# Higher Levels of Neanderthal Ancestry in East Asians than in Europeans

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**ABSTRACT** Neanderthals were a group of archaic hominins that occupied most of Europe and parts of Western Asia from ~30,000 to 300,000 years ago (KYA). They coexisted with modern humans during part of this time. Previous genetic analyses that compared a draft sequence of the Neanderthal genome with genomes of several modern humans concluded that Neanderthals made a small (1–4%) contribution to the gene pools of all non-African populations. This observation was consistent with a single episode of admixture from Neanderthals into the ancestors of all non-Africans when the two groups coexisted in the Middle East 50–80 KYA. We examined the relationship between Neanderthals and modern humans in greater detail by applying two complementary methods to the published draft Neanderthal genome and an expanded set of high-coverage modern human genome sequences. We find that, consistent with the recent finding of Meyer *et al.* (2012), Neanderthals contributed more DNA to modern East Asians than to modern Europeans. Furthermore we find that the Maasai of East Africa have a small but significant fraction of Neanderthal DNA. Because our analysis is of several genomic samples from each modern human population considered, we are able to document the extent of variation in Neanderthal ancestry within and among populations. Our results combined with those previously published show that a more complex model of admixture between Neanderthals and modern humans is necessary to account for the different levels of Neanderthal ancestry among human populations. In particular, at least some Neanderthal–modern human admixture must postdate the separation of the ancestors of modern European and modern East Asian populations.

**N**EANDERTHALS were a group of archaic hominins that occupied large parts of Europe and West Asia from ~30,000 to 300,000 years ago (KYA) (Stringer and Hublin 1999; Hublin 2009). Their disappearance in the fossil record often coincides with the first appearance of anatomically modern humans (AMH) in that region (Finlayson 2004). Where, when, and how often Neanderthals interbred with expanding AMH populations is still an open question. Morphological studies have generally concluded that Neanderthals made little or no contribution to present-day human populations (Stringer and Andrews 1988; Lahr 1994), but others have suggested there was some admixture (Duarte

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*et al.* 1999; Trinkaus 2007). Initial comparisons of Neanderthal and modern human DNA found no evidence for a Neanderthal contribution to the modern human gene pool (Krings *et al.* 1997; Serre *et al.* 2004; Noonan *et al.* 2006). However, indirect studies of patterns of linkage disequilibrium (LD) in contemporary human populations have consistently found support for admixture between "archaic" human groups (such as Neanderthals) and modern humans (Garrigan *et al.* 2005a,b; Plagnol and Wall 2006; Wall *et al.* 2009; Hammer *et al.* 2011; Lachance *et al.* 2012).

A detailed analysis of a draft Neanderthal genome and five low-coverage (4×) human sequences estimated that Neanderthals made a 1–4% contribution to the gene pool of modern non-African populations (Green *et al.* 2010). The presence of "Neanderthal DNA" in East Asians and Melanesians was initially surprising because the archaeological record shows that Neanderthals and early modern humans coexisted only in Europe and western Asia. Green and colleagues hypothesized that Neanderthals and modern humans came into contact and interbred in the Middle East

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 ${\sim}50{-}80$  KYA, prior to the divergence of modern-day European and Asian populations.

Green *et al.* (2010) presented three kinds of evidence in favor of interbreeding. First, they found (using *D*-statistics, a new measure of genetic similarity introduced in that article) that the three sampled non-African genome sequences (from a French, a Han Chinese, and a Papua New Guinean) are more similar to the Neanderthal sequence than is either of the two sampled African sequences (from a San and a Yoruban). Second, they identified several haplotypes that are in low frequency in Europeans, absent from Africans, and present in the Neanderthal sequence, which suggests those haplotypes were derived from Neanderthals. Third, they found many more genomic fragments in a European genome than in an African genome that have low divergence to the Neanderthal genome.

Admixture between modern humans and Neanderthals within the past 100,000 years (Kyr) is only one possible explanation for these D-statistic patterns. Green et al. noted that another potential explanation is ancient population subdivision within Africa before both Neanderthals and modern humans left Africa (cf. Green et al. 2010, figure 6). If there had been long-lived (e.g., >500 Kyr) population structure within Africa, and both Neanderthals and non-African AMH came from the same "source" subpopulation, then Neanderthals would be more similar to non-Africans in the absence of any recent admixture between AMH and Neanderthals (see Figure 1A). This intuitive argument was confirmed by the simulation studies of Durand et al. (2011) and Eriksson and Manica (2012), but these studies did not account for the other two lines of evidence summarized above. Two other studies have shown that the ancient-subdivision model is incompatible with other aspects of the data. Yang et al. (2012) demonstrated that recent admixture (Figure 1B) could be distinguished from ancient subdivision (Figure 1A) by computing the frequency spectrum of modern humans, conditioned on the Neanderthal sequence having the derived allele and an African sequence having the ancestral allele. This double conditioning enriches for alleles introduced by recent admixture if it occurred. Yang and colleagues found that the doubly conditioned frequency spectrum in Europeans and in East Asians is consistent with recent admixture, not with ancient subdivision. Separately, an analysis of the extent of LD at closely linked sites also concluded that the data were consistent with recent admixture and not with ancient subdivision (Sankararaman et al. 2012).

In this study, we revisit the question of Neanderthal admixture, using an expanded data set of 42 high-coverage (>45×) modern human genomic sequences, and we take advantage of the recent high-coverage Denisova genome (Meyer *et al.* 2012) to obtain more refined estimates of admixture proportions. We use two complementary methods of analysis. One is the *D*-statistic method introduced by Green *et al.* (2010). *D*-statistics reflect site-by-site differences. Because we have multiple individuals from each of several populations, we can quantify the extent of variation in



**Figure 1** Simplified versions of models of ancient population structure (A) or recent admixture (B) that can explain the observed levels of divergence between modern human genomes and the draft Neanderthal genome. Here T1 is the time when Neanderthals and modern humans first split, T2 is the time when African and non-African modern human populations split, and T3 is the time when Neanderthals mixed with modern humans.

