

Great ape genetic diversity and population history

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Most great ape genetic variation remains uncharacterized^{1,2}; however, its study is critical for understanding population history³⁻⁶, recombination⁷, selection⁸ and susceptibility to disease^{9,10}. Here we sequence to high coverage a total of 79 wild- and captive-born individuals representing all six great ape species and seven subspecies and report 88.8 million single nucleotide polymorphisms. Our analysis provides support for genetically distinct populations within each species, signals of gene flow, and the split of common chimpanzees into two distinct groups: Nigeria-Cameroon/western and central/eastern populations. We find extensive inbreeding in almost all wild populations, with eastern gorillas being the most extreme. Inferred effective population sizes have varied radically over time in different lineages and this appears to have a profound effect on the genetic diversity at, or close to, genes in almost all species. We discover and assign 1,982 loss-of-function variants throughout the human and great ape lineages, determining that the rate of gene loss has not been different in the human branch compared to other internal branches in the great ape phylogeny. This comprehensive catalogue of great ape genome diversity provides a framework for understanding evolution and a resource for more effective management of wild and captive great ape populations.

We sequenced great ape genomes to a mean of 25-fold coverage per individual (Table 1, Supplementary Information and Supplementary Table 1) sampling natural diversity by selecting captive individuals of known wild-born origin as well as individuals from protected areas in Africa (Fig. 1a). We also included nine human genomes—three African and six non-African individuals¹¹. Variants were called using the software package GATK (ref. 12) (Methods), applying several quality

filters, including conservative allele balance filters, and requiring that genomes showed <2% contamination between samples (Methods and Supplementary Information). In order to assess the quality of single nucleotide variant (SNV) calls, we performed three sets of independent validation experiments with concordance rates ranging from 86% to 99% depending on allele frequency, the great ape population analysed and the species reference genome used (Supplementary Information and Supplementary Table 2). In total, we discovered 84.0 million fixed substitutions and 88.8 million segregating sites of high quality (Table 1 and Supplementary Table 3), providing the most comprehensive catalogue of great ape genetic diversity to date. From these variants we also constructed a list of potentially ancestry-informative markers (AIMs) for each of the surveyed populations, although a larger sampling of some subspecies is still required (Supplementary Information).

We initially explored the genetic relationships between individuals by constructing neighbour-joining phylogenetic trees from both autosomal and mitochondrial genomes (Supplementary Information). The autosomal tree identified separate monophyletic groupings for each species or subspecies designation (Supplementary Fig. 8.5.1) and supports a split of extant chimpanzees into two groups. Nigeria-Cameroon and western chimpanzees form a monophyletic clade (>97% of all autosomal trees); central and eastern chimpanzees form a second group (72% of all autosomal trees).

Genome-wide patterns of heterozygosity (Fig. 1b) reveal a threefold range in single nucleotide polymorphism (SNP) diversity. Non-African humans, eastern lowland gorillas, bonobos and western chimpanzees show the lowest genetic diversity ($\sim 0.8 \times 10^{-3}$ heterozygotes per base pair (bp)). In contrast, central chimpanzees, western lowland gorillas

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Table 1 | Genetic variation summary by species and subspecies

