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Origin of the human malaria parasite *Plasmodium falciparum* in gorillas

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Plasmodium falciparum is the most prevalent and lethal of the malaria parasites infecting humans, yet the origin and evolutionary history of this important pathogen remain controversial. Here we develop a single-genome amplification strategy to identify and characterize *Plasmodium* spp. DNA sequences in faecal samples from wild-living apes. Among nearly 3,000 specimens collected from field sites throughout central Africa, we found *Plasmodium* infection in chimpanzees (*Pan troglodytes*) and western gorillas (*Gorilla gorilla*), but not in eastern gorillas (*Gorilla beringei*) or bonobos (*Pan paniscus*). Ape plasmodial infections were highly prevalent, widely distributed and almost always made up of mixed parasite species. Analysis of more than 1,100 mitochondrial, apicoplast and nuclear gene sequences from chimpanzees and gorillas revealed that 99% grouped within one of six host-specific lineages representing distinct *Plasmodium* species within the subgenus *Laverania*. One of these from western gorillas comprised parasites that were nearly identical to *P. falciparum*. In phylogenetic analyses of full-length mitochondrial sequences, human *P. falciparum* formed a monophyletic lineage within the gorilla parasite radiation. These findings indicate that *P. falciparum* is of gorilla origin and not of chimpanzee, bonobo or ancient human origin.

Malaria is a blood infection caused by mosquito (Anopheles spp.)borne apicomplexan parasites of the genus *Plasmodium*¹⁻³. Of five Plasmodium species known to infect humans, P. falciparum causes by far the greatest morbidity and mortality, with several hundred million cases of clinical malaria and more than one million deaths occurring annually^{1,2}. Although much progress has been made in the treatment and prevention of P. falciparum⁴, the origin and natural reservoirs of this and related plasmodial pathogens remain controversial. Until recently, the closest known relative of P. falciparum was a chimpanzee parasite, Plasmodium reichenowi, which was assumed to have diverged from its human counterpart at the same time as the ancestors of chimpanzees and humans, more than 5,000,000 years ago⁵⁻⁸. Within the past year, other closely related *Plasmodium* strains have been detected in chimpanzees, western gorillas and bonobos, raising the possibility that P. falciparum in humans could have arisen as a consequence of cross-species transmission from one or more of these apes⁹⁻¹². However, all of these studies were limited by an analysis of only few apes, many of which were captive and living in close proximity to humans. In addition, all prior studies used non-limiting dilution polymerase chain reaction (PCR) amplification methods that are prone to generating artefactual mosaic sequences by recombination between genetically distinct templates. Here we used conventional and single-template PCR amplification methods to screen and analyse wild-living chimpanzee, bonobo and gorilla populations across sub-Saharan Africa for parasites related to P. falciparum.

Prevalence of ape Plasmodium infections

To determine the geographic distribution, species association and prevalence of ape *Plasmodium* spp. infections, we adapted a previously described PCR-based diagnostic method¹⁰ to amplify a 956-base-pair (bp) fragment of *Plasmodium* cytochrome b (cytb) sequences from faecal DNA (Supplementary Fig. 1a). Ape faecal samples were selected from existing specimen banks that we had collected earlier for molecular epidemiological studies of simian retrovirus infections^{13–16}. Except for 28 samples from one habituated gorilla community at field site DS (Fig. 1), all other specimens were derived from non-habituated apes living in remote forest areas (Supplementary Table 1). Chimpanzee (n = 1,827), gorilla (n = 805) and bonobo (n = 107) samples were subjected to diagnostic PCR, and all amplification products were sequenced to confirm Plasmodium infection. In addition, we subjected a subset of samples (n = 1,027), including all specimens from eastern gorillas and bonobos, to microsatellite analysis of host genomic DNA14-16 to determine the number of individuals tested at particular field sites (Supplementary Table 1). Microsatellite analysis also provided quantitative estimates of specimen integrity (Supplementary Table 2) and redundant sampling (Supplementary Table 3), thereby allowing us to determine the sensitivity of the non-invasive diagnostic test by identifying the proportion of PCR-positive specimens from infected apes that were sampled more than once. From 32 such individuals, we estimated the test sensitivity to be 57% (Supplementary Table 4) and calculated the prevalence of ape infection at each field

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Figure 1 | **Location of ape study sites.** Field sites are shown in relation to the ranges of three subspecies of the common chimpanzee (*P. t. ellioti, P. t. troglodytes* and *P. t. schweinfurthii*), the western (*G. gorilla*) and eastern (*G. beringei*) gorillas, and the bonobo (*P. paniscus*) in sub-Saharan Africa (coloured as indicated). Forested areas are indicated in green, and arid areas are

site (Supplementary Table 1). The results revealed widespread *Plasmodium* infection in chimpanzees and western gorillas, but not in eastern gorillas or bonobos.

Ape malaria parasites were detected at 32 of 45 chimpanzee collection sites, and at 17 of 20 western gorilla collection sites (Fig. 1), including every site where at least ten individuals were estimated to have been sampled. Plasmodium infection was endemic in Nigeria-Cameroon chimpanzees (Pan troglodytes ellioti), central (Pan troglodytes troglodytes) and eastern (Pan troglodytes schweinfurthii) chimpanzees, and in western lowland gorillas (Gorilla gorilla gorilla), with estimated prevalence rates ranging from 32% to 48% (Table 1). The true infection rates are likely to be even higher, because Plasmodium detection in faecal samples can be expected to be less sensitive than detection in blood, as is the case for urine and saliva samples¹⁷. Although wild-living western chimpanzees (Pan troglodytes verus) and Cross River gorillas (Gorilla gorilla diehli) were not tested in this study, these two subspecies have previously been shown to harbour *Plasmodium* parasites in the wild^{9,10}. On the basis of these data, it is clear that chimpanzees and western gorillas represent substantial Plasmodium reservoirs. Unexpectedly, we did not find this to be true for eastern gorillas or bonobos. Screening 71 and 58 members of these respective species at multiple field sites, we failed to detect Plasmodium infection in any of them (Supplementary Table 1). These findings suggest that malaria parasites are rare or absent in some wild-living ape communities, possibly reflecting regional, ecological or seasonal differences in the distribution and/ or host specificities of the transmitting mosquito vectors. Additional field studies are needed to determine whether eastern gorillas and bonobos are infected by Plasmodium parasites at other locations or if they harbour divergent parasites not detected by current diagnostic assays.

shown in yellow (map courtesy L. Pintea, The Jane Goodall Institute). Circles, squares and hexagons identify field sites where chimpanzees, gorillas or both were sampled, respectively; ovals indicate bonobo collection sites. Sites where ape malaria was detected are highlighted in yellow, with red lettering indicating that both chimpanzees and gorillas were infected.

Analysis of ape *Plasmodium* species by single-genome amplification

To examine the evolutionary relationships of the newly identified Plasmodium parasites, we constructed phylogenetic trees for a subset of the diagnostic cytb sequences. This analysis showed that all sequences, except for one Plasmodium ovale-like strain, fell into one large monophyletic clade that also included P. reichenowi and P. falciparum (Supplementary Fig. 2). Parasites related to P. reichenowi and P. falciparum have previously been classified into a subgenus, termed Laverania, to distinguish them from more-divergent Plasmodium species¹⁸. Our results thus indicated that parasites from this subgenus were common and widespread among wild ape populations. However, the topology of the Laverania clade was highly unusual, characterized by only few discrete clades and multiple sequences with very short branches attached to internal branches. Moreover, repeated PCR analysis of the same faecal samples yielded sequences that clustered variably in different parts of the tree (Supplementary Fig. 2). These findings indicated simultaneous infection with genetically diverse Plasmodium parasites and suggested that conventional (bulk) PCR amplification had generated in vitro recombinants. To examine this possibility, we reanalysed the same Plasmodium-positive faecal samples by single-genome amplification (SGA), a molecular strategy that has been used extensively to characterize the genetic identity and quasispecies complexity of human and simian immunodeficiency viruses¹⁹⁻²³. Faecal DNA was diluted so that fewer than 30% of all PCR reactions yielded an amplification product, which ensured amplification of single Plasmodium templates in most reactions¹⁹⁻²³. All amplicons were sequenced directly and sequences containing mixed bases indicative of more than one amplified template were discarded. Using this approach to characterize the genetic complexity of malaria parasites in faecal samples, we could

Table 1 | Prevalence of Plasmodium spp. infection in wild-living African apes

Species or subspecies	Field sites*	Faecal samples tested	Faecal samples positive†	Plasmodium spp. prevalence (CI)‡
Nigeria–Cameroon chimpanzee (Pan troglodytes ellioti)	9	119	19	32% (23–46%)
Central chimpanzee (Pan troglodytes troglodytes)	13	612	147	48% (44-53%)
Eastern chimpanzee (Pan troglodytes schweinfurthii)	24	1,096	187	34% (30-40%)
Western lowland gorilla (Gorilla gorilla gorilla)	20	659	120	37% (32-41%)
Eastern lowland gorilla (Gorilla beringei graueri)	3	146	0	0% (0-4%)
Bonobo (Pan paniscus)	2	107	0	0% (0–6%)

* Field sites are listed in Supplementary Table 1 and their location is shown in Fig. 1.

+ Faecal samples were tested for Plasmodium mitochondrial (cytb) DNA by diagnostic PCR; all amplification products were sequenced to confirm Plasmodium infection.