*D*-statistics among pairs of individuals from the same two populations and obtain greater statistical power by combining estimates among all pairs. The second method is an LD-based method similar to one introduced by Wall (2000) and Plagnol and Wall (2006) for identifying putatively introgressed regions in modern human genomes. We use the draft Neanderthal genome to identify segments in the modern human genome that were derived from admixture with Neanderthals. This method is similar to the one used by Green *et al.* (2010) but is less restrictive and allows quantification of the differences in the number of admixed segments in different populations.

Using both of these methods, we show there was more Neanderthal admixture into East Asian populations than into European populations. This conclusion is consistent with that of Meyer *et al.* (2012), which was based on the analysis of a smaller number of modern human sequences. By using the high-coverage Denisova genome, we are able to show that the admixture rate into East Asians is 40% higher than into Europeans. We conclude that admixture between Neanderthals and modern humans did not occur at a single time and place, as suggested by Green *et al.* (2010). Some of it had to have occurred after the separation of East Asians and Europeans. Further, we show that there was significant Neanderthal admixture into the Maasai population of East Africa, probably because of secondary contact with a non-African population rather than admixture directly from Neanderthals.

#### **Materials and Methods**

#### Complete genomics data

We downloaded data from 69 publicly available genome sequences from the Complete Genomics (CGI) website (http://www.completegenomics.com/public-data/). Complete Genomics sequenced a Yoruba (YRI) trio, a Centre d'Etude du Polymorphisme Humain (CEPH)/Utah (CEU) pedigree family of 17 family members, a Puerto Rican (PUR) trio, and a diversity panel from 10 different populations. Combining these data sets and using only nonrelated, nonadmixed individuals, we have a sample size of 42 individuals representing nine different populations (Table 1). In addition to 36 members of the diversity panel, we also used the parents from the YRI trio and the maternal and paternal grandparents in the CEU pedigree. The individual genomes were sequenced to a minimum 45-fold coverage (Drmanac et al. 2010). The eight populations are Utah residents with Northern and Western European ancestry from the CEPH collection (CEU); Han Chinese from Beijing, China (CHB); Gujarati Indians from Houston (GIH); Japanese from Tokyo (JPT); Luhya from Webuye, Kenya (LWK); Maasai from Kinyawa, Kenya (MKK); Toscani from Italy (TSI); and Yoruba from Ibadan, Nigeria (YRI). Samples from three other populations were also available from Complete Genomics, those of Mexican ancestry in Los Angeles (MXL), African-Americans from southwest Arizona (ASW), and Puerto Ricans from Puerto Rico (PUR), but these were excluded from our analysis because of recent intercontinental admixture. All genomic data were downloaded from Complete Genomics' ftp site (ftp://ftp2.completegenomics.com/). We used two separate pipelines for filtering and processing the data, optimized for the different analyses performed (see below).

# D-statistic filtering

For the *D*-statistic analyses, each individual genome was aligned with the human genome assembly hg19 for consistency with the available assembly of the Neanderthal genome. Since our results were somewhat unexpected, we prepared the data for analysis in two different ways to check for consistency. We denote these analysis A and analysis B.

For analysis A, we used the release of the file format version 2.0 (software version 2.0.0.26) that was generated

Table 1 Forty-two individual genome sequences from Complete Genomics included in our study

ID	Population	ID	Population
NA06985	CEU	NA21732	MKK
NA06994	CEU	NA21733	MKK
NA07357	CEU	NA21737	MKK
NA10851	CEU	NA21767	MKK
NA12004	CEU	NA18940	JPT
NA12889	CEU	NA18942	JPT
NA12890	CEU	NA18947	JPT
NA12891	CEU	NA18956	JPT
NA12892	CEU	NA20502	TSI
NA18526	СНВ	NA20509	TSI
NA18537	СНВ	NA20510	TSI
NA18555	СНВ	NA20511	TSI
NA18558	СНВ	NA18501	YRI
NA20845	GIH	NA18502	YRI
NA20846	GIH	NA18504	YRI
NA20847	GIH	NA18505	YRI
NA20850	GIH	NA18508	YRI
NA19017	LWK	NA18517	YRI
NA19020	LWK	NA19129	YRI
NA19025	LWK	NA19238	YRI
NA19026	LWK	NA19239	YRI

in September 2011. This version was mapped to the human reference genome hg19. We also downloaded the chimpanzee genome pantro2 aligned to hg19 from the University of California, Santa Cruz (UCSC) Genome Browser (http:// hgdownload.cse.ucsc.edu/goldenPath/hg18/vsPanTro2/). The Neanderthal sequence was obtained by pooling reads from the three Vindija bones (SLVi33.16, SLVi33.25, and SL Vi33.26) that were aligned to the reference human genome (Green et al. 2010). The Neanderthal data were downloaded from the UCSC genome browser (http://genome. ucsc.edu/Neandertal/). To match the filtering used in the original Green et al. (2010) study, we used only sites with a mapping quality score (MAPQ) of at least 90 and a sequence quality >40. On average, the coverage of the Neanderthal genome was  $\sim$ 1.3-fold. We kept only sites that had one, two, or three reads.

After filtering out any insertions, deletions, or ambiguously called sites in the Complete Genomics data, we merged them with the chimpanzee and Neanderthal genomes. We kept only sites that had no more than two alleles in any of the human genomes and at which alleles were called for each human, the chimp, and the Neanderthal. Furthermore, we considered only transversion differences.