Genus	Scientific name species/subspecies	Common name	N	Mean coverage	Fixed sites to human reference	No. of SNVs*	Mean SNVs per individual*	No. of singletons†	Ancestry informative markers (AIMs)‡	N_e (10^{-3})§
<i>Homo</i>	<i>Homo sapiens</i>	Non-African	6	18.3	386,974	5,887,443	2,639,546	1,379,448	12,316	9.7–19.5
		African	3	20.9	632,253	6,309,453	3,203,178	2,448,454	12,316	13.9–27.9
		Humans	9	19.2	224,660	9,172,573	3,061,604	3,827,902	NA	13.1–16.2
<i>Pan</i>	<i>Pan troglodytes</i>	Nigeria–Cameroon	10	16.7	25,017,403	12,605,585	4,816,435	2,695,109	2,213	18.5–37.0
		Eastern	6	28.7	25,126,506	11,264,879	4,843,530	2,228,396	1,265	19.7–39.5
		Central	4	23.8	25,080,750	11,820,858	4,983,933	3,948,347	619	24.4–48.7
		Western	4	27.3	26,832,247	4,729,933	2,411,501	1,481,079	145,548	9.8–19.5
		Common Chimpanzees	24	22.5	24,087,088	27,153,659	5,693,903	10,352,931	149,645	30.9–61.8
<i>Pan paniscus</i>	Bonobos	13	27.5	27,068,299	8,950,002	2,738,755	3,159,889	NA	11.9–23.8	
<i>Gorilla</i>	<i>Gorilla beringei</i>	Eastern lowland	3	22.8	34,537,496	3,866,117	2,578,328	484,482	317,028	12.2–24.3
		Cross river	1	17.6	35,553,861	2,585,360	2,585,360	165,482	35,693	14.9–29.8
		Western lowland	23	17.8	31,602,620	17,314,403	6,410,662	2,797,388	19,902	26.8–53.5
	Gorillas	27	18.3	31,376,203	19,177,989	6,492,831	3,447,352	372,623	28.4–56.9	
<i>Pongo</i>	<i>Pongo abellii</i>	Sumatran	5	28.7	62,880,923	14,543,573	7,263,256	5,681,303	1,132,808	27.5–55.0
		Bornean	5	25.8	64,249,235	10,321,213	5,763,354	3,555,596	1,132,808	19.5–39.0
		Orangutans	10	27.3	60,661,869	24,309,920	9,338,148	6,409,648	NA	42.3–84.6
		All	83	23.0	83,954,672	88,764,143	NA	NA	NA	NA

*Polymorphic variants found in each species/subspecies after subtracting fixed sites.

†Singletons and doubletons calculated combining all the samples within the species.

‡Variants only found in a single group within each species.

§Calculated from Θ_w . $\mu = 1 \times 10^{-9}$ to 0.5×10^{-9} mut bp⁻¹ yr⁻¹ and $g = 25$ for *Homo* and *Pan*, 19 for *Gorilla* and 26 for *Pongo*.

||Hybrid sample Donald and 4 related gorillas were excluded.

The combined data for groups is shown in bold.

NA, not applicable.

and both orangutan species show the greatest genetic diversity (1.6×10^{-3} – 2.4×10^{-3} heterozygotes per bp). These differences are also reflected by measures of inbreeding from runs of homozygosity¹³ (Fig. 1c and Supplementary Information). Bonobos and western lowland gorillas, for example, have similar distributions of tracts of homozygosity as human populations that have experienced strong genetic bottlenecks (Karitiana and Papuan). Eastern lowland gorillas appear to represent the most inbred population, with evidence that they have been subjected to both recent and ancient inbreeding.

To examine the level of genetic differentiation between individuals we performed a principal component analysis (PCA) of SNP genotypes (Supplementary Information). Chimpanzees were stratified between subspecies with PC1 separating western and Nigeria–Cameroon chimpanzees from the eastern and central chimpanzees and PC2 separating western and Nigeria–Cameroon chimpanzees. In gorillas, PC1 clearly separates eastern and western gorillas, whereas the western lowland gorillas are distributed along a gradient of PC2, with individuals from the Congo and western Cameroon positioning in opposite directions along the axis. The isolated Cross River gorilla is genetically more similar to Cameroon western lowland gorillas and can be clearly differentiated with PC3 (Supplementary Fig. 8.2.9).

We explored the level of shared ancestry among individuals within each group¹⁴ using an admixture model (FRAPPE). In chimpanzees, the four known subspecies are clearly distinguished when fitting the model using four ancestry components ($K = 4$) (Fig. 1d). Additional substructure is identified among the eastern chimpanzees Vincent and Andromeda ($K = 6$), who hail from the most eastern sample site (Gombe National Park, Tanzania). As in Gonder *et al.*², we have identified three Nigeria–Cameroon samples (Julie, Tobi and Banyo, $K = 3$ – 5) with components of central chimpanzee ancestry. However, taking central chimpanzees and the remaining Nigeria–Cameroon chimpanzees as ancestral populations shows no evidence of gene flow by either the F3 statistic or HapMix. This indicates that these three samples are not the result of a recent admixture and may represent a genetically distinct population (Supplementary Information).

In gorillas, following the separation of eastern and western lowland species ($K = 2$), an increasing number of components further subdivide western lowland populations distinguishing Congolese and Cameroonian gorillas—a pattern consistent with the structure observed in the PCA analysis (Supplementary Fig. 8.2.9). One striking observation is the extent of admixed ancestry predicted for captive individuals when

compared to wild-born. Our analysis suggests that most captive individuals included in this study are admixed from two or more genetically distinct wild-born populations leading to an erosion of phylogeographic signal. This finding is consistent with microsatellite analyses of captive gorillas¹⁵ and the fact that great ape breeding programs have not been managed at the subspecies level.

As great apes have been evolving on separate lineages since the middle Miocene, we attempted to reconstruct the history of these various species and subspecies by applying methods sensitive to branching processes, changes in effective population size (N_e), and gene flow occurring at different time scales. Using a combination of speciation times inferred from a haploid pairwise sequential Markovian coalescent (PSMC) analysis¹⁶, a coalescent hidden Markov model (CoalHMM)³ and incomplete lineage sorting approaches, we were able to estimate the most ancient split times and effective population sizes among the great ape species. By combining these estimates with an approximate Bayesian computation (ABC)¹⁷ analysis applied to the more complex chimpanzee phylogeny, we constructed a composite model of great ape population history over the last ~15 million years (Fig. 2). This model presents a complete overview of great ape divergence and speciation events in the context of historical effective population sizes.

PSMC analyses of historical N_e (Fig. 3) suggests that the ancestral *Pan* lineage had the largest effective population size of all lineages >3 million years ago (Myr), after which time the population of the common ancestor of both bonobos and chimpanzees experienced a dramatic decline. Both PSMC and ABC analyses support a model of subsequent increase in chimpanzee N_e starting ~1 Myr, before their divergence into separate subspecies. Following an eastern chimpanzee increase in N_e (~500 thousand years ago, kyr), the central chimpanzees reached their zenith ~200–300 kyr followed by the western chimpanzee ~150 kyr. Although the PSMC profiles of the two subspecies within each of the major chimpanzee clades (eastern/central and Nigeria–Cameroon/western) closely shadow each other between 100 kyr and 1 Myr, the western chimpanzee PSMC profile is notable for its initial separation from that of the other chimpanzees, followed by its sudden rise and decline (Fig. 3 and Supplementary Information). The different gorilla species also show variable demographic histories over the past ~200 kyr. Eastern lowland gorillas have the smallest historical N_e , consistent with smaller present-day populations and a history of inbreeding (Fig. 1c). A comparison of effective population sizes with the ratio of non-synonymous to synonymous substitutions finds that selection has

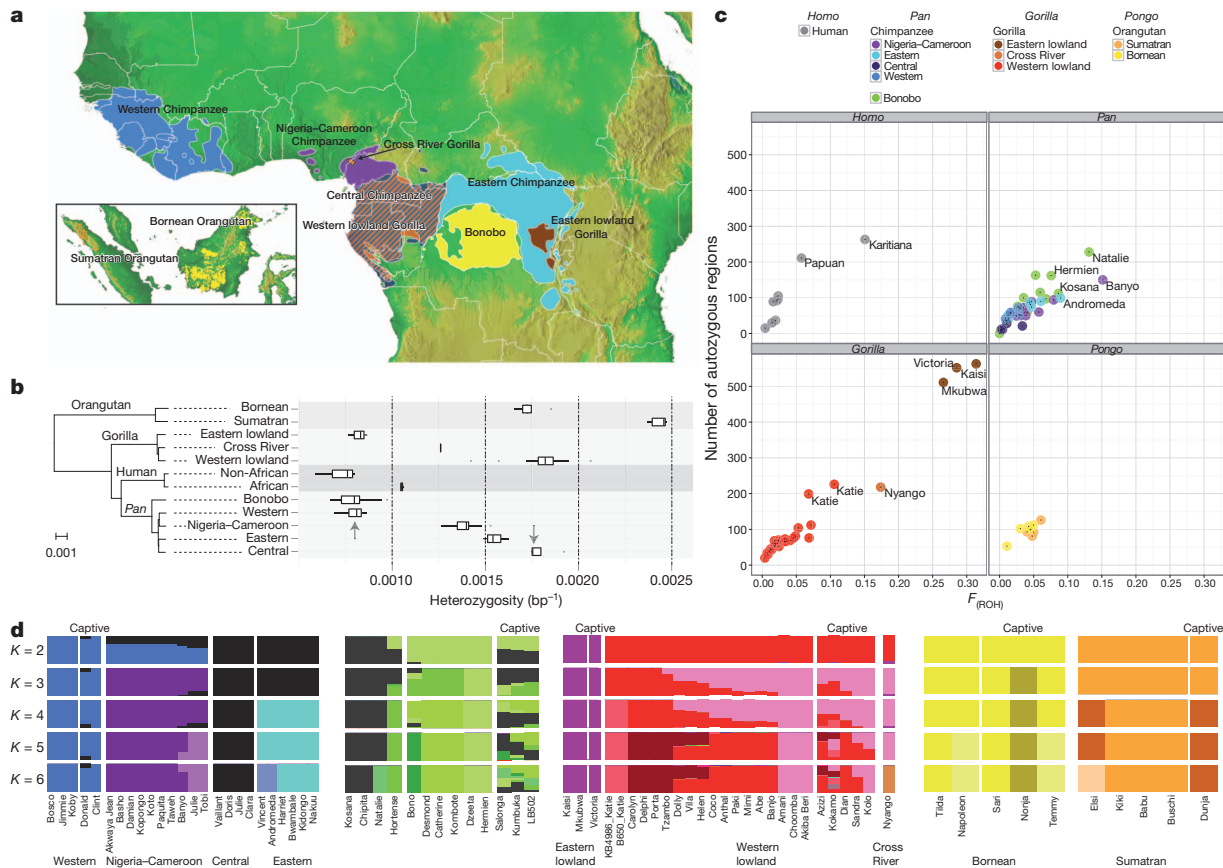


Figure 1 | Samples, heterozygosity and genetic diversity. **a**, Geographical distribution of great ape populations across Indonesia and Africa sequenced in this study. The formation of the islands of Borneo and Sumatra resulted in the speciation of the two corresponding orangutan populations. The Sanaga River forms a natural boundary between Nigeria–Cameroon and central chimpanzee populations whereas the Congo River separates the bonobo population from the central and eastern chimpanzees. Eastern lowland and western lowland gorillas are both separated by a large geographical distance. **b**, Heterozygosity estimates of each of the individual species and subspecies are superimposed onto a neighbour-joining tree from genome-wide genetic distance estimates (branch lengths in units of substitutions per bp). Arrows indicate heterozygosities previously reported³⁰ for western and central chimpanzee populations. **c**, Runs of homozygosity among great apes. The relationship

between the coefficient of inbreeding (F_{ROH}) and the number of autozygous >1 megabase segments is shown. Bonobos and eastern lowland gorillas show an excess of inbreeding compared to the other great apes, suggesting small population sizes or a fragmented population. **d**, Genetic structure based on clustering of great apes. All individuals (columns) are grouped into different clusters ($K = 2$ to $K = 6$, rows) coloured by species and according to their common genetic structure. Most captive individuals, labelled on top, show a complex admixture from different wild populations. A signature of admixture, for example, is clearly observed in the known hybrid Donald, a second-generation captive predicted to be a 15% admixture of central chimpanzee on a western background consistent with its pedigree. A grey line at the bottom denotes new groups at $K = 6$ in agreement with the location of origin or ancestral admixture.

acted more efficiently in populations with higher N_e , consistent with neutral theory (Supplementary Information).

Although the phylogeny of bonobos and western, central and eastern common chimpanzees has been well established based on genetic data¹⁸, there is still uncertainty regarding their relationship to Nigeria–Cameroon chimpanzees^{2,19}. Regional neighbour-joining trees and a maximum-likelihood tree estimated from allele frequencies both show that Nigeria–Cameroon and western chimpanzees form a clade. A complex demographic history has been previously reported for chimpanzees with evidence of asymmetrical gene flow among different subspecies. For instance, migration has been identified from western into eastern chimpanzees⁴, two subspecies that are currently geographically isolated. We find support for this using the D-statistic, a model-free approach that tests whether unequal levels of allele sharing between an outgroup and two populations that have more recently diverged ($D(H,W;E,C) > 16$ s.d.). However, no previous genome-wide analysis that has examined gene flow included chimpanzees from the Nigeria–Cameroon subspecies and a comparison of them with eastern chimpanzees results in a highly significant D-statistic ($D(H,E;W,N) > 25$ s.d.). Furthermore, TreeMix, a model-based approach that identifies gene flow events to explain allele frequency patterns not captured by a simple branching phylogeny, infers a signal of gene flow between Nigeria–Cameroon and

eastern chimpanzees ($P = 2 \times 10^{300}$). A more detailed treatment of gene flow applying different models and methods may be found in the Supplementary Information.

Genetic diversity is depressed at or close to genes in almost all species (Supplementary Fig. 11.1) with the effect less pronounced in subspecies with lower estimated N_e , consistent with population genetic theory. When we compare the relative level of X chromosome and autosomal (X/A) diversity across great apes as a function of genetic distance from genes, the eastern lowland gorillas and Bornean orangutans are outliers, with substantially reduced X/A diversity compared to the neutral expectation of 0.75, regardless of the distance to genes. This pattern is consistent with a recent reduction in effective population size²⁰, clearly visible in the PSMC analysis for both species (Fig. 3). However, bonobos also demonstrate a relatively constant level of X/A diversity regardless of distance from genes, with values very much in line with neutral expectations. All other subspecies demonstrate a pattern consistent with previous studies in humans²¹ where X/A diversity is lower than 0.75 close to genes and higher farther away from genes.

It has been proposed that loss of gene function may represent a common evolutionary mechanism to facilitate adaptation to changes in an environment²². There has been speculation that the success of humans may have, in part, been catalysed by an excess of beneficial

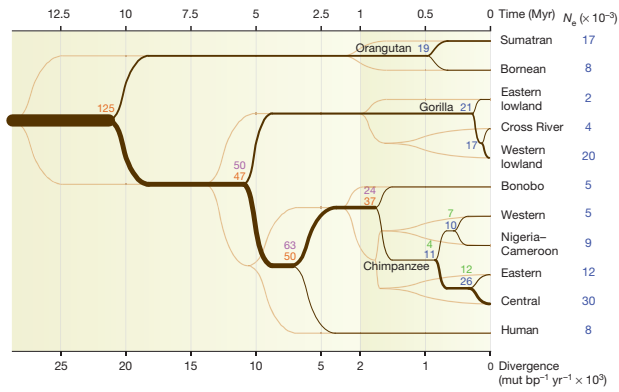


Figure 2 | Inferred population history. Population splits and effective population sizes (N_e) during great ape evolution. Split times (dark brown) and divergence times (light brown) are plotted as a function of divergence (d) on the bottom and time on top. Time is estimated using a single mutation rate (μ) of 1×10^{-9} mut bp $^{-1}$ year $^{-1}$. The ancestral and current effective population sizes are also estimated using this mutation rate. The results from several methods used to estimate N_e (COALHMM, ILS COALHMM, PSMC and ABC) are coloured in orange, purple, blue and green, respectively. The chimpanzee split times are estimated using the ABC method. The x axis is rescaled for divergences larger than 2×10^{-3} to provide more resolution in recent splits. All the values used in this figure can be found in Supplementary Table 5. The terminal N_e correspond to the effective population size after the last split event.

loss-of-function mutations²³. We thus characterized the distribution of fixed loss-of-function mutations among different species of great apes identifying nonsense and frameshift mutations resulting from SNVs ($n = 806$) and indels ($n = 1080$) in addition to gene deletion events ($n = 96$) (Supplementary Table 4). We assigned these events to the phylogeny and determined that the number of fixed loss-of-function mutations scales proportionally to the estimated branch lengths ($R^2 = 0.987$ SNVs, $R^2 = 0.998$ indels). In addition, we found no evidence of distortion on the terminal branches of the tree compared to point mutations based on a maximum likelihood analysis (Supplementary Information). Thus, the human branch in particular showed no excess of fixed loss-of-function mutations even after accounting for human-specific pseudogenes²⁴ (Supplementary Information).

Our analysis provides one of the first genome-wide views of the major patterns of evolutionary diversification among great apes. We have generated the most comprehensive catalogue of SNPs for chimpanzees (27.2 million), bonobos (9.0 million), gorillas (19.2 million) and orangutans (24.3 million) (Table 1) to date and identified several thousand AIMs, which provides a useful resource for future analyses of ape populations. Humans, western chimpanzees and eastern gorillas all show a remarkable dearth of genetic diversity when compared to other great apes. It is striking, for example, that sequencing of 79 great ape genomes identifies more than double the number of SNPs obtained from the recent sequencing of more than a thousand diverse humans²⁵—a reflection of the unique out-of-Africa origin and nested phylogeny of our species.

We provide strong genetic support for distinct populations and subpopulations of great apes with evidence of additional substructure. The common chimpanzee shows the greatest population stratification when compared to all other lineages with multiple lines of evidence supporting two major groups: the western and Nigeria–Cameroon and the central and eastern chimpanzees. The PSMC analysis indicates a temporal order to changes in ancestral effective population sizes over the last two million years, previous to which the *Pan* genus suffered a dramatic population collapse. Eastern chimpanzee populations reached their maximum size first, followed by the central and western chimpanzee. The Nigeria–Cameroon chimpanzee population size appears much more constant.

Despite their rich evolutionary history, great apes have experienced drastic declines in suitable habitat in recent years²⁶, along with declines in local population sizes of up to 75% (ref. 27). These observations highlight the urgency to sample from wild ape populations to more fully understand reservoirs of genetic diversity across the range of each species and to illuminate how basic demographic processes have affected it. The >80 million SNPs we identified in this study may now be used to characterize patterns of genetic differentiation among great apes in sanctuaries and zoos and, thus, are of great importance for the conservation of these endangered species with regard to their original range. These efforts will greatly enhance conservation planning and management of apes by providing important information on how to maintain genetic diversity in wild populations for future generations.

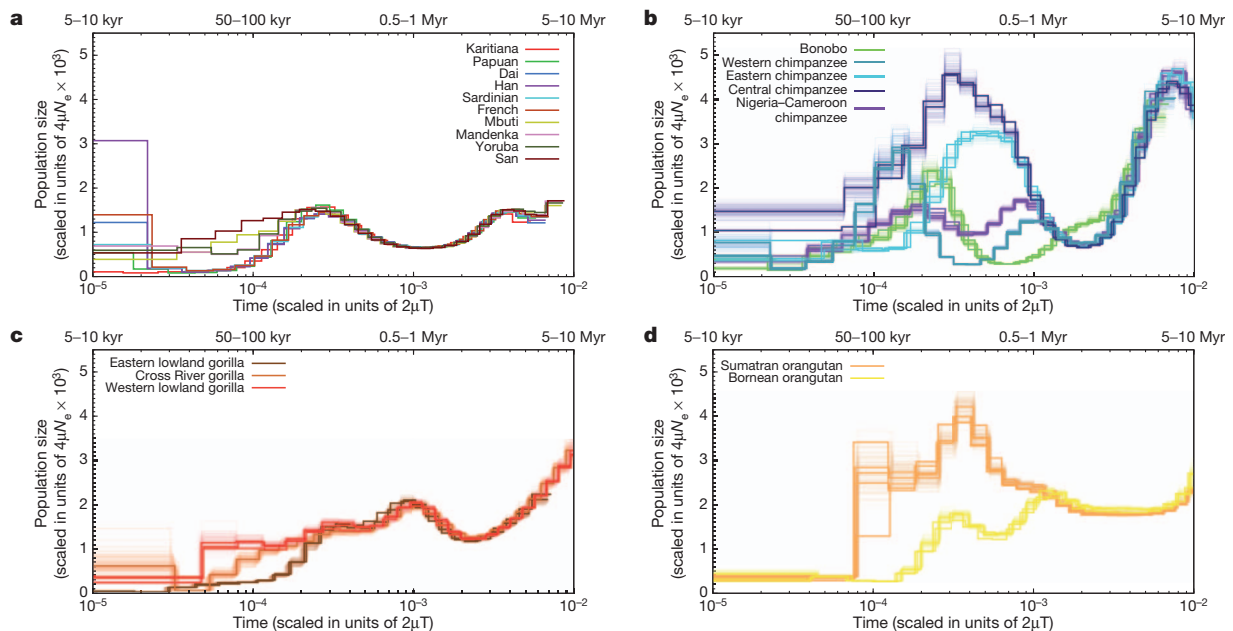


Figure 3 | PSMC analysis. Inferred historical population sizes by pairwise sequential Markovian coalescent analysis. The lower x axis gives time measured by pairwise sequence divergence and the y axis gives the effective population size measured by the scaled mutation rate. The upper x axis indicates scaling in

years, assuming a mutation rate ranging from 10^{-9} to 5×10^{-10} per site per year. The top left panel shows the inference for modern human populations. In the rest of the three panels, thin light lines of the same colour correspond to PSMC inferences on 100 rounds of bootstrapped sequences.

METHODS SUMMARY

We sequenced to a mean coverage of $25\times$ (Illumina HiSeq 2000) a total of 79 great ape individuals, representing 10 subspecies and four genera of great apes from a variety of populations across the African continent and Southeast Asia. SNPs were called using GATK¹² after BWA²⁸ mapping to the human genome (NCBI Build 36) using relaxed mapping parameters. Samples combined by species were realigned around putative indels. SNP calling was then performed on the combined individuals for each species. For indels, we used the GATK Unified Genotyper to produce an initial set of indel candidates applying several quality filters and removing variants overlapping segmental duplications and tandem repeats. We also removed groups of indels clustering within 10 bp to eliminate possible artefacts in problematic regions. Conservative allelic imbalance filters were used to eliminate false heterozygotes that may affect demographic analyses, some of which are sensitive to low levels of contamination. We estimate that the application of this filter resulted in a 14% false negative rate for heterozygotes. Our multispecies study design facilitated this assessment of contamination, which may remain undetected in studies focused on assessing diversity within a single species. The amount of cross-species contamination was estimated from the amount of non-endogenous mitochondrial sequence present in an individual. Because we wished to compare patterns of variation between and within species, we report all variants with respect to coordinates of the human genome reference. For FRAPPE analyses, we used MAF0.06 (human, orangutan and bonobo) and 0.05 (chimpanzee and gorilla) to remove singletons. For most of the analyses, we only used autosomal markers, except in the X/A analysis. To determine the amount of inbreeding, we calculated the heterozygosity genome-wide in windows of 1 megabase with 200-kilobase sliding windows. We then clustered together the neighbouring regions to account for runs of homozygosity. For the PSMC analyses, we called the consensus bases using SAMtools²⁹. Underlying raw sequence data are available through the Sequence Read Archive (SRA) (PRJNA189439 and SRP018689). Data generated in this work are available from (<http://biologiaevolutiva.org/greatape/>). A complete description of the materials and Methods is provided in the Supplementary Information.

Received 30 December 2012; accepted 26 April 2013.

Published online 3 July 2013.

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Supplementary Information is available in the online version of the paper.

Acknowledgements We thank the following funding agencies: ERC Starting Grant (260372) to T.M.-B.; NIH grants HG002385 to E.E.E., R01_HG005226 to K.R.V., A.E.W., M.F.H., L.S. and J.D.W., GM100233 and NSF HOMINID grant 1032255 to D.R. and He.Li.; MICINN (Spain) BFU2011-28549 to T.M.-B., BFU2010-19443 to Ja.Be., Spanish Government and FEDER for grants BFU2009-13409-C02-02 and BFU2012-38236 to A.N. and J.P.-M., Direcció General de Recerca, Generalitat de Catalunya (Grup de Recerca Consolidat 2009 SGR 1101) to Ja.Be., D.C., A.N. and T.M.-B.; ERC Advanced Grant (233297) and Max Planck Society to S. Paabo; Danish Council for Independent Research Natural Sciences to H.S.; Spanish Grant (CGL-2010-20170) and Zoo de Barcelona (Beca PRIC) to A.R.-H.; EUPRIM-Net to BPRC; DP1ES022577-04 NIH grant to S.A.T.; NSF Grant 0755823 to M.K.G.; P.G. is supported by the G. Harold and Leila Y. Mathers Foundation. A.N. and T.M.-B. are ICREA Research Investigators (Institut Català d'Estudis i Recerca Avancats de la Generalitat de Catalunya). J.P.-M. is supported by the Zoo de Barcelona and l'Ajuntament de Barcelona. P.H.S. is supported by an HHMI International Student Fellowship. E.E.E. is an investigator with the Howard Hughes Medical Institute. We are especially grateful to all those who generously provided the samples for the project: O. Thalmann and H. Siedel from Limbe Sanctuary; R. Garriga from Tacugama Sanctuary; W. Schempp (University of Freiburg), Burgers' Zoo; Zoo of Antwerp; Wilhelma Zoo; Givskud Zoo; Ngamba Island Chimpanzee Sanctuary and Centre de Primatologie; Centre International de Recherches Médicales de Franceville; North Carolina Zoological Park; Zoo Atlanta; the Lincoln Park Zoo (Chicago); the Antwerp Zoo and the Limbe Wildlife Centre (Cameroon); D. Travis from University of Minnesota and M. Kinsel from University of Illinois Urbana-Champaign and S. Paabo and L. Vigilant, Max Planck Institute for Evolutionary Anthropology. We thank T. Brown for revising the manuscript, L. Capilla and E. Eyraas for technical support, and M. Dierssen for comments on genes expressed in the brain.

Author Contributions E.E.E. and T.M.-B. designed the study. J.P.-M., P.H.S., J.M.K., J.L.K., B.L.-G., M.D., M.F.-C., J.C.M., C.D.B., E.E.E. and T.M.-B. analysed the raw data and performed the variant calling. J.P.-M., P.H.S., Ma.Ma., J.H.-R., I.H.-H., T.C., C.B., L.V., A.R.-H. and C.C. validated the different variants. J.P.-M., P.H.S., B.L.-G., C.A., F.H., E.E.E. and T.M.-B. analysed large deletions. K.R.V., L.J., A.E.W. and M.F.H. analysed the X/Autosome diversity. D.T., G.S., A.C., C.T., F.C., Ha.La., K.P., M.P., M.L., N.P., D.C., Ja.Be., A.N. and A.M.A. performed selection analyses. J.P.-M., P.H.S., J.M.K., J.L.K., T.D.O'C., He.Li., D.R., K.M., A.H., A.E.H., M.H.S., C.H., J.M.A., T.M., C.D.B., E.E.E. and T.M.-B. analysed different aspects of demography. M.L.W., L.S., T.A., I.K., A.P., F.L., J.A.K., E.L., P.G., H.S., M.K.G., S.A.T., R.A.B., R.E.B., O.A.R. and B.H.H. provided critical samples and participated in the discussion of phylogeny. L.F., R.K.W., Ju.Bl., E.E.E., Ma.Ma., L.A.-C., M.G. and I.G.G. generated genome libraries and produced the genome sequence associated with this project. All authors contributed to data interpretation. J.P.-M., P.H.S., E.E.E. and T.M.-B. drafted the manuscript with input from all authors.

Author Information Underlying raw sequence data are available through the Sequence Read Archive (SRA) (PRJNA189439 and SRP018689). Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to T.M.-B. (tomas.marques@upf.edu) or E.E.E. (eee@gs.washington.edu).



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