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Accessibility
A direct characterization of human mutation based on microsatellites

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Abstract

Mutations are the raw material of evolution, but have been difficult to study directly. We report the largest study of new mutations to date: 2,058 germline changes discovered by analyzing 85,289 Icelanders at 2,477 microsatellites. The paternal-to-maternal mutation rate ratio is 3.3, and the rate in fathers doubles from age 20 to 58 whereas there is no association with age in mothers. Longer microsatellite alleles are more mutagenic and tend to decrease in length, whereas the opposite is seen for shorter alleles. We use these empirical observations to build a model that we apply to individuals for whom we have both genome sequence and microsatellite data, allowing us to estimate key parameters of evolution without calibration to the fossil record. We infer that the sequence mutation rate is 1.4–2.3×10−8 per base pair per generation (90% credible interval), and that human-chimpanzee speciation occurred 3.7–6.6 million years ago.
Here we focus on microsatellites: 1–6 base pair motifs that vary in the number of times they repeat. Due to DNA polymerase slippage during replication, the mutation rate of microsatellites is around $10^{-4}$ to $10^{-3}$ per locus per generation\textsuperscript{8–12}, far higher than the nucleotide substitution rate of $10^{-8}$. We analyzed 2,477 autosomal microsatellites that had been genotyped as part of linkage-based disease gene mapping studies and were ascertained to be highly polymorphic\textsuperscript{13}. The data set included 85,289 Icelanders from 24,832 father-mother-child trios, after restricting to individuals genotyped for at least half of these loci and without evidence of inaccurate parental assignment (Online Methods, Supplementary Figure 1). The median genotype error rate was $1.8\times10^{-3}$ per allele (Supplementary Figure 2, Supplementary Note), high compared to the mutation rate, and thus we took additional steps to reduce the error.

To distinguish genuine mutations from genotype errors, we used two approaches (Online Methods, Supplementary Note). In the ‘trio’ approach (Figure 1A), we identified 1,695 mutations in 5,085,672 transmissions by restricting to instances in which each member of the trio was genotyped more than once. In the ‘family’ approach (Figure 1B), we identified 363 mutations in 952,632 transmissions, validating new mutations by requiring them to be seen in at least one of the proband’s children, and validating ancestral alleles by requiring them to be seen in all of the proband’s siblings (in the family approach, we also used haplotypes of nearby microsatellites to determine the parental origin of mutations; Online Methods, Supplementary Note). The trio and family approaches produced indistinguishable inferences about the mutation process (Table 1, Supplementary Figure 3, Supplementary Figure 4), and hence we combined the data for subsequent analysis (62 mutations were counted twice due to overlap).

To estimate the proportion of candidate mutations that are real, we re-genotyped samples of 103 trio mutations and 99 family mutations leading to false-positive rate estimates of 2.9% and 2.6% respectively (Supplementary Table 1, Supplementary Figure 5). We also estimated the false-positive rate due to errors in the allele-calling algorithm to be 4.3% by manually re-scoring the electropherograms of 316 individuals from the family dataset, and declaring a false-positive if there was disagreement. Combining the two modes, we estimate a 7.2% false-positive rate (Supplementary Table 1). We also obtained an entirely independent estimate by analyzing next-generation sequencing data from the proband and the transmitting parent for 14 candidate mutations for which we had such data, allowing us to validate all but one leading to an estimated false positive rate of 7.1% (Supplementary Table 2). The false-negative rate (probability of an undetected real mutation) was estimated to be 9.0% by simulating mutations and recording the fraction that we failed to detect (Online Methods).

The estimated mutation rate of tetra-nucleotides is $10.01\times10^{-4}$ per locus per generation, 3.7 times higher than the di-nucleotide rate of $2.73\times10^{-4}$ (Table 1). Estimates are nearly unchanged after correcting for false-positives and false-negatives by $(1-0.072)/(1-0.090)$, and thus we quote unadjusted rates in what follows. Our estimate of the male-to-female mutation rate ratio is $\alpha=3.3$ (95% CI 2.9–3.7) (Supplementary Table 3), within the range of 2–7 previously inferred for sequence substitutions\textsuperscript{4,14,3,15}. Paternal age is correlated with mutation rate ($P=9.3\times10^{-5}$), whereas maternal age is not ($P=0.47$; Figure 2A, Supplementary Figure 6), consistent with observations based on disease-causing mutations, and the fact that male germ cells undergo numerous mitoses as a man ages, whereas female oocytes do not undergo postnatal cell division\textsuperscript{4}.

These data allow the first high resolution direct characterization of the mutation process for the highly polymorphic di- and tetra-nucleotide microsatellites that are typically genotyped\textsuperscript{8}. First, 32% of mutations at di-nucleotide microsatellites are multi-step, compared to 1% in...
tetra-nucleotides (Figure 2B, Supplementary Figure 3) (this explains why the variance of allele length distribution at the tetra-nucleotides is similar to that of di-nucleotides despite their 3.7-fold higher mutation rate\textsuperscript{16,17}). Second, mutation rate increases with allele length\textsuperscript{18,19}, quadrupling between 30–70 bp for di-nucleotides and 40–120 bp for tetra-nucleotides (both tests significant at P<0.002) (Figure 2C). Third, loci with uniform repeat structures (e.g. CACACACA) have a 40\% higher rate (P=3×10\textsuperscript{-7}) than compound repeat structures (e.g. CACATCACA), consistent with less DNA polymerase slippage for interrupted tandem repeats\textsuperscript{8,20} (Supplementary Figure 7). Fourth, we detect length constraints\textsuperscript{21,22}, with shorter alleles tending to mutate to become longer and vice versa (P=2×10\textsuperscript{-15}) (Figure 2D, Supplementary Figure 8, Supplementary Figure 9\textsuperscript{20,23,24}). This pattern contrasts with tri-nucleotide repeat disorders, where long alleles tend to get even longer\textsuperscript{19}. Fifth, mutation rate correlates (P<10\textsuperscript{-4}) with motif length, repeat number, allele-size, distance from exons, gender, and age, but not with recombination rate, distance from telomeres, human-chimpanzee divergence, or parental heterozygosity (Supplementary Table 4; Supplementary Table 5; Supplementary Note).

Microsatellites have been widely used for making inferences about evolutionary history. However, the accuracy of these inferences has been limited by a poor understanding of the mutation process. We developed a new model of microsatellite evolution (Supplementary Note). This model can estimate the time to the most recent common ancestor (TMRCA) between two samples at a microsatellite by taking into account: (1) the dependence of mutation rate on allele length and parental age (Figure 2A,C); (2) the step-size of mutations (Figure 2B); (3) the size constraints on allele length (Figure 2D, Supplementary Figure 8, Supplementary Figure 9); and (4) the variation in generation interval over history. In contrast to the Generalized Stepwise Mutation Model (GSMM), which predicts a linear increase of average squared distance (ASD) between microsatellite alleles over time, the new model predicts a sub-linear increase (Figure 3) and saturation of the molecular clock, due to the constraints on allele lengths. We also extended the model to estimate the sequence mutation rate, using the per-nucleotide diversity flanking each microsatellite as an additional datum. To implement the model, we used a Bayesian hierarchical approach, first generating global parameters common to all loci, followed by locus-specific parameters, and finally the microsatellite alleles at each locus (Online Methods). We used Markov Chain Monte Carlo to infer TMRCA and sequence mutation rate.

We validated the model in three ways (Online Methods). First, we simulated datasets in which we know the true sequence mutation rate and TMRCA, and found that our model is unbiased in estimating sequence mutation rate while producing accurate estimates of the standard error (Supplementary Note). Second, we carried out sensitivity analyses by perturbing model parameters and found that our key inferences are robust (Supplementary Note, Supplementary Figure 10). Third, we empirically validated the model by analyzing 23 individuals for whom we had both microsatellite genotypes and whole genome sequence (WGS) data\textsuperscript{2}, and comparing the ASD to the surrounding sequence heterozygosity as a surrogate for TMRCA. The ASD predicted by our model is similar to the empirical curve (Figure 3, Supplementary Figure 11)

Our approach allows inference of evolutionary parameters without calibration to the fossil record. Using the empirical ASD at the di-nucleotide microsatellites in each of the 23 individuals of European, East Asian and sub-Saharan African ancestry for whom we had WGS sequence data (Online Methods), and comparing the ASD to local heterozygosity and human-macaque divergence (as a surrogate for the local mutation rate; Online Methods), we inferred a sequence mutation rate and the TMRCA averaged across the genome (Table 2). We also inferred a 90\% credible interval (CI) based on a Bayesian approach integrating over uncertainty in the model parameters (Online Methods; Supplementary Note, Supplementary Note).
Table 6). Empirically, mutation rates tend to be more similar within than between populations (Supplementary Table 7; Supplementary Figure 12). The differences across populations are not likely to be due to poor modeling of demographic history, as when we model more realistic histories involving two bottlenecks in non-Africans we obtain the same results (Supplementary Figure 13). The mutation rate differences between populations may be due to shared history, but they are not significant, so we pooled our data across the 23 individuals to produce a sequence mutation rate estimate of $1.82 \times 10^{-8}$ per bp per generation (90% CI 1.40–2.28 $\times 10^{-8}$/bp/generation, Table 2) (the confidence interval takes into account correlations in the 23 peoples’ histories through a jackknife) (Online Methods).

Our inference of the sequence mutation rate is consistent with Nachman and Crowell’s estimate of $\mu_{seq} = 1.3–2.7 \times 10^{-8}$/bp/generation based on calibration to the fossil record6. It is also consistent with Kondrashov’s direct estimate of $\mu_{seq} = 1.8 \times 10^{-8}$/bp/generation from studies of disease causing genes. However, the lower bound of our 90% CI is higher than two recent studies based on whole-genome sequencing (WGS): $\mu_{seq} = 1.1 \times 10^{-8}$/bp/generation based on 28 sequence mutations detected in a four member family1, and $\mu_{seq} = 1.0 \times 10^{-8}$ and $1.2 \times 10^{-8}$/bp/generation based on 84 sequence mutations detected in two trios2,3. We considered the possibility that this discrepancy might be due to ascertainment bias, because the microsatellites we analyzed were selected to be highly polymorphic (for disease gene mapping) which could cause a too-high ASD. However, this would overestimate TMRCA at the loci we analyzed and thus underestimate the mutation rate, opposite to what would be necessary to explain the discrepancy (Supplementary Figure 12 and Supplementary Note). We hypothesize that the lower mutation rate estimates from the WGS studies might be due to: (i) the limited number of mutations detected in the WGS studies which explains why their confidence intervals overlap ours, (ii) possible underestimation of the false-negative rate in the WGS studies, or (iii) variability in the mutation rate across individuals so that a few families cannot provide a reliable estimate of the population-wide rate. There is already empirical evidence for variability in the mutation process across individuals: in one trio analyzed in the 1000 Genomes Project study8, the father transmitted 92% of the mutations, while in the other trio the father transmitted 36%. Studies of sequence substitution in many families are important, as they will make it possible to measure population-wide rates and study features of the sequence substitution process not accessible to microsatellite analysis.

Our direct estimation of the microsatellite mutation rate, combined with comparative genomics data, also allows us to estimate the date of human-chimpanzee speciation $\tau_{HC}$, which we define as the date of last gene flow between human and chimpanzee ancestors26,27. We estimate a genome-wide average human-chimpanzee genetic divergence time $t_{HC} = 5.80–9.77$ Mya28 (Online Methods; Table 2). By definition, this must be older than the speciation date $\tau_{HC}$. We then inferred the human-chimpanzee speciation to be $\tau_{HC} = 3.75–6.57$ Mya by integrating our posterior distribution on $t_{HC}$ with a prior distribution on $\tau_{HC}$ of 0.663±0.041 whose mean we obtained by modeling-based estimates of $\tau_{HC}/t_{HC} = 0.61–0.68$29,30, and whose 95th percentile upper bound of 0.73 we obtained by analyzing human-chimpanzee sequence data in regions with a reduced divergence compared to the autosomal average due to being (1) on chromosome X, (2) in proximity to genes, and (3) near divergent sites that cluster humans and chimpanzees to the exclusion of gorilla (Supplementary Note). Our upper bound of $\tau_{HC} < 6.57$ Mya is lower than the 6.8–7.2 Mya31 date estimate for Sahelanthropus tchadensis, a fossil interpreted as being on the human lineage after the final separation of human and chimpanzee ancestors32 because it shares derived features with other hominins such as bipedal posture, reduced canines and expanded post-canines with thicker enamel33. In the Supplementary Note, we also obtain an independent upper bound on the human-chimpanzee speciation date of $\tau_{HC} < 6.3$ Mya based on calibration to the fossil record of human-orangutan speciation26. A possible explanation
for the discrepancy between the genetic and fossil dates is that *Sahelanthropus* was not a hominin, but instead shared independently-derived similarities (homoplasies)\(^{34}\). Alternatively, populations with hominin traits may have continued to exchange genes with *chimpanzee* ancestors after *Sahelanthropus*\(^{26}\). Finally, the age of *Sahelanthropus*\(^{31}\) may be overestimated.

**URL:** Complete Genomics data: [http://www.completegenomics.com/](http://www.completegenomics.com/).

### Online Methods

#### Data sets

Microsatellite genotypes were obtained at deCODE Genetics using DNA extracted from blood, and multiplexed capillary gel electrophoresis with automated allele calling\(^{13}\). We restricted to 2,477 autosomal loci that were genotyped most heavily (all had a minimum repeat length of 5 units). We analyzed 85,289 individuals genotyped for at least half of these loci, from which we identified 25,067 mother-father-offspring trios using the deCODE Genetics genealogical database (Íslendingabók).

To filter out trios with inaccurate parental assignments, we computed the fraction of loci where both alleles differ between a parent and a child. We empirically set the threshold to filter out almost all known uncle-proband and aunt-proband pairs while retaining almost all known parent-proband pairs (Supplementary Figure 1).

To estimate the per-locus genotyping error rate, we used discordance rates in cases of repeated genotypes (Supplementary Note).

Deep whole genome sequencing data was obtained from two sources. We downloaded 9 sequences generated using Illumina technology, mapping reads using BWA\(^{35}\) and calling SNPs with SAMtools\(^{36}\). We also downloaded 20 Complete Genomics sequences, 6 of which overlap the Illumina sequences (Supplementary Table 7, Supplementary Figure 14). To estimate heterozygosity around each microsatellite, we extracted a window of data centered around it (in most cases, 0.001 centimorgans masking out the central 1kb\(^{37}\); Supplementary Figure 15),

#### Detecting mutations

For the trio approach, we restricted to transmissions in which all members of the trio were genotyped at least twice and searched for Mendelian inheritance incompatibilities. There were some detected mutations in which the parental origin was ambiguous (Supplementary Note), and we included these for analyses of the mutation rate but not for analyses requiring parental origin (Fig 2B). For mutations with unambiguous parental origin, the ancestral allele was defined as the one that was closer in length to the mutant allele (we randomly chose the ancestral allele if both were equally close). We filtered out 49 loci that harbored many more mutations from homozygous parents to homozygous children than expected based on Hardy-Weinberg equilibrium, a phenomenon that affected the trio, but not the family data. We determined that this is a real error mode due to polymorphisms under the PCR primer sites\(^{38–40}\) by sequencing primer sites from 15 mutations and identifying 5 with SNPs in the primer region (Supplementary Note).

For the family approach, we restricted to transmissions where genotyping was available not just for a proband’s two parents, but also for at least one child and one sibling (Figure 1B). We identified putative mutations by searching for Mendelian incompatibilities between the proband and their parents. We used Allegro 2.0\(^{41}\) to phase the family masking out the mutant locus, using all available loci from the same chromosome. We then assigned the
haplotype carrying the mutation to one of the proband’s parents (Supplementary Note). To validate the mutation, we required at least one sibling to carry the haplotype with the ancestral allele, no sibling to carry the mutant, and at least one child to carry the haplotype with the mutant.

The trio and family approaches provide complementary information. A bias that only affects the trio approach is somatic mutations in the lineage of genotyped cells but not germline cells transmitted to offspring (this is minimized since the DNA we analyzed was extracted from blood, but is still a concern). A bias that only affects the family-based approach is that mutations in progenitor germ cells might cause a mutation to be observed simultaneously in the proband and its siblings, causing us to reject a real mutation. The fact that both approaches produce consistent inferences despite the difference biases increases our confidence in the results.

False-positive and false-negative rates

To estimate the false-positive rate, we re-genotyped candidate mutations. In the trio dataset, we randomly re-genotyped 103 mutations. In the family dataset, we targeted mutations that had a higher a priori chance of being in error (Supplementary Table 1). To provide an entirely independent estimate of the false-positive rate, we identified 14 candidate mutations where we had at least 7-fold whole genome sequencing data from the proband as well as (at least) the transmitting parent. We then manually examined the data, failing to validate only 1 of the 14 mutations (Supplementary Table 2).

To estimate the false-negative rate (the proportion of genuine mutations that were missed), we randomly distributed mutations on the genealogy and then tested whether they gave rise to detectable inheritance errors. As an example, suppose that the father-mother-proband trio has genotypes of allele-lengths (6, 10), (8, 10), (8, 10), respectively. If the mother passed allele 10 to the proband, and the father passed a 6 → 8 mutation, then this mutation would not be detected.

Statistical characterization of the microsatellite mutation process

To infer the standard error of the mutation rate, taking into account rate variation across loci, we used a hierarchical Bayesian model (Supplementary Note). To infer the number of microsatellite repeats, we started with the amplicon size, which includes not just the repeats but all the sequence between the PCR primers. We then subtracted the span of the flanking sequence inferred from the human genome reference (Supplementary Figure 16). To compute the relative length of an allele, we measured the mean and standard deviation over all individuals at that locus, and report the standard deviations from the mean (Z-score). To estimate motif impurity (Supplementary Figure 7), we applied the Tandem Repeat Finder software to the human genome reference (Supplementary Figure 16). To test for association between the microsatellite mutation process and genomic features (Supplementary Table 4), we performed a logistic regression to mutation rate and directionality, and a Poisson regression to step size. To test for interaction, we performed multivariate logistic regression (Supplementary Table 5).

Prior distributions on evolutionary parameters

For our Bayesian modeling of sequence mutation rate and genetic divergence times, we required prior distributions on evolutionary parameters (Supplementary Table 6):

- *Generation interval*: Based on interviews with experts on chimpanzee and gorilla demographic structure (L. Vigilant and K. Langergraber), we assume that the ancestral generation time was 22.5 ± 4.2 (mean ± SD) years. Based on the
literature\(^\text{42,43}\) (Supplementary Figure 17), we assume that present-day generation time is 29±2 years. We also assume that the difference between the male and female generation time was 0.5±3.3 years in the ancestral population and 6.0±2.0 today (Supplementary Note). We sample the transition between ancestral to present-day generation time to be a mixture of 3 equally weighted exponential distributions, with means of 50Kya, 200Kya, and 2Mya, corresponding to hypothetical changes around the Upper Paleolithic revolution, evolution of modern humans, and evolution of *Homo erectus*.

- **Human-ape relative genetic divergences**: From the literature, we assume that the ratio of human-chimpanzee to European-European genetic divergence per base pair is\(^\text{28}\) 15.400±0.356, and that the ratio of human-orangutan to human-chimpanzee genetic divergence of 2.650±0.075\(^\text{26,29}\). We assume that the molecular process of mutation has been constant over great ape history.

- **Human-chimpanzee speciation time (\(\tau_{HC}\))**: Human-chimpanzee speciation time is by definition less than human-chimpanzee genetic divergence time. Our prior on \(\tau_{HC}/t_{HC}\) has a normal distribution with mean 0.663, within the range of 0.61–0.68 from model-based analyses\(^\text{29,30}\). We use a standard deviation of 0.041 based on an analysis in the Supplementary Note suggesting showing that an upper bound on the ratio of human-chimpanzee speciation to genetic divergence is 0.73 (our prior distribution is set that 95% of its density is below this).

### A model of microsatellite evolution assisted by flanking sequence heterozygosity

As a metric of microsatellite allelic divergence between two samples, we use the Average Squared Distance (ASD). Given allele lengths \(x_1, x_2, \ldots, x_n\),

\[
ASD = \frac{1}{n(n-1)} \sum_{i,j} (x_i-x_j)^2.
\]

To model ASD along with flanking sequence heterozygosity, we simulate the evolution of a pair of chromosomes from a common ancestor, over multiple loci and individuals. The model is hierarchical: At the top level, global parameters (Supplementary Table 6) common to all loci are simulated, such as the genome-wide present-day sequence and microsatellite mutation rates, and generation-time effects. One level down, locus-specific mutation rates are computed based on global parameters and locus-specific information. At the third level, for each individual, a two-sample coalescent tree is generated (Supplementary Note).

A potential pitfall in inferring TMRCA with our data is that the microsatellites we analyzed were ascertainment to be highly polymorphic in Europeans. This raises two complications. First, the sequence flanking the microsatellites may have a different mutation rate than the genome average, and to correct for this, we compared ASD to the ratio of sequence heterozygosity and human-macaque divergence at each locus (as a surrogate for local mutation rate). Second, ascertainment of highly polymorphic microsatellites can bias toward deeper genealogies than the genome average which in turn can bias average TMRCA to be too high. By studying the sequence flanking the microsatellites, we determined that the trees were on average 1.04 times deeper than the genome average, and we corrected Table 2’s estimate of genome-wide average TMRCA by this factor (Supplementary Note, Supplementary Table 8).

To infer sequence mutation rate and TMRCA using the microsatellite evolution model, we use a Markov Chain Monte Carlo (MCMC) following the method of ref. 44 (Supplementary Note). Combining data across individuals is not trivial because of shared history across individuals. To obtain proper standard errors for the combined mutation rate, we performed
a jackknife\(^5\), where each locus is removed at a time. This gives the final set of standard errors.

**Editorial Summary (AOP and Month, same)**

David Reich and colleagues report a direct characterization of the human mutation rate based on analysis of 85,289 Icelandic individuals genotyped at 2,477 autosomal microsatellite loci. They use this to build a model of microsatellite evolution and estimate key evolutionary parameters.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Examples of verified mutations from a trio and a family
The proband is the individual inheriting a mutation, and all individuals are named relative to the proband. All alleles are given in repeat units and shifted so that the ancestral allele has length 0. The mutating allele is underlined. (A) We show a mutation detected using the trio approach. Confirmation of the mutation is from multiple genotyping of the trio: the father, mother, and proband are genotyped 3×, 3×, and 4×, respectively. (B) We show a mutation detected using the family approach. One sibling verified the ancestral allele, and one child verified the mutant allele. The phasing of alleles from the mutant locus and other loci from the same chromosome shows that the sibling with alleles (0,-2) did not inherit the ancestral ‘0’ but rather the other ‘0’ allele from the father.
Figure 2. Characteristics of the microsatellite mutation process

(A) Paternal (blue) and maternal (red) mutation rates. The x-axis shows the parental age at child-birth. The data points are grouped into 10 bins (vertical bars show 1 standard error). The paternal rate shows a positive correlation with age (logistic regression of raw data: P=9.3×10^{-5}; slope = 1.1×10^{-5}/yr), with an estimated doubling of rate from age 20 to 58. The maternal rate shows no evidence of increasing with age (P=0.47). (B) Mutation length distributions differ between di- and tetra-nucleotides (upper and lower histograms), with the x-axis in units of step-size. While the di-nucleotide loci experience multi-step mutations in 32% of instances, tetra-nucleotides mutate almost exclusively by a single-step of 4 bases. (C) Mutation rate increases with allele length: di-nucleotides (blue) have a slope of 1.65×10^{-5} per repeat unit (P=1.3×10^{-5}) and tetra-nucleotides (red) have a slope of 6.73×10^{-5} per repeat unit (P=1.8×10^{-3}). (D) Constraints on allele lengths: When the parental allele is relatively short, mutations tend to increase in length, and when the parental allele is relatively long, mutations tend to decrease in length. Di- and tetra-nucleotides are shown in blue crosses and red circles, respectively. Probit regression of the combined di- and tetra-data shows highly significant evidence of an effect (P=2.8×10^{-18}).
Figure 3. Empirical validation of our model with sequence-based estimates of TMRCA

In red is the simulation of ASD as a function of TMRCA for the standard random walk (GSMM) model. In blue is the simulation of our model, in which the non-linearity compared to GSMM is primarily due to the length constraint that we empirically observed in microsatellites. In black is the empirically observed ASD at microsatellites in 23 HapMap individuals as a function of sequence-based estimates of TMRCA, which is estimated using $\frac{\theta_{seq}}{\mu_{seq}}$, where $\theta_{seq}$ is the local sequence diversity surrounding each microsatellite locus, and $\mu_{seq}$ is $1.82 \times 10^{-8}$ (obtained from Table 2). The close match of the empirical curve to our model simulations suggests that our model works, and motivates the analysis in which we use the sequence substitution rate in small windows around the microsatellites to make inferences about evolutionary parameters like the sequence mutation rate.
Figure 4. Human-chimpanzee speciation date inferred without a fossil calibration
In the square panel, we give the 90% Bayesian credible interval for human-chimpanzee speciation time (gray), for a range of values of the ratio of speciation time to divergence time $\tau_{HC}/t_{HC}$. The blue curve shows our prior probability distribution for $\tau_{HC}/t_{HC}$, justified in Supplementary Note. The red horizontal lines are the dates of fossils that are candidates for being on the hominin lineage post-dating the speciation of humans and chimpanzees. *Australopithecus amanensis, Orrorin tugenensis* and *Ardipithecus kadabba* are within our plausible speciation times, while *Sahelanthropus tchadensis* pre-dates the inferred speciation time for all plausible values of $\tau_{HC}/t_{HC}$. Our prior distribution for $\tau_{HC}/t_{HC}$ is shown in the bottom histogram, and our posterior distribution of human-chimpanzee speciation time is shown in the left histogram.
Table 1
Direct estimates of microsatellite mutation rates

|                      | Mutations | Transmissions | Mutation rate ($\times 10^{-4}$)* |
|----------------------|-----------|---------------|-----------------------------------|
|                      |           |               | mean 5th – 95th percentile         |
| di-nucleotide loci†  |           |               |                                   |
| Trio-approach        | 1,218     | 4,578,348     | 2.66                              |
| Family-approach      | 269       | 861,204       | 3.12                              |
| Combined             | 1,487     | 5,439,552     | 2.73                              |
| tetra-nucleotide loci|           |               |                                   |
| Trio-approach        | 380       | 393,072       | 9.67                              |
| Family-approach      | 86        | 72,516        | 11.86                             |
| Combined             | 466       | 465,588       | 10.01                             |

*The 90% credible interval is calculated based on a Bayesian hierarchical beta-binomial model (Supplementary Note), which allows for the mutation rate to vary across loci.

†The breakdown of the mutation rate by motif type for di-nucleotides is in Supplementary Table 9.
# Table 2

Estimates of mutation rates and human-ape divergence times

|                                      | Mean (units: per generation per site) | 5th – 95th percentile (units: per year per site) | mean (units: thousand generations ago) | 5th – 95th percentile (units: million years ago) |
|--------------------------------------|---------------------------------------|-------------------------------------------------|----------------------------------------|-------------------------------------------------|
| Present-day mutation rates           |                                       |                                                 |                                        |                                                 |
| di-nucleotide microsatellite rate (per locus) | $2.73 \times 10^{-4}$ | $2.56 – 2.91 \times 10^{-4}$ | $9.47 \times 10^{-6}$ | $8.29 – 10.82 \times 10^{-6}$ |
| $\mu_{\text{seq}}$ nucleotide substitution rate (per base) | $1.82 \times 10^{-8}$ | $1.40 – 2.28 \times 10^{-8}$ | $6.76 \times 10^{-10}$ | $5.11 – 8.41 \times 10^{-10}$ |
| Genetic divergence times             |                                       |                                                 |                                        |                                                 |
| $t_{\text{CEU}}$: Western Europeans  | 22.8                                  | $17.8 – 29.6$                                  | 0.546                                  | 0.426 – 0.709                                   |
| $t_{\text{YRI}}$: Yoruba (African)   | 30.2                                  | $23.6 – 39.2$                                  | 0.720                                  | 0.562 – 0.933                                   |
| $t_{\text{HC}}$: human-chimpanzee    | 352                                   | $272 – 459$                                    | 7.49                                  | 5.80 – 9.77                                    |
| $\tau_{\text{HO}}$: human-orangutan  | 932                                   | $717 – 1220$                                   | 19.8                                  | 15.2 – 25.9                                    |
| $\tau_{\text{HC}}$: human-chimpanzee speciation time | 233 | $176 – 309$ | 4.97 | 3.75 – 6.57 |

*90% Bayesian credible interval obtained from the Bayesian posterior distribution.