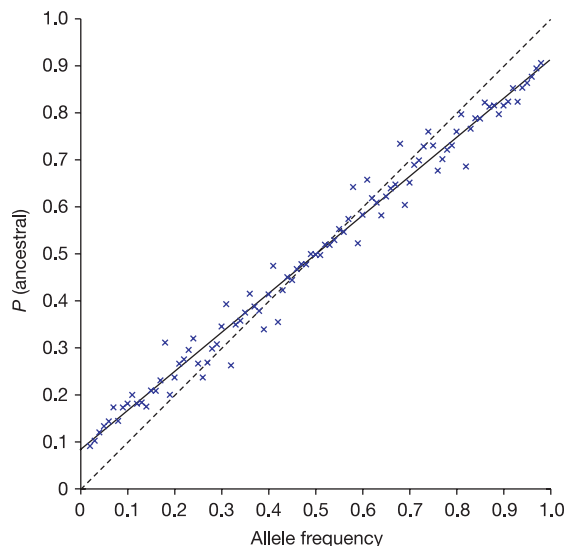
§ Polymorphic in chimpanzee.

and derived alleles using a database of allele frequencies for  $\sim 120,000$  SNPs (see Supplementary Information 'Human population genetics'). As expected, ancestral alleles tend to have much higher frequencies than derived alleles (Supplementary Fig. S9). Nonetheless, a significant proportion of derived alleles have high frequencies: 9.1% of derived alleles have frequency  $\geq 80\%$ .

An elegant result in population genetics states that, for a randomly interbreeding population of constant size, the probability that an allele is ancestral is equal to its frequency<sup>143</sup>. We explored the extent to which this simple theoretical expectation fits the human population. We tabulated the proportion  $p_a(x)$  of ancestral alleles for various frequencies of  $x$  and compared this with the prediction  $p_a(x) = x$  (Fig. 13).

The data lie near the predicted line, but the observed slope (0.83) is substantially less than 1. One explanation for this deviation is that some ancestral alleles are incorrectly assigned (an error rate of  $\varepsilon$  would artificially decrease the slope by a factor of  $1-2\varepsilon$ ). However, with  $\varepsilon$  estimated to be only 1.6%, errors can only explain a small part of the deviation. The most likely explanation is the presence of bottlenecks during human history, which tend to flatten the distribution of allele frequencies. Theoretical calculations indicate that a recent bottleneck would decrease the slope by a factor of  $(1-b)$ , where  $b$  is the inbreeding coefficient induced by the bottleneck (see Supplementary Information 'Human population genetics' and Supplementary Fig. S10). This suggests that measurements of the slope in different human groups may shed light on population-specific bottlenecks. Consistent with this, preliminary analyses of allele frequencies in several regions for SNPs obtained by systematic uniform sampling indicate that the slope is significantly lower than 1 in European and Asian samples and close to 1 in an African sample (see Supplementary Information 'Human population genetics' and Supplementary Fig. S11).

**Signatures of strong selective sweeps in recent human history.** The pattern of human genetic variation holds substantial information about selection events that have shaped our species. Strong positive selection creates the distinctive signature of a 'selective sweep', whereby a rare allele rapidly rises to fixation and carries the haplotype on which it occurs to high frequency (the 'hitchhiking' effect). The surrounding region should show two distinctive signatures: a significant reduction of overall diversity, and an excess of derived alleles with high frequency in the population owing to hitchhiking of



**Figure 13 | The observed fraction of ancestral alleles in 1% bins of observed frequency.** The solid line shows the regression ( $b = 0.83$ ). The dotted line shows the theoretical relationship  $p_a(x) = x$ . Note that because each variant yields a derived and an ancestral allele, the data are necessarily symmetrical about 0.5.

derived alleles on the selected haplotype (see Supplementary Information 'Human population genetics'). The pattern might be detectable for up to 250,000 years after a selective sweep has ended<sup>144</sup>. Notably, the chimpanzee genome provides crucial baseline information required for accurate assessment of both signatures.

The size of the interval affected by a selective sweep is expected to scale roughly with  $s$ , the selective advantage due to the mutation. Simulations can be used to study the distribution of the interval size (see Supplementary Information 'Human population genetics'). With  $s = 1\%$ , the interval over which heterozygosity falls by 50% has a modal size of 600 kb and a probability of greater than 10% of exceeding 1 Mb.

We undertook an initial scan for large regions ( $>1$  Mb) with the two signatures suggestive of strong selective sweeps in recent human history. We began by identifying regions in which the observed human diversity rate was much lower than the expectation based on the observed divergence rate with chimpanzee. The human diversity rate was measured as the number of occurrences from a database of 1.92 million SNPs identified by shotgun sequencing in a panel of African-American individuals (see Supplementary Information 'Genome sequencing and assembly'). The comparison with the chimpanzee eliminates regions in which low diversity simply reflects a low mutation rate in the region. Regions were identified based on a simple statistical procedure (see Supplementary Information 'Human population genetics'). Six genomic regions stand out as clear outliers that show significantly reduced diversity relative to divergence (Table 8; see also Supplementary Fig. S12).

We next tested whether these six regions show a high proportion of SNPs with high-frequency derived alleles (defined here as alleles with frequency  $\geq 80\%$ ). Within each region, we focused on the 1-Mb interval with the greatest discrepancy between diversity and divergence and compared it to 1-Mb regions throughout the genome. For the database of 120,000 SNPs with allele frequencies discussed above, the typical 1-Mb region in the human genome contains  $\sim 40$  SNPs, and the proportion  $p_h$  of SNPs with high-frequency derived alleles is  $\sim 9.1\%$ . All six regions identified by our scan for reduced diversity have a higher than average fraction of high-frequency derived alleles; all six fall within the top 10% genome-wide and three fall within the top 1%. Although this is not definitive evidence for any particular region, the joint probability of all six regions randomly scoring in the top 10% is  $10^{-6}$ . The results indicate that the six regions are candidates for strong selective sweeps during the past 250,000 years<sup>144</sup>. The regions differ notably with respect to gene content, ranging from one containing 57 annotated genes (chromosome 22) to another with no annotated genes whatsoever (chromosome 4). We have no evidence to implicate any individual functional element as a target of recent selection at this point, but the regions contain a number of interesting candidates for follow-up studies. Intriguingly, the chromosome 4 gene desert, which flanks a protocadherin gene and is conserved across vertebrates<sup>15</sup>, has been implicated in two independent studies as being associated with obesity<sup>145,146</sup>.

In addition to the six regions, one further genomic region deserves mention: an interval of 7.6 Mb on chromosome 7q (see Supplementary Information 'Human population genetics'). The interval contains several regions with high scores in the diversity-divergence analysis (including the seventh highest score overall) as well as in the proportion of high-frequency derived alleles. The region contains the *FOXP2* and *CFTR* genes. The former has been the subject of much interest as a possible target for selection during human evolution<sup>147</sup> and the latter as a target of selection in European populations<sup>148</sup>.

Convincing proof of past selection will require careful analysis of the precise pattern of genetic variation in the region and the identification of a likely target of selection. Nonetheless, our findings suggest that the approach outlined here may help to unlock some of the secrets of recent human evolution through a combination of within-species and cross-species comparison.

**Table 8 | Human regions with strongest signal of selection based on diversity relative to divergence**

Chromosome	Start (Mb)	End (Mb)	Regression log-score	Skew P-value	Genes
1	48.58	52.58	103.3	0.071	Fourteen known genes from <i>ELAVL4</i> to <i>GPX7</i>
2	144.35	148.47	84.8	0.074	<i>ARHGAP15</i> (partial), <i>GTDC1</i> and <i>ZFH1B</i>
22	36.15	40.22	81.8	0.00022	Fifty-seven known genes from <i>CARD10</i> to <i>PMM1</i>
12	84.69	89.01	80.9	0.031	Ten known genes from <i>PAMC1</i> to <i>ATP2B1</i>
8	34.91	37.54	76.9	0.00032	<i>UNC5D</i> and <i>FKSG2</i>
4	32.42	35.62	55.9	0.00067	No known genes or Ensembl predictions

## Discussion

Our knowledge of the human genome is greatly advanced by the availability of a second hominid genome. Some questions can be directly answered by comparing the human and chimpanzee sequences, including estimates of regional mutation rates and average selective constraints on gene classes. Other questions can be addressed in conjunction with other large data sets, such as issues in human population genetics for which the chimpanzee genome provides crucial controls. For still other questions, the chimpanzee genome simply provides a starting point for further investigation.

The hardest such question is: what makes us human? The challenge lies in the fact that most evolutionary change is due to neutral drift. Adaptive changes comprise only a small minority of the total genetic variation between two species. As a result, the extent of phenotypic variation between organisms is not strictly related to the degree of sequence variation. For example, gross phenotypic variation between human and chimpanzee is much greater than between the mouse species *Mus musculus* and *Mus spretus*, although the sequence difference in the two cases is similar. On the other hand, dogs show considerable phenotypic variation despite having little overall sequence variation (~0.15%). Genomic comparison markedly narrows the search for the functionally important differences between species, but specific biological insights will be needed to sift the still-large list of candidates to separate adaptive changes from neutral background.

Our comparative analysis suggests that the patterns of molecular evolution in the hominids are typical of a broader class of mammals in many ways, but distinctive in certain respects. As with the murids, the most rapidly evolving gene families are those involved in reproduction and host defence. In contrast to the murids, however, hominids appear to experience substantially weaker negative selection; this probably reflects their smaller population size. Consequently, hominids accumulate deleterious mutations that would be eliminated by purifying selection in murids. This may be both an advantage and a disadvantage. Although decreased purifying selection may tend to erode overall fitness, it may also allow hominids to 'explore' larger regions of the fitness landscape and thereby achieve evolutionary adaptations that can only be reached by passing through intermediate states of inferior fitness<sup>149,150</sup>.

Although the analyses presented here focus on protein-coding sequences, the chimpanzee genome sequence also allows systematic analysis of the recent evolution of gene regulatory elements for the first time. Initial analysis of both gene expression patterns and promoter regions suggest that their overall patterns of evolution closely mirror that of protein-coding regions. In an accompanying paper<sup>83</sup>, we show that the rates of change in gene expression among different tissues in human and chimpanzee correlate with the nucleotide divergence in the putative proximal promoters and even more interestingly with the average level of constraint on proteins in the same tissues. Another study<sup>151</sup> has similarly used the chimpanzee sequence described here to show that gene promoter regions are also evolving under markedly less constraint in hominids than in murids.

The draft chimpanzee sequence here is sufficient for initial analyses, but it is still imperfect and incomplete. Definitive studies of gene and genome evolution—including pseudogene formation, gene family expansion and segmental duplication—will require high-

quality finished sequence. In this regard, we note that efforts are already underway to construct a BAC-based physical map and to increase the shotgun sequence coverage to approximately sixfold redundancy. The added coverage alone will not affect the analysis greatly, but plans are in place to produce finished sequence for difficult to sequence and important segments of the genome.

Our close biological relatedness to chimpanzees not only allows unique insights into human biology, it also creates ethical obligations. Although the genome sequence was acquired without harm to chimpanzees, the availability of the sequence may increase pressure to use chimpanzees in experimentation. We strongly oppose reducing the protection of chimpanzees and instead advocate the policy positions suggested by an accompanying paper<sup>152</sup>. Furthermore, the existence of chimpanzees and other great apes in their native habitats is increasingly threatened by human civilization. More effective policies are urgently needed to protect them in the wild. We hope that elaborating how few differences separate our species will broaden recognition of our duty to these extraordinary primates that stand as our siblings in the family of life.

## METHODS

**Sequencing and assembly.** Approximately 22.5 million sequence reads were derived from both ends of inserts (paired end reads) from 4-, 10-, 40- and 180-kb clones, all prepared from primary blood lymphocyte DNA. Genomic resources available from the source animal include a lymphoid cell line (S006006) and genomic DNA (NS06006) at Coriell Cell Repositories (<http://locus.umdj.edu/ccr/>), as well as a BAC library (CHORI-251)<sup>153</sup> (see also Supplementary Information 'Genome sequencing and assembly').

**Genome alignment.** BLASTZ<sup>154</sup> was used to align non-repetitive chimpanzee regions against repeat-masked human sequence. BLAT<sup>155</sup> was subsequently used to align the more repetitive regions. The combined alignments were chained<sup>156</sup> and only best reciprocal alignments were retained for further analysis.

**Insertions and deletions.** Small insertion/deletion (indel) events (<15 kb) were parsed directly from the BLASTZ genome alignment by counting the number and size of alignment gaps between bases within the same contig. Sites of large-scale indels (>15 kb) were detected from discordant placements of paired sequence reads against the human assembly. Size thresholds were obtained from both human fosmid alignments on human sequence ( $40 \pm 2.58$  kb) and chimpanzee plasmid alignments against human chromosome 21 ( $4.5 \pm 1.84$  kb). Indels were inferred by two or more pairs surpassing these thresholds by more than two standard deviations and the absence of sequence data within the discordancy.

**Gene annotation.** A total of 19,277 human RefSeq transcripts<sup>157</sup>, representing 16,045 distinct genes, were indirectly aligned to the chimpanzee sequence via the genome alignment. After removing low-quality sequences and likely alignment artefacts, an initial catalogue containing 13,454 distinct 1:1 human–chimpanzee orthologues was created for the analyses described here. A subset of 7,043 of these genes with unambiguous mouse and rat orthologues were realigned using Clustal W<sup>158</sup> for the lineage-specific analyses. Updated gene catalogues can be obtained from <http://www.ensembl.org>.

**Rates of divergence.** Nucleotide divergence rates were estimated using baseml with the REV model. Non-CpG rates were estimated from all sites that did not overlap a CG dinucleotide in either human or chimpanzee.  $K_A$  and  $K_S$  were estimated jointly for each orthologue using codeml with the F3x4 codon frequency model and no additional constraints, except for the comparison of divergent and polymorphic substitutions where  $K_A/K_S$  for both was estimated as  $(\Delta A/N_A)/(\Delta S/N_S)$ , with  $N_S/N_A$ , the ratio of synonymous to non-synonymous sites, estimated as 0.36 from the orthologue alignments. Unless otherwise specified,  $K_A/K_S$  for a set of genes was calculated by summing the number of substitutions and the number of sites to obtain  $K_A$  and  $K_S$  for the concatenated set before taking



the ratio. Hominid and murid pairwise rates were estimated independently from codons aligned across all four species. Human and chimpanzee lineage-specific  $K_A$  and  $K_S$  were estimated on an unrooted tree with both mouse and rat included. Lineage-specific rates were also estimated by parsimony, with essentially identical results (see Supplementary Information).  $K_1$  was estimated from all interspersed repeats within 250 kb of the mid-point of each gene.

**Accelerated evolution in GO categories.** The binomial probability of observing  $X$  or more non-synonymous substitutions, given a total of  $X + Y$  substitutions and the expected proportion  $x$  from all orthologues, was calculated by summing substitutions across the orthologues in each GO category. For the absolute rate test,  $Y$  = the number of synonymous substitutions in orthologues in the same category. For the relative rate tests,  $Y$  = the number of non-synonymous substitutions on the opposite lineage. Note that this binomial probability is simply a metric designed to identify potentially accelerated categories, it is not a  $P$ -value that can be used to reject directly the null hypothesis of no acceleration in that particular category. For each test, the observed number of categories with a binomial probability less than 0.001 was compared to the expected distribution of such outliers by repeating the procedure 10,000 times on randomly permuted GO annotations. The significance of the number of observed outliers  $n$  was estimated as the proportion of random trials yielding  $n$  or more outliers.

**Detection of selective sweeps.** The observed number of human SNPs,  $u_i$ , human bases,  $m_i$ , human–chimpanzee substitutions,  $v_i$ , and chimpanzee bases,  $n_i$ , within each set of non-overlapping 1-Mb windows along the human genome were used to generate two random numbers,  $x_i$  (adjusted human diversity) and  $y_i$  (adjusted human–chimpanzee divergence), from the two beta-distributions:

$$x_i \approx \text{Beta}(u_i + a, m_i - u_i + b)$$

$$y_i \approx \text{Beta}(v_i + c, n_i - v_i + d)$$

where  $a = 1$ ,  $b = 1,000$ ,  $c = 1$  and  $d = 100$ . These numbers were then fit to a linear regression:

$$x|y \approx N(\alpha_0 + \alpha_1 y, \beta^2)$$

A  $P$ -value for each window was calculated for each window based on  $(x_i, y_i)$  and the regression line. This was repeated 100 times and the average of the  $P$ -values taken as the  $P$ -value for diversity given divergence in each window. Overlapping windows with  $P < 0.1$  containing at least one window of  $P < 0.05$  were coalesced and scored as the sum of their  $-\log(p)$  scores.

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**Supplementary Information** is linked to the online version of the paper at [www.nature.com/nature](http://www.nature.com/nature).

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