* Percentage prevalence of *Plasmodium* infection with brackets indicating 95% confidence intervals (CI). Values were estimated on the basis of the proportion of PCR-positive faecal samples, correcting for specimen degradation (Supplementary Table 2), repeat sampling (Supplementary Table 3), and the sensitivity of the diagnostic test (Supplementary Table 4). We note that these prevalence rates represent minimum estimates, because the extent to which infected apes shed *Plasmodium* DNA into their faeces is not known.



eliminate both Taq-polymerase-induced recombination (template switching) and nucleotide misincorporations in finished sequences, thereby ensuring an accurate representation of plasmodial variants as they existed *in vivo*^{21–23}.

Figure 2 depicts the phylogenetic relationships of a subset of SGAderived mitochondrial cytb sequences (the entire set of 697 sequences is analysed in Supplementary Fig. 3). As in the corresponding tree of bulk-PCR-derived sequences (Supplementary Fig. 2), all SGA-derived sequences, except for seven P. ovale, Plasmodium vivax and Plasmodium malariae-like strains, grouped within the Laverania radiation. However, unlike in the bulk PCR tree, Laverania sequences in the SGA tree clustered in a strictly host-species-specific manner, forming three chimpanzee-specific (C1-C3) and three gorilla-specific (G1-G3) clades, each supported by high bootstrap values. This host specificity did not extend to the subspecies level, because P. t. ellioti, P. t. troglodytes and P. t. schweinfurthii-derived sequences were interspersed; however, cytb sequences from P. t. schweinfurthii segregated into distinct subclades within two of the three chimpanzee lineages (C1 and C2), suggesting a phylogeographic distribution of certain Plasmodium variants (Supplementary Fig. 3a, b). None of 363 chimpanzee-derived Plasmodium cytb sequences was closely related to human P. falciparum. Instead, all human sequences grouped within a single clade of parasites (G1) that infected western gorillas at numerous sites in Cameroon (LB, BB, CP, NK, BQ, DD, MM and LM), the Central African Republic (DS and ND) and the Republic of the Congo (GT) (Fig. 2 and Supplementary Fig. 3d). A notable finding of the SGA analysis, which was obscured by bulk PCR analysis, was that most apes were co-infected with parasites representing multiple different plasmodial lineages, including variants from the same Laverania clade, from different Laverania clades or from Laverania and non-Laverania clades (Supplementary Table 5). Of 65 chimpanzee and 53 gorilla samples characterized, 48 (74%) and 37 (70%), respectively, harboured more than one genetically distinct parasite strain, and 36 (55%) and 23 (43%) contained members of two or more major *Plasmodium* clades (Supplementary Fig. 3). Given this high frequency of co-infection with divergent parasites, conventional recombinationprone PCR approaches are not appropriate for generating ape Plasmodium sequences for phylogenetic analysis. Moreover, previously reported ape *Plasmodium* sequences⁹⁻¹² must be interpreted with caution because they were subject to these same confounding factors.

To test the robustness of the phylogenetic relationships depicted in Fig. 2, we used SGA to amplify additional genomic regions from *cytb*-positive faecal samples, targeting loci in the mitochondrial, apicoplast and nuclear *Plasmodium* genomes. These regions included 390 bp of the caseinolytic protease C (*clpC*) gene (n = 126), 772 bp of the lactase dehydrogenase (*ldh*) gene (n = 46), and 3.4-kilobase (kb) (n = 165) and 3.3-kb (n = 127) fragments that together spanned the entire mitochondrial genome (Supplementary Fig. 1a). Phylogenetic analyses of each of these genomic loci revealed very similar topologies. In trees of *clpC* (Supplementary Fig. 4), *ldh* (Supplementary Fig. 5) and mitochondrial sequences (Supplementary Figs 6 and 7), *Laverania* sequences formed the same number of chimpanzee-specific (C1–C3) and

Figure 2 Phylogeny of *Plasmodium* parasites from wild-living chimpanzees and western gorillas. A representative subset of 146 SGAderived *Plasmodium* mitochondrial cytochrome *b* sequences (956 bp) is shown in relation to human and simian *Plasmodium* reference sequences (for accession numbers, see Supplementary Tables 6 and 7). The full set of 697 SGAderived ape *Plasmodium* sequences is analysed in Supplementary Fig. 3. Sequences are colour-coded, with capital letters indicating the field site (Fig. 1) and lower-case letters denoting species and subspecies origin (ptt: *P. t. troglodytes*, red; pte: *P. t. ellioti*, orange; pts: *P. t. schweinfurthii*, blue; gor: *G. g. gorilla*, green). C1–C3 and G1–G3 represent chimpanzee- and gorilla-specific *Plasmodium* species, respectively, all of which are included within the subgenus *Laverania*¹⁸. The tree was inferred using maximum-likelihood methods³⁰. Bootstrap values (>70%) are indicated for major nodes only (the scale bar represents 0.01 substitutions per site). gorilla-specific (G1–G3) clades, albeit with some variations in the relationships among these lineages. Importantly, there was no evidence of recombination between chimpanzee- and gorilla-specific parasites, although many of them infected apes at the same field sites. This suggested that *Laverania* parasites are largely host specific (recombination between parasites infecting the same host species could not be assessed because of mixed *Plasmodium* infections). These findings, together with the extent of genetic diversity that distinguishes the various clades, argue strongly for the existence of six distinct *Plasmodium* species within the *Laverania* subgenus (Supplementary Figs 3–8). Formal classification of these lineages must await additional taxonomic evaluation.

Origin of human P. falciparum

The new SGA-derived ape Plasmodium sequences call for a reassessment of the origin of human P. falciparum. Among over 600 sequences derived from ape samples spanning most of central Africa, we failed to find a single chimpanzee parasite that was sufficiently closely related to P. falciparum to represent a progenitor (Fig. 2 and Supplementary Figs 3-8). Thus, P. reichenowi, as well as other chimpanzee Plasmodium species, can be excluded as precursors of P. falciparum. Instead, all new phylogenetic evidence points to a western gorilla origin of human P. falciparum (Fig. 2 and Supplementary Figs 3-8). To investigate how often gorilla parasites might have colonized humans, we constructed phylogenetic trees from complete mitochondrial genome equivalents of the closest Plasmodium relatives of human P. falciparum (Fig. 3). In a tree of concatenated CYTB, COXI and COXIII protein sequences (980 amino acids), all available human *P. falciparum* sequences (n = 105) coalesced to a single common ancestor nested within the G1 clade of gorilla parasites (Fig. 3a). Nucleotide sequences from the remaining

(non-coding) portions of the mitochondrial genome yielded a very similar topology, again showing that human P. falciparum formed a monophyletic lineage within the gorilla P. falciparum radiation (Fig. 3b). These findings, together with the observation that human parasites exhibit substantially less sequence diversity than the various ape Plasmodium species, including the closest gorilla relatives (Table 2), provide compelling evidence for a gorilla origin of human P. falciparum. Moreover, the monophyly of the human parasite sequences (Fig. 3) may indicate that all extant human strains evolved from a single gorilla-to-human cross-species transmission event. Notably, four recently reported Plasmodium sequences from captive bonobos¹¹ also clustered closely with *P. falciparum*. However, unlike the gorilla sequences, the bonobo sequences were interspersed with the human sequences (Fig. 3). This finding, together with the fact that the bonobo parasites encoded dihydrofolate reductase/thymidylate synthase (dhfr-ts) drug-resistance mutations prevalent in the local human population¹¹, suggests that the bonobos became infected with human parasites while housed in an urban sanctuary. In fact, the topologies in Fig. 3 are consistent with more than one human-tobonobo transmission, although some (or all) of the substitutions that distinguish bonobo and human sequences could represent PCR misincorporations because they were not generated by SGA methods¹¹.

Discussion

Using single-template amplification strategies and a much larger collection of ape specimens than previously analysed, we show here that wild-living chimpanzees and western gorillas are naturally infected with at least nine *Plasmodium* species. Among more than 1,100 SGAderived mitochondrial, apicoplast and nuclear gene sequences from 80 chimpanzee and 55 gorilla samples, we found a total of nine





bonobos¹¹ (Bo), which are shown in magenta. Human *P. falciparum* (Hu) and chimpanzee reference sequences are depicted in black, with the number of human sequences representing the same haplotype shown in parentheses (for accession numbers, see Supplementary Table 7). The trees were inferred using maximum-likelihood methods³⁰; numbers at nodes indicate bootstrap (above) and posterior probability (below) values, respectively (only bootstrap values >70 and posterior probabilities >0.7 are shown). The scale bars represent one amino acid (aa; **a**) and one nucleotide (nt) substitution per site (**b**).

Table 2 Genetic diversity within different Laverania species

Species	cytb*			mtDNA-3.4 kb*			mtDNA-3.3 kb*					
	п	Mean	Median	Max.	n	Mean	Median	Max.	n	Mean	Median	Max.
C1	58	0.0055	0.0031	0.0136	25	0.0069	0.0024	0.0148	9	0.0035	0.0052	0.0060
G1(gor/all)†	51	0.0085	0.0021	0.0251	31	0.0058	0.0015	0.0193	19	0.0029	0.0016	0.0108
G1(gor/main)†	40	0.0018	0.0011	0.0073	26	0.0012	0.0012	0.0027	17	0.0015	0.0012	0.0040
G1(hum)†	105	0.0005	0	0.0021	105	0.0003	0.0003	0.0015	105	0.0002	0	0.0012
C2	109	0.0077	0.0021	0.0262	39	0.0078	0.0024	0.0202	16	0.0044	0.0020	0.0112
G2	92	0.0031	0.0031	0.0115	25	0.0032	0.0036	0.0071	36	0.0015	0.0016	0.0036
C3	48	0.0029	0.0031	0.0094	13	0.0035	0.0030	0.0071	8	0.0026	0.0028	0.0044
G3	23	0.0017	0.0021	0.0042	14	0.0010	0.0009	0.0033	4	0	0	0

* Pairwise distances were determined for SGA-derived cytb, mtDNA-3.4 kb and mtDNA-3.3 kb sequences for each of the six *Laverania* species as shown in Supplementary Figs 3, 6 and 7, respectively. The number of sequences (n) analysed for each clade is indicated. Sequences were aligned and gap-stripped to remove ambiguous sites. Alignments were then used to determine the mean, median and maximum sequence distances. Identical sequences form different samples were included (identical sequences from the same sample were excluded). The sequence overlap between the mtDNA-3.3 kb and mtDNA-3.4 kb fragments was removed from the mtDNA-3.3 kb comparisons.

† The *P. falciparum* clade (G1) was subdivided into three separate groups: G1(gor/all) included all gorilla *P. falciparum* sequences, G1(gor/main) included only the closest gorilla relatives of human *P. falciparum* and G1(hum) included only human *P. falciparum* sequences. The comparisons show that human *P. falciparum* strains have much lower diversity than the various clades of ape-derived strains (except for G3, where only very few sequences were available for the 3.3-kb fragment); in particular, the G1(gor/main) clade has substantially higher diversity than the G1(hum) clade.

sequences that were related to P. malariae, P. ovale or P. vivax (Supplementary Table 5). All others grouped within one of six chimpanzeeor gorilla-specific lineages representing distinct Plasmodium species, three of which had not previously been described. Significantly, all currently available human P. falciparum sequences constitute a single lineage nested within the G1 clade of gorilla parasites. This indicates that human P. falciparum is of gorilla origin, and not of chimpanzee9,10,12, bonobo¹¹ or ancient human⁵ origin, and that all known human strains may have resulted from a single cross-species transmission event. What is still unclear is when gorilla P. falciparum entered the human population and whether present-day ape populations represent a source for recurring human infection. It has been suggested that the limited levels of genetic diversity seen at many loci in human P. falciparum reflect a relatively recent selective sweep⁸. Our data suggest that this bottleneck or 'Eve event' was instead the consequence of cross-species transmission of a gorilla parasite. It is difficult to date this event without having reliable dates with which to calibrate the *Plasmodium* phylogenetic trees. Previous estimates of dates in the evolution of *Plasmodium* have relied largely on the belief that P. falciparum and P. reichenowi diverged at the same time as the ancestors of humans and chimpanzees^{6-8,24}, an assumption that is now groundless. Others have proposed a much shorter timescale coincident with the emergence of agricultural societies in sub-Saharan Africa, the incomplete penetration of protective human gene polymorphisms (for example haemoglobin C) that are selected by P. falciparum infection, or the speciation of African mosquito vectors^{3,25}. Complete sequence analysis of members of the ape Plasmodium species identified here may help to resolve this conundrum. The second question, of whether additional cross-species transmissions of Laverania parasites have given rise to human infections, is more immediately addressable. An alignment of over 100 ape Plasmodium mitochondrial genome sequences reveals ape-specific single nucleotide polymorphisms (Supplementary Fig. 1b), which can now be used to screen plasmodial sequences from humans living in close proximity to wild gorillas and chimpanzees. Such studies can inform malaria eradication efforts about potential zoonotic Plasmodium reservoirs and provide insights into adaptive changes that might be required for ape Plasmodium infection of humans²⁶.

METHODS SUMMARY

Ape samples. We selected faecal samples from wild-living chimpanzees, gorillas and bonobos from existing specimens banks¹³⁻¹⁶ on the basis of their geographic location, available host genetic information, and species and subspecies origin (Supplementary Table 1).

Conventional PCR. *Plasmodium* mitochondrial, apicoplast and nuclear gene sequences were amplified as previously described^{9,10,27,28}, but using modified primers and PCR conditions suitable for faecal DNA. Bulk-PCR-positive faecal samples were subsequently subjected to SGA analysis.

Single-genome amplification. We performed SGA analysis of *Plasmodium* sequences from faecal DNA as previously described^{21,22}. All amplicons were sequenced directly, and sequences containing double peaks were discarded.

Prevalence estimations. The prevalence rates of *Plasmodium* infection were estimated on the basis of the proportion of PCR-positive faecal samples, correcting for specimen degradation, repeat sampling and the sensitivity of the diagnostic test.

Phylogenetic analysis. We inferred the phylogenetic trees of newly derived ape *Plasmodium* mitochondrial, apicoplast and nuclear gene sequences by Bayesian²⁹ and maximum-likelihood methods³⁰ (GenBank accession numbers are listed in Supplementary Table 6).

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

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Author Contributions All authors contributed to the acquisition, analysis and interpretation of the data; W.L., M.P., J.C.R., P.M.S. and B.H.H. initiated and designed the study; W.L., Y.L. and J.D.R. performed non-invasive *Plasmodium* testing and SGA analyses; B.F.K, R.S.R and J.D.R. performed microsatellite analyses; P.M.S. calculated *Plasmodium* prevalence rates; G.H.L. and P.M.S performed phylogenetic analyses; J.-B.N.N., C.M.S., D.B.M., S.L., M.K.G., P.J.K., P.D.W., E.D., E.M.-N., A.V.G. and M.N.M. conducted and supervised all fieldwork; and W.L., G.M.S., M.P., P.M.S., J.C.R. and B.H.H. coordinated the contributions of all authors and wrote the paper.

Author Information SGA-derived *Plasmodium* nucleotide sequences have been deposited in GenBank under accession numbers HM234976–HM235117 and HM237301 (*cytb*), HM235118–HM235143 (*ldh*), HM235144–HM235170 (*clpC*), HM235171–HM235268 (mtDNA-3.3 kb) and HM235269–HM235404 (mtDNA-3.4 kb) (also see Supplementary Table 6). 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 this article at www.nature.com/nature. Correspondence and requests for materials should be addressed to B.H.H. (bhahn@uab.edu).

METHODS

Ape faecal samples. To screen wild ape populations for *Plasmodium* infection, we selected 2,739 faecal samples from an existing bank of chimpanzee (P. troglodytes), western gorilla (G. gorilla), eastern gorilla (G. beringei) and bonobo (P. paniscus) specimens previously collected for molecular epidemiological studies of simian immunodeficiency virus $({\rm SIV}_{\rm CPZ} \text{ and } {\rm SIV}_{\rm GOR})^{14-16,31}$ and simian foamy virus (SFV_{CPZ})¹³. All of these specimens, except for 28 samples from a group of habituated western gorillas (Makumba group) at the DS field site, were collected from non-habituated apes living in remote forest areas. Faecal samples were first subjected to host mitochondrial DNA analysis to determine their species and subspecies origin^{13–16,31}. A subset was then selected for host microsatellite analysis to determine the number of individuals at particular field sites (Supplementary Table 1). These included 198 chimpanzee samples from the GT field site, 189 eastern gorilla samples from the KE, LU and OP field sites, and 119 bonobo samples from the LK and KR field sites (Supplementary Table 2). For estimates of sample degradation (Supplementary Table 2) and oversampling (Supplementary Table 3), we also included microsatellite results that we had obtained earlier for specimens collected from central chimpanzees14-16 and western gorillas¹⁵ in Cameroon, as well as from eastern chimpanzees³¹ in the Democratic Republic of the Congo.

Microsatellite analyses. Faecal DNA was extracted¹⁴ and used to amplify four (GT) or eight polymorphic microsatellite loci (KE, LU, OP, LK and KR) as previously described^{14,15}. Amplification products were analysed on an automated sequencer (Applied Biosystems) and sized using GENEMAPPER 4.0 (Applied Biosystems). For individual identification, samples were first grouped by field site and mitochondrial DNA haplotype. Within each haplotype, samples were then grouped by microsatellite genotypes, but allowing for one allelic mismatch between samples. Specimens were classified as degraded if they failed to amplify two or more (GT) or three or more (all other sites) microsatellite loci. Samples with evidence of DNA admixture (multiple peaks for the same locus) were discarded. Amplification of ape Plasmodium sequences by conventional (bulk) PCR. Faecal samples were first screened for Plasmodium cytb sequences (956 bp) as previously described^{10,27}, but using modified PCR conditions as well as a different secondround reverse primer to generate an amplicon 166 bp longer. Nested PCR was performed using DW2 (5'-TAATGCCTAGACGTATTCCTGATTATCCAG-3') and DW4 (5'-TGTTTGCTTGGGAGCTGTAATCATAATGTG-3') in the first round of PCR and Pfcytb1 (5'-CTCTATTAATTTAGTTAAAGCACA-3') and PLAS2a (5'-GTGGTAATTGACATCCWATCC-3') in the second round. For the first round, 2.5 µl faecal DNA was used in a 25-µl reaction volume, containing 0.5 µl dNTPs (10 mM of each dNTP), 20 pmol of each primer (DW2 and DW4), 2.5 µl PCR buffer and 0.25 µl Expand Long Template enzyme mix (Roche). Cycling conditions included an initial denaturation step of 2 min at 94 °C, followed by 15 cycles of denaturation (94 °C, 10 s), annealing (45 °C, 30 s) and elongation (68 °C, 2 min), followed by 35 cycles of denaturation (94 $^\circ\text{C},$ 10 s), annealing (48 $^\circ\text{C},$ 30 s) and elongation (68 $^{\circ}\text{C},$ 2 min; with 15-s increments for each successive cycle), followed by a final elongation step of 10 min at 68 °C. For the second round of PCR, 1 µl of the first-round product was used in a 25-µl reaction volume containing 0.5 µl dNTPs (10 mM of each dNTP), 20 pmol of each primer (Pfcytb1 and PLAS2a), 2.5 µl PCR buffer and 0.25 µl Expand Long Template enzyme mix. Cycling conditions included an initial denaturation step of 2 min at 94 °C, followed by 60 cycles of denaturation (94 °C, 10 s), annealing (52 °C, 30 s) and elongation (68 °C 1 min), followed by a final elongation step of 10 min at 68 °C. Amplified products were gel-purified and sequenced directly to confirm Plasmodium infection.

Samples positive for Plasmodium cytb sequences were then screened for apicoplast clpC (390 bp) and nuclear ldh (772 bp) sequences. Amplification of the clpC fragment was performed as previously described9, but using modified PCR conditions and a different second-round reverse primer to generate an amplicon 117 bp longer. Nested PCR was performed using primers TFM1421+ (5'-AAAAC TGAATTAGCAAAAATATTA-3') and TFM1423RC (5'-CGAGCTCCATATAA AGGAT-3') in the first round of PCR and CLPCF1 (5'-TCTAAACAATTATTTG GTTCTG-3') and CLPCR2 (5'-GTTAATCTATTTARTAATTCHGGTTTAA-3') in the second round. Amplification of the *ldh* fragment was also performed as previously described^{28,32,33}, but using different PCR conditions. Nested PCR was performed using primers JNB272 (5'-ATGGCACCAAAAGCAAAAAT-3') and JNB273 (5'-GCCTTCATTCTSYTAGTTTCAGC-3') for the first round and LDH1 (5'-GGNTCDGGHATGATHGGAGG-3') and Fv2n (5'-AACRASAGG WGTACCACC-3') for the second round. PCR conditions were the same as described for the cytb fragment. Amplified products were gel-purified and sequenced directly to confirm Plasmodium infection.

Finally, *cytb*-positive samples were subjected to nested PCR with the aim of amplifying larger fragments (3.4 kb and 3.3 kb in length, respectively), which together spanned the entire *Plasmodium* mitochondrial genome (Supplementary Fig. 1a). The 3.4-kb fragment was amplified using Pf936p (5'-GAGAAAAA

TGYAATCCWGTWACACAATA-3') and DW4 in the first round of PCR and Pf1031p (5'-GATGCAAAACATTRWCCTAATAAGTA-3') and PLAS2a in the second round. The 3.3-kb fragment was amplified using McytP (5'-TATCCAA ATCTATTAAGTCTTG-3') and Pf1916n (5'-GCGTTCGTTCTTATAGTGTAG GC-3') in the first round of PCR and Pf4450p (5'-CTGTTCCTATTATGGTTT ATGTGTGC-3') and Pf1880n (5'-CCTTTAATGTAGTTTCCTCACAGCTT-3') in the second round. For the first round of amplification, 2.5 µl faecal DNA was used in a 25-µl reaction volume containing 0.5 µl dNTPs (10 mM of each dNTP), 20 pmol of each first-round primer, 2.5 µl PCR buffer and 0.25 µl Expand Long Template enzyme mix (Roche). Cycling conditions included an initial denaturation step of 2 min at 94 °C, followed by 15 cycles of denaturation (94 °C, 10 s), annealing (45 °C, 30 s) and elongation (68 °C, 4 min), followed by 35 cycles of denaturation (94 °C, 10 s), annealing (48 °C, 30 s) and elongation (68 °C, 4 min; with 15-s increments for each successive cycle), followed by a final elongation step of 10 min at 68 $^\circ \rm C.$ For the second round of amplification, 2 µl of the first-round PCR product was used in a 50 µl volume containing 1 µl dNTPs (10 mM of each dNTP), 40 pmol of each second-round primer, 5 µl PCR buffer and 0.5 µl Expand Long Template enzyme mix. Cycling conditions included an initial denaturation step of 2 min at 94 °C, followed by 60 cycles of denaturation (94 °C, 10 s), annealing (52 °C, 30 s) and elongation (68 °C, 4 min), followed by a final elongation step of 10 min at 68 °C. Amplified products were gel-purified, but only a small fragment was sequenced to confirm Plasmodium infection.

SGA of *Plasmodium* sequences from faecal DNA. To derive sequences suitable for phylogenetic analyses, a subset of bulk-PCR-positive chimpanzee (n = 80) and gorilla (n = 55) faecal samples was subjected to SGA analyses^{21,22}. Following a Poisson distribution, the DNA dilution that yields PCR products in no more than 30% of wells contains one amplifiable template per positive PCR more than 80% of the time. Faecal DNA was thus end-point diluted in 96-well plates, and the dilution for which fewer than 30% of the wells were positive was used to generate between one and 40 different SGA sequences per sample (Supplementary Table 5). The same primers and PCR conditions used for bulk amplification of *cytb*, mtDNA-3.4 kb, mtDNA-3.3 kb, *clpC* and *ldh* fragments were also used for cSGA analyses. Amplification products were gel-purified, and sequenced directly using SEQUENCHER 4.9 (Gene Codes Corporation). Sequences that contained double peaks as an indicator of more than one amplified template were discarded.

Sensitivity and specificity of Plasmodium nucleic-acid detection in faecal samples. To estimate the sensitivity of the diagnostic cytb PCR test, we determined the proportion of PCR-positive specimens from Plasmodium-infected apes that were sampled more than once on the same day. Other replicate samples were excluded because the duration of natural ape Plasmodium infections is unknown. The sensitivity of Plasmodium nucleic-acid detection was then calculated as the fraction of positive tests per total number of samples tested (Supplementary Table 4). Including data from 32 such apes, we estimated the sensitivity to be 57% (with confidence limits determined assuming binomial sampling). It should be noted that this approach led to a systematic (albeit small) overestimation of the assay sensitivity, because it did not account for infected apes that yielded only negative replicate samples. Moreover, Plasmodium detection in faecal samples is very probably less sensitive than in blood, as is the case in urine and saliva¹⁷. Thus, the prevalence rates in Table 1 and Supplementary Table 1 should be interpreted as minimum estimates of Plasmodium infection rates in wild apes. The specificity of faecal Plasmodium detection was 1.00, because all amplification products were sequence-confirmed.

Ape Plasmodium prevalence estimations. For sites where the number of sampled chimpanzees was known (Supplementary Table 1), Plasmodium prevalence rates were estimated on the basis of the proportion of infected individuals. For each ape, the probability that it would be detected as being infected, if it was truly infected, was calculated taking into consideration the sensitivity of the diagnostic PCR test and the number of samples analysed, with 95% confidence limits determined assuming binomial sampling. For the remaining field sites, where the number of sampled individuals was not known, prevalence rates were estimated on the basis of the number of faecal samples, but correcting for specimen degradation and oversampling. As shown in Supplementary Table 2, microsatellite analysis of 1,027 faecal samples indicated an average degradation factor of 13%. Microsatellite analyses also provided a quantitative estimate of oversampling. Because of regional differences in sample collection, oversampling values were calculated separately for the different ape species and subspecies. As shown in Supplementary Table 3, central chimpanzees, western gorillas, eastern chimpanzees, eastern gorillas and bonobos were each assumed to have been sampled on average 1.77, 1.84, 3.74, 2.01 and 1.84 times, respectively. Using these corrections, the proportion of Plasmodium-infected chimpanzees was estimated for each field site, again taking into account the sensitivity of the diagnostic test. From these determinations, prevalence rates and their confidence limits were calculated.

Phylogenetic analyses. Ape-derived Plasmodium sequences were aligned with human and simian reference sequences using CLUSTAL W³⁴. Sites that could not be aligned unambiguously were excluded. Trees were constructed from mitochondrial cytb sequences (956 bp, Supplementary Figs 2 and 3; 240 bp, Supplementary Fig. 8), apicoplast *clpC* sequences (390 bp, Supplementary Fig. 4), nuclear ldh sequences (772 bp, Supplementary Fig. 5) and mitochondrial halfgenomes (3,361 bp, Supplementary Fig. 6; 3,277 bp, Supplementary Fig. 7). In addition, trees were constructed from mitochondrial coding (Fig. 3a) and noncoding regions (Fig. 3b). Deduced COXI, COXIII, and CYTB protein sequences were concatenated into a single 980-amino-acid sequence. The non-proteincoding portion of the mtDNA-3.3kb fragment comprised 2,447 nucleotides following the removal of ambiguous sites. Phylogenetic trees were inferred using PHYML³⁰. The class of evolutionary model was chosen using MODELTEST³⁵, and parameters were iteratively estimated in PHYML, using the GTR+I+G model for nucleotide sequence trees and the LG+I+G model (ref. 36) for amino-acid sequence trees. Bootstrap values were calculated with 100 replicates³⁷. Posterior probability values were calculated with MRBAYES²⁹, using an average standard deviation of partition frequencies <0.01 as a convergence diagnostic. A neighbour-joining phylogenetic tree (Supplementary Fig. 8) was calculated with

CLUSTAL W, using the Kimura two-parameter model of evolution with bootstrap support based on 1,000 bootstrap replicates^{34,37}.

Nucleotide sequence accession numbers. All new SGA-derived ape *Plasmodium* sequences have been submitted to GenBank, with accession numbers listed in Supplementary Table 6.

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