We also obtained the high-coverage Denisova genome from Meyer *et al.* (2012). The genome was aligned to the human reference genome (hg19) and the average coverage was  $\sim$ 30x. We filtered out all sites that had <16 reads or >46 reads. We merged these data with the data from analysis A to compute the *D*-statistic and *f*-statistic.

For analysis B, we redownloaded the genomic data from the Complete Genomics website (ftp://ftp2.completegenomics. com/, software version 2.0.2.15, file format version 2.0,

February 2012). These sequences were aligned to hg18. We applied a less stringent filter of the Neanderthal data: the filtering for mapping quality and sequence quality remained the same as in analysis A, but there were no restrictions on the number of reads per site. Finally, instead of considering the chimp genome as the outgroup, we used the ancestral alleles defined by the 1000 Genomes Project from the Enredo-Pecan-Ortheus (EPO) pipeline (Paten *et al.* 2008a,b) (data downloaded from ftp://ftp.1000genomes.ebi.ac.uk/). We refer to this outgroup as the reconstructed common ancestor (RCA).

For samples from any two populations compared, we filtered out any insertions, deletions, or ambiguously called sites. These genomic samples were then merged with the Neanderthal genome and the RCA outgroup. This differs from analysis A, where all populations were merged with the Neanderthal, Denisova, and chimp genome prior to any comparisons between populations. We considered only sites where the difference between the ancestral allele from the RCA and the alternate allele is a transversion, as we did in analysis A.

#### LD-based analysis filters

Since the LD-based analyses primarily utilize patterns of extant genetic variation (and only secondarily use the draft Neanderthal genome), we aligned variant calls to the updated human genome assembly (hg19), included both transitions and transversions, and imposed more stringent filters to throw out repetitive regions. Specifically, a custom series of Perl/C scripts and cgatools v1.3.0.9 were used to get a common set of variants from each individual. Using the CGI's variant file, all polymorphic regions containing SNPs were identified and reconstructed according to CGI's descriptions. These regions were then filtered for SNPs in such a way that both alleles were known for a given individual and were not part of a complex variant (for example, a SNP on one haploid phase and a deletion on the other phase). We then pooled all unique SNP positions from the full panel of samples and removed all SNPs located within repeats and segmental duplications with a minimum size of 50 bp. Structural variants (dgv track on UCSC), self chain (identity <90%, UCSC self-chain track), segmental duplications (UCSC), microsatellites (UCSC), simple tandem repeats (UCSC), and repeat masked sequence (UCSC) were also excluded. The final list of SNPs was then used by CGI's "snpdiff" tool to extract each sample's base calls relative to the human reference genome (hg19, Build 37). The snpdiff output was then reformatted to ms, PLINK, and other text-based formats for further analyses.

Subsequently, we identified numerous regions where all/ most individuals had heterozygous SNP calls but only one homozygous genotype was present. These regions likely reflect either alignment errors due to the Complete Genomics short-read sequencing technology or errors in the human reference genome sequence. We excluded all regions that included sites where over half of the individuals are heterozygous and only one homozygous genotype is present. The coordinates for these regions are available from the authors upon request.

Denisova sequence reads (Reich et al. 2010), mapped to the human reference genome hg18, were downloaded from the UCSC genome browser (http://genome.ucsc.edu/ cgi-bin/hgTrackUi?db=hg18&c=chrX&g=bamSLDenisova). Consensus Neanderthal sequence generated from three bones and aligned to the human reference genome hg18 was downloaded from the Ensembl genome browser (http://neandertal. ensemblgenomes.org/data info.html). Samtools 0.1.18 (Li et al. 2009) was used to convert the BAM files into a pileup alignment (mpileup arguments: -B -q5 -Q30) of each ancient hominin genome and hg18 for the region of interest. To compare modern human sequence tracks to ancient hominin sequences, hg19 coordinates of interest were converted to hg18 coordinates using the UCSC genome browser tool liftOver and extracted from the pileup alignments via custom perl scripts. To further compare the human sequences to sequences of other primate genomes, another custom perl script was used to extract the same hg19 coordinates of interest from a subset of the genomes in the UCSC MultiZ alignments found at http://hgdownload. cse.ucsc.edu/goldenPath/hg19/multiz46way/. Computations were performed using the University of California, San Francisco, Biostatistics High-Performance Computing System.

#### D-statistics and estimates of admixture rates

D-statistics, introduced by Green et al. (2010), are summary statistics for genome sequences from four populations. Two populations,  $P_1$  and  $P_2$ , are compared to a test population,  $P_3$ . The fourth population  $P_4$  is used as an outgroup to determine which allele is ancestral at each site. In our case,  $P_4$ is the chimpanzee reference sequence (pantro2) denoted by C, and  $P_3$  is the Neanderthal sequence, denoted by N.  $P_1$  and  $P_2$  are two human sequences. The chimp reference sequence is assumed to have the ancestral allele, denoted by A. D is computed only for sites at which both of the Neanderthal and one but not both of the human sequences have a different allele, assumed to be derived and denoted by *B*. That is, only those sites with configurations ABBA and BABA are used, where the order is  $P_1$ ,  $P_2$ ,  $P_3$ ,  $P_4$ . The requirement that two copies of both the derived and the ancestral alleles be present greatly reduces the effect of sequencing error (Durand et al. 2011).

When only a single sequence from each population is available,

$$D(P_1, P_2, P_3, P_4) = \frac{n_{ABBA} - n_{BABA}}{n_{ABBA} + n_{BABA}},$$
(1)

where  $n_{ABBA}$  and  $n_{BABA}$  are the numbers of sites with each of the two configurations. When diploid sequences from each individual from  $P_1$  and  $P_2$  are available, then

$$D(P_1, P_2, P_3, P_4) = \frac{\sum_i (1 - p_i^{(1)}) p_i^{(2)} - \sum_i p_i^{(1)} (1 - p_i^{(2)})}{\sum_i (1 - p_i^{(1)}) p_i^{(2)} + \sum_i p_i^{(1)} (1 - p_i^{(2)})}, \quad (2)$$

where  $p_i^{(1)}$  and  $p_i^{(2)}$  are the frequencies of the derived allele (0, 0.5, 1) in the individual in  $P_1$  and  $P_2$ , respectively at site *i*. Equation 2 is equivalent to sampling one of the chromosomes at random from  $P_1$  and  $P_2$  and then using Equation 1.

Green *et al.* (2010) and Durand *et al.* (2011) showed that the expected value of *D* is 0 if  $P_1$  and  $P_2$  form a clade and  $P_3$ is the outgroup. These articles also showed that if there was admixture from  $P_3$  into  $P_2$ , then E(D) > 0. The magnitude of *D* depends on the admixture proportion *f* and on the population divergence times and various effective population sizes.

Reich *et al.* (2010) showed that if there is a sister group of  $P_3$ , which we call  $P_5$ , that has not admixed with  $P_1$ ,  $P_2$ , or  $P_3$ , then it is possible to estimate *f* directly. In our case,  $P_5$  is the Denisovan genome. To estimate *f*, we define  $S(P_1, P_2, P_3, P_4)$  to be the numerator of either Equation 1 or Equation 2. Then

$$\hat{f} = \frac{S(P_1, P_2, P_5, P_4)}{S(P_1, P_3, P_5, P_4)}.$$
(3)

The intuition behind this estimator is that the denominator quantifies the excess coalescent events that occur between lineages in  $P_3$  and  $P_5$  because they are sister groups. Lineages in  $P_2$  that are introduced by admixture have the same coalescent history as all lineages from  $P_3$ . Hence, the ratio is the fraction of lineages in  $P_2$  that trace their ancestry to  $P_3$  because of admixture (Reich et al. 2010). In our application of this method, we are assuming that there is no admixture from Denisovans  $(P_5)$  into the other populations  $(P_1, \ldots, P_4)$ . Although Skoglund and Jacobsson (2011) have argued that there was admixture from Denisovans into East Asians, our results described below did not find evidence of this admixture for the Han Chinese and Japanese samples we analyzed. For analysis A, we explored the variation in estimated D-statistics and admixture rates (f) for all pairs of individuals of different human populations. For analysis B, since we did not include the Denisova genome, we estimated only Dstatistics.

#### Randomization tests

We computed D for each pair of individuals, both within populations and between populations. We developed two randomization tests of statistical significance. Both are similar to the Mantel test. Test 1 tests whether the average Dcomputed for one pair of populations is significantly larger than for another pair, and test 2 tests whether the average Dfor a pair of populations differs significantly from 0.

For test 1, we start with sequences from three human populations,  $G_1$ ,  $G_2$ , and  $G_3$ , each containing  $k_1$ ,  $k_2$ , and  $k_3$  diploid sequences. We compute two matrices of *D* values.



**Figure 2** Schematic of a model of recent and ancient population structure without admixture used in simulations. See text for details.

The elements of  $M_1$  are  $D(G_{1,i}, G_{3,j}, N, C)$ , where  $G_{1,i}$  and  $G_{3,j}$ are the *i*th and *j*th individuals in  $G_1$  and  $G_3$  ( $i = 1, ..., k_1$ ;  $j = 1, ..., k_3$ ). The elements of  $M_2$  are  $D(G_{2,i}, G_{3,j}, N, C)$ .  $M_1$ has  $k_3$  rows and  $k_1$  columns, and  $M_2$  has  $k_3$  rows and  $k_2$ columns. From  $M_1$  and  $M_2$  the average D's are computed,  $D_1$ and  $D_2$ . The problem is to test whether  $D_1 = D_2$ . A *t*-test cannot be used because the elements within each matrix are not independent of each other and because the same reference population  $(G_3)$  is used to compute both matrices. Instead, we combine  $M_1$  and  $M_2$  into a single matrix with  $k_3$ rows and  $k_1 + k_2$  columns. Then we randomize the columns and compute  $D_1$  for the matrix containing the first  $k_1$  columns and  $D_2$  for the matrix containing the last  $k_2$  columns. Then we compare the observed  $D_1 - D_2$  with the distribution of differences from the randomized matrices. We used a two-tailed test and 1 million replicates for each test.

Test 2 is similar to test 1, but because we compare only  $G_1$  and  $G_2$ , a subset of one population is used in place of the reference population,  $G_3$ . For the population with the larger sample size (say  $G_1$ ), we create a random partition  $(G_1^a, G_1^b)$  subject to the constraint that they differ in number by no more than one. For  $M_1$ , we compute D for all pairs of individuals in  $G_1^a$  and  $G_2$ . The elements of  $M_2$  are  $D(G_{1,i}^a, G_{1,j}^b, N, C)$ , where  $G_{1,i}^a$  and  $G_{1,j}^b$  are the *i*th and *j*th individuals in the two subpopulations created by the partition. Test 1 is then applied to  $M_1$  and  $M_2$ .

We also calculated the *f*-statistics for each pair of individuals. Using the same randomization tests as described above, we determined whether there were significant differences between populations in estimates of the admixture rate. Significant differences observed using the admixture rate suggest that the effect is truly due to the Neanderthal and not admixture with Denisovans.

#### Identifying putative archaic human regions

Previous work has shown that archaic admixture often leads to long, divergent haplotypes at low frequency (Wall 2000; Plagnol and Wall 2006). We define two SNPs to be "congruent" if their diploid allele counts (*i.e.*, zero, one, or two counts of a particular allele) across individuals are completely correlated (*i.e.*,  $r^2 = 1$ ). We define the maximum number of pairwise congruent SNPs to be  $l_d$  and denote the collection of rarer (minor allele frequency  $\leq 0.5$ ) alleles at each of these pairwise congruent sites to be the putative archaic haplotype. From the filtered Complete Genomics data, we then identified all regions from 8 to 100 kb in length where  $l_d \geq 30$  and  $l_d/S \geq 0.1$ , where *S* is the total number of polymorphic sites in the region. When identified regions overlapped, we took the region with the largest value of  $l_d/S$ . We also required that neighboring regions with putative archaic haplotypes congruent with each other be separated by at least 200 kb, to avoid double counting long archaic haplotypes. A total of 2254 regions were identified. Of these, 411 were private to the non-African samples.

To estimate what proportion of these regions might be false positives, we simulated whole-chromosome sequence data (Chen et al. 2009) under a model that incorporated both recent (intracontinental) and ancient (intercontinental) population structure (Figure 2). Specifically, we assume a panmictic ancestral population split into two daughter populations at time  $T_0 = 0.6$  (using the standard coalescent scaling of 4N generations), with (symmetric) scaled migration rate of  $M_0 = 5$ . At time  $T_1 = 0.05 - 0.053$ , one of the ancestral populations (i.e., the "non-African" one) experiences a population bottleneck resulting in a 100-fold reduction in population size. Then, at time  $T_2 = 0.045$ , each population splits into two descendant populations, connected by migration rate  $M_1 = 8$ . While arbitrary, this model attempts to incorporate the major features of human demographic history, including intra- and intercontinental population structure and a bottleneck in the history of non-African populations, and is similar to the model used by Yang et al. (2012). The results described below are qualitatively similar if other plausible values for the times and migration rates are used (results not shown). Using N = 10,000 and an average generation time of 25 years, each unit of scaled time corresponds to 1 million years.

We simulated 30 different 100-Mb chromosomes, using the model described above with mutation parameter  $\theta$  =  $3.5 \times 10^{-4}$ /bp, recombination parameter  $\rho = 4 \times 10^{-4}$ /bp, and 10 individuals sampled from each of the four extant populations. The simulated number of segregating sites was substantially higher than the actual number in our filtered data. Since average  $l_d$  values are positively correlated with levels of diversity, the simulated  $l_d$  values are higher on average than expected in real data, and our choice of  $\theta$  is conservative. Also, standard estimates of  $\rho$  are generally higher than the value we took (Myers et al. 2005), which is also conservative for our purposes. We then tabulated the total number of regions with  $l_d \ge 30$ ,  $l_d/S \ge 0.1$ , and with divergent haplotype SNPs private to the simulated non-African samples. We identified a total of 3 regions that satisfied these criteria, compared with 411 regions that were identified from the actual data. This leads to an estimate of a false discovery rate of q < 0.01.

#### Identifying putative Neanderthal regions

To identify which of the 2254 regions described above were likely to reflect recent Neanderthal admixture, we imposed the following additional criteria on the putative archaic human haplotypes:

- 1. The Neanderthal allele must be called at  $\geq$ 12 SNPs and match the putative archaic haplotype at  $\geq$ 70% of these SNPs.
- 2. The Neanderthal allele and the chimp allele must be called at  $\geq$ 8 SNPs and the Neanderthal allele must be derived (relative to chimp) at  $\geq$ 60% of these sites.
- 3. The putative archaic haplotype must be at low frequency (<5%) in the sub-Saharan African samples.

The motivation for criterion 1 is obvious, and we note that a more stringent cutoff was not used due to the poor quality of the Neanderthal genome sequence. Criterion 2 was implemented to cut down on regions that reflect shared ancestral polymorphism between modern humans and Neanderthals; it is based on an observation of Noonan et al. (2006) that recent Neanderthal admixture will lead to an increase in SNPs where Neanderthals have the derived allele. Finally, criterion 3 reflects our prior belief that admixture with Neanderthals did not occur in Africa and that the presence of Neanderthal alleles in Africa could reflect only more recent migration patterns. A total of 226 regions were identified that meet these additional criteria. We note in passing that the specific cutoffs used in criteria 1-3 are somewhat arbitrary, but our qualitative conclusions are unchanged under a range of similar criteria (results not shown).

We implemented a simple permutation test to assess the statistical significance of the observed difference in frequencies of Neanderthal regions in East and South Asians and Europeans. Specifically, we kept the presence/absence of Neanderthal regions for each individual constant and randomly permuted the geographic label (*i.e.*, "European" *vs.* "East Asian") of the sample 100,000 times. Similar analyses were used to compare the frequency of Neanderthal regions in Maasai *vs.* other sub-Saharan African samples.

#### Identifying putative Denisovan regions

Excluding the 226 Neanderthal regions identified above, we screened the remaining 2028 putative archaic regions for Denisovan admixture, using the same criteria as for Neanderthals. Thirty total regions fit these criteria.

#### Estimating local ancestry in the Maasai

We took the filtered Complete Genomics data described at the start of this section and estimated SNP allele frequencies separately in the 13 European samples and the 13 non-Maasai African samples. These were used as proxies for the (unknown) non-African and African ancestral populations. We then included only those SNPs with allele frequencies that differ by at least 0.3 in our analyses. We calculated the



Figure 3 Summary of significance tests for average values of D. Positive values indicate that the second sequence is more similar to the Neanderthal genome than the first sequence. In all parts, the box plots indicate the range of D values obtained for pairs of individuals from the populations indicated. A and B are box plots of individual D-statistics computed for each individual from the specified population compared with each Yoruban. P-values are from the randomization test, test 1, of significant differences in the average D values for different pairs of populations. C and D show box plots of individual D-statistics computed for every pair of individuals in the specified populations. P-values are from the randomization test, test 2, of significant differences of the average D from 0. See also Table S2.

likelihood of each ancestral configuration (*i.e.*, zero, one, or two alleles inherited from the non-African population) separately for each SNP. Then, over sliding windows of 1 Mb, we formed a composite likelihood by multiplying together all of the single-SNP likelihoods contained in the window and tabulated which ancestral configuration had the highest (composite) likelihood. For each SNP, we then used majority rule to make ancestry calls, using all windows containing the SNP in question. See Wall *et al.* (2011) for further details.

## Results

#### D-statistics and estimates of f

The D-statistics and estimates of f we computed are summarized in Figure 3 and Supporting Information, File S1, Table S1, Table S2, Table S3, Table S4, Table S5, Table S6, Table S7, Table S8, Table S9, Figure S1, Figure S2, Figure S3, Figure S4, Figure S5, Figure S6, Figure S7, and Figure S8. Several features of the results are notable. First, we find evidence for more Neanderthal admixture into the East Asian samples than into the European samples (P =0.001)—consistently higher D values result when East Asians are compared to one of the African populations than when Europeans are compared (Figure 3A, Table S4), and the average D is positive when East Asians are compared to Europeans (Figure 3C, Table S5). In analysis B, comparisons with the South Asian samples are intermediate with respect to the European and East Asian samples but not in analysis A, indicating that the South Asian sample differs from the East Asian ones but the degree of similarity to Europeans remains to be established. Also, we find evidence for a small but significant amount of Neanderthal admixture into the Maasai genomes ( $P \sim 0.03$ , Table S4). When compared to the Yoruba, the Maasai have a higher average D than the Luhya (Figure 3B, Table S4). When the Maasai are compared to all other African samples, the average D is positive (Figure 3D). In addition, when East Asians and Europeans are compared to the Maasai, the average D's are somewhat lower than when they are compared to either the Yoruba or the Luhya. The *P*-values shown in Figure 3, A and B are from test 1 and those in Figure 3, C and D are from test 2.

Table S1, Table S2, and Table S3 show estimated values of *f*. The estimates of the admixture rate show that when we incorporate the Denisovan genome into our analysis, the admixture rate between East Asians and Neanderthals remains significantly higher than the admixture rate between Europeans and Neanderthals ( $P \sim 0.001$ , Table S7). The Maasai remain significantly more genetically similar to the Neanderthals when compared to the Luhya ( $P \sim 0.03$ , Table S7), but the observed significant difference for the *D*-statistic ( $P \sim 0.34$ , Table S7), which probably reflects the lower power of using *f* as a test statistic. The admixture rates for the South Asians give the same results as those for the *D*-statistic (Table S9).

### Identifying "Neanderthal haplotypes"

Our new method for identifying introgressed Neanderthal fragments in human populations detected 226 different putative Neanderthal regions. The relative frequencies of



**Figure 4** Distribution of the number of putative Neanderthal regions for each Eurasian individual. European genomes are colored in green, East Asian genomes are colored in red, and South Asian genomes are colored in black.

these putative Neanderthal haplotypes in the 42 sampled modern human individuals then provide estimates of the relative contributions of Neanderthal DNA to the gene pools of contemporary human populations. We found that on average the "Neanderthal haplotypes" were at higher frequency in the East Asians than in the Europeans (9.6% vs. 6.4%;  $P = 3.0 \times 10^{-4}$ , permutation test), consistent with the *D*-statistic results presented in Figure 3 (Figure 4). We also found evidence for a small, but statistically significant, Neanderthal contribution to the genomes of the Maasai ( $P = 4.9 \times 10^{-4}$ ), but did not find a significant difference in Neanderthal haplotype frequency between the East Asian and South Asian samples (P > 0.05).

#### Additional test of ancient population structure

As reviewed in the Introduction, there is already evidence against the hypothesis that the extra similarity of non-African populations to Neanderthals is accounted for by ancient population subdivision. To explore this point further, we took the 411 regions from our whole-genome analyses that were identified purely on the basis of their LD patterns (*i.e.*, without using any information from the Neanderthal genome sequence). Then, for each non-African individual, we calculated the D-statistic for those regions where the individual contained a rare, diverged haplotype. If this haplotype were recently inherited from Neanderthals, we would expect the D values to be strongly positive. If instead there were no recent admixture between modern humans and Neanderthals, then there is no a priori reason why these regions would show D values significantly different from 0. Recombination acting over the past 300 Kyr would break up local patterns due to shared ancestral polymorphisms to scales <0.01 cM (i.e., <10 kb on average). The *D* values that we observe are strongly positive (average D = 0.594, compared with an average D = 0.068 for the whole genome), providing additional evidence that most of the unusual haplotypes from these 411 regions are indeed the result of recent introgression from the Neanderthal gene pool ( $P \ll 10^{-8}$ , Figure 5).

# Identifying "Denisovan haplotypes"

Excluding the 226 Neanderthal regions described above, we used the same criteria to identify regions likely inherited

from Denisovans. We identified a total of 30 regions, all at low frequency, with no significant difference in frequency between populations.

#### Maasai admixture

Previous genetic studies have suggested that the Maasai may be an admixed population with a substantial proportion of non-African ancestry (Henn et al. 2011). If the non-African ancestry were due to recent (i.e., post-Neanderthal) admixture, then the observation of Neanderthal ancestry in the Maasai would not be unexpected. Alternatively, spatially explicit models of ancient population structure might explain the greater similarity between Maasai and Neanderthals relative to other sub-Saharan African groups (A. Manica, personal communication). One difference between these alternative explanations is what they predict about the patterns of similarity across the genomes of Maasai individuals. Under a model of recent admixture, we expect Maasai genomes to show large, distinct blocks of sequence with different genetic patterns, corresponding to blocks with non-African vs. African ancestry. The average size of the non-African blocks (in morgans) is roughly the inverse of the time (in generations) since admixture. In contrast, under a model of ancient admixture the similarity of Maasai genomes to the Neanderthal genome will be spread throughout the genome because the admixture happened much longer ago.

To distinguish between these two possibilities, we employed a composite-likelihood-based approach to identifying African and non-African regions of ancestry across the genomes of the four Maasai samples (Wall *et al.* 2011). Briefly, we used the European (CEU and TSI) and other African (YRI and LWK) samples (Table 1) to estimate allele frequencies in non-African and African ancestral populations and then estimated the number of alleles inherited from each ancestral population at each SNP in the genome. These extant samples may not be perfect proxies for the true ancestral populations, but the qualitative results presented below are likely to be valid.

In summary, we estimate an average of  $\sim$ 30% non-African ancestry in each Maasai genome, and the sizes of the ancestral blocks are consistent with admixture that



**Figure 5** Box plot showing the average D across the whole genomes of the non-African individuals compared with the average D (for the same individuals) across regions identified as having unusual patterns of LD (*i.e.*, putative archaic regions).

happened ~100 generations ago (Figure 6A). We then partitioned each Maasai genome into regions with zero, one, or two inferred African alleles and calculated *D* separately for each partition. We found that the *D* values are significantly more negative with increasing numbers of inferred non-African alleles ( $P = 2.0 \times 10^{-4}$ ; Figure 6B). This observation provides strong support for recent non-African gene flow into the Maasai, with the non-African alleles bringing with them low levels of Neanderthal ancestry.

#### Discussion

Our results confirm and reinforce several conclusions about admixture between Neanderthals and the ancestors of modern humans. Using a much larger number of highcoverage genome sequences than were previously analyzed for this purpose and using two complementary methods of analysis (D-statistics and detection of introgressed Neanderthal segments), we confirm the conclusion of Meyer et al. (2012) that East Asians (Han Chinese and Japanese) are more similar to the published Neanderthal sequence than are Europeans. Because we have analyzed more modern human sequences than Meyer et al. (2012) did, we are able to show the extent of variation within both Asian and African populations. We also confirm the conclusions of Yang et al. (2012) and Sankararaman et al. (2012) that the similarity of both Europeans and East Asians to Neanderthals is the result of recent admixture and not ancient population subdivision. Finally, we used the high-coverage Denisova sequence of Meyer et al. (2012) to determine that the admixture rate (f) into East Asians is  $\sim$ 40% higher than into Europeans.



**Figure 6** Recent and ancient admixture in the Maasai. (A) Representative plot of the number of estimated "African" alleles across the first 30 Mb of chromosome 1 in one of the Maasai genomes. (B) Estimated values of *D* for portions of the genome estimated to contain zero, one, or two "non-African" alleles.

We were not able to confirm the conclusion of Skoglund and Jakobsson (2011) that there was Denisovan admixture into East Asians. We did not detect any difference in the number of apparent Denisovan segments in Europeans and East Asians. The East Asian genomes analyzed, however, were from northern East Asia (Beijing and Tokyo), not from southern East Asia where Skoglund and Jakobsson found the strongest signal of admixture with Denisovans.

Our results and those of Meyer *et al.* (2012) imply that the relatively simple admixture scenario proposed by Green *et al.* (2010) needs to be altered. At least two separate episodes of admixture between Neanderthals and modern humans must have occurred, and at least one of those episodes must have occurred after the separation of the ancestors of modern Europeans and East Asians. Rather than have two distinct episodes of admixture, it seems more plausible that admixture took place over a protracted period 50–80 KYA. During that period the ancestors of Europeans diverged and subsequently experienced less admixture than the ancestors of East Asians. This scenario is consistent with the simulation models of Currat and Excoffier (2011) and Skoglund and Jakobsson (2011).

If this scenario is correct, the time of separation of the ancestors of modern European and East Asian populations is constrained. Since there is no archaeological record of Neanderthals in the past  $\sim$ 30 Kyr, it follows that the separation of Europeans from East Asians had to have occurred before Neanderthals went extinct. Consequently, estimates of East Asian–European population divergence of <30 KYA (Gutenkunst *et al.* 2009; Gravel *et al.* 2011) are unlikely to be correct. This timeframe is also supported by a 40- to 50-KYA modern human fossil recently found in China (Fu *et al.* 2013).

Our two analyses yielded slightly different results for the Gujarati (South Asian) samples. However, it would not be surprising if the true level of Neanderthal ancestry in South Asians was intermediate between Europeans and East Asians because previous studies have shown gradients in genetic ancestry across Eurasia (Rosenberg *et al.* 2002).

Our finding of Neanderthal admixture into the Maasai was initially surprising, given the lack of evidence that Neanderthals ever crossed into Africa or that the ancestors of the Maasai were ever in the Middle East. Although direct contact between the two groups in the past is theoretically possible, our results are more consistent with a scenario involving recent admixture between the ancestors of the Maasai and one or more (historically) non-African groups with Neanderthal ancestry several thousand years ago. This interpretation is broadly consistent with recent findings of African admixture into Middle Eastern and Southern European populations during the same timescale (Moorjani et al. 2011) and a greater genetic similarity between East African and non-African samples than between West African and non-African samples (Tishkoff et al. 2009). Together these studies provide additional support for the hypothesis that admixture between genetically diverged groups is a common feature of human demographic history.

The new picture of human and Neanderthal ancestry that emerges from our results is almost certainly not complete, and our results suggest that intracontinental variation in levels of Neanderthal ancestry may be common. With the current rate of progress in whole-genome sequencing and the possibility of additional draft genomes from specimens of archaic individuals, we will soon learn more about the admixture process. In particular, the construction of "archaic admixture maps" detailing the distribution of archaic DNA segments in different modern human populations will help us to infer the timing, locations, and exact numbers of introgression events and the role that archaic admixture may have played in the evolution of the AMH genome.

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# GENETICS

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# Higher Levels of Neanderthal Ancestry in East Asians than in Europeans

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#### File S1

#### Additional D-statistic results

We computed D(P1, P2, Neanderthal, Outgroup) for all pair of individuals (P1,P2) from the Complete Genomics data, as described in the Materials and Methods. The D-statistics were averaged over all combinations of individuals for each given pair of populations. The means and the standard deviations for all twenty-eight pairwise population comparisons are given in Tables S1 and S2. We also show regional comparisons, where some populations are grouped into East Asian, European, or African (Tables S2 and S3). We also computed the admixture rate (*f*) for each of these combinations (Tables S1-S3) for the data from Analysis A. Where we have results from both Analysis A and Analysis B, we show the results in the text in curly braces, such that the results from the two analyses are given as {Analysis A, Analysis B}.

#### Comparison between non-Africans and Africans

The averaged D-statistics are consistently positive when comparing African populations and non-African populations (Average D-statistic range = {[0.0429, 0.0891], [0.0530, 0.0750]}, Table S1, Figure S1). These results confirm the previous findings that the non-African populations are more closely related to Neanderthal than African populations (Green et al. 2010). The admixture rate estimated also gives positive values ranging from 0.0191 to 0.0417 (Table S1, Figure S2).

#### Comparison between Europeans and East Asians (Figure S3-S4)

When we compared the set of D-statistics for the pair (Afr, East Asia) and the pair (Afr, Europe) using Test 1, we find that the D-statistics for the East Asian individuals tend to be higher than the D-statistics for the European individuals (mean difference = {0.0083, 0.0096}, two-tailed p-value = {0.0010, 0.0006}, Table S4). This suggests that East Asians may have a greater signal for genetic admixture with Neanderthals than Europeans. These results are consistent when we consider each African population separately and across both Analysis A and Analysis B. The difference between Europeans and East Asians is always significant using Test 1 (Table S4).

This trend is further supported by the set of positive D-statistics estimated for the pair (Europe, East Asia). The values averaged over each population are given in Table S2. The average D-statistic for the merged East Asian group compared to the merged European group is {0.0110, 0.0131}, which is significantly different from zero (twotailed p-value = {0.0037, 0.0009}, Table S5).

The *f*-statistics also show the same trend, with a higher signal between Neanderthals and East Asians, compared to Neanderthals and Europeans. Test 1 shows that the difference in *f*-statistic of 0.0098 is significant (two-

tailed p-value = 0.0011, Table S7). Test 2 compares Europeans and East Asians directly, and shows that the *f*-statistic computed is 0.0100, which is significantly different from zero (two-tailed p-value = 0.0072, Table S8).

#### Gujarati population (Figure S5-S6)

We also studied whether the South Asian population GIH was more similar to the Europeans or the East Asians in term of admixture from Neanderthal. When compared to the African individuals, the GIH individuals have an average D-statistic of {0.0712, 0.0656} (Table S3), which is higher than the average D-statistic for Europeans (average D-statistic for (Afr, Europe) = {0.0644, 0.0604}, Table S3) and lower than the average D-statistic for East Asians (average D-statistic for (Afr, East Asia) = {0.0727, 0.0699}. The same results are observed when considering each African population separately (Table S3). We applied both Test 1 and Test 2 to investigate the significance of these observations. When we use Test 1, we find that the difference in the estimates of D for the pair (Afr ,GIH) and the pair (Afr, East Asia) are significant when comparing against all Africans (two-tailed p-value = {0.0101, 0.0259}, Table S6). However, the difference in the estimates of D for (Afr, GIH) and (Afr, Europe) is not significant (two-tailed p-value = {0.4232,0.1343}, Table S6). Thus, the average D-statistics found for GIH are closer to the estimates of D for the European samples than for the East Asian samples. When we use Test 2, we find that D-statistics for (Europe, GIH) are not significantly different from 0 (D={0.0035, 0.0067}, p-values = {0.4386, 0.2345}, Table S5). D for (GIH, East Asia) are significantly different from zero in Analysis A (two-tailed p-value=0.0346), but must be taken with caution as the estimate is not significantly different in Analysis B (two-tailed p-value=0.0867). The results from Test 2 cannot distinguish if the GIH samples group more closely with East Asians or Europeans, while Test 1 does. Test 1 and Test 2 for the *f*-statistic show similar results (Table S8-S9).

#### Maasai population (Figure S7-S8)

The Maasai individuals (MKK) seem to share more genetic similarity with Neanderthals than other African populations. The average D-statistic for (Afr, MKK), with Afr =YRI or LWK, were positive (average D-statistic = {(0.0110, 0.0075), (0.0102, 0.0145)}, Table S2). Using Test 2, the average D-statistic for (YRI+LWK, MKK) is significantly different from zero (D={0.0123,0.0116}, two-tailed p-value={0.0101, 0.0135}). However, the significant difference from zero is lost when the Maasai are compared separately to the Yoruba or Luhya, except for the (YRI,MKK) comparison in Analysis A (two-tailed p-value=0.0418, Table S5). Notably, the (LWK, YRI) pair is not significantly different from zero (two-tailed p-value = {0.3457, 0.3611}, Table S5).

When we compare the estimates of *D* for the pair (YRI, East Asia) to (MKK, East Asia) using Test 1, the estimates of *D* for (MKK, East Asia) were significantly different from the estimates of *D* for (YRI, East Asia) by a small

3 SI

amount (difference = {0.0074, 0.0051}, two-tailed p-value = {0.0054, 0.0352}, Table S4), indicating that the signal of archaic admixture in non-Africans is weaker when we use the MKK as a reference. We also see a significant difference using the Europeans instead of East Asians (two-tailed p-value = {0.0098, 0.0404}, Table S4). A significant difference is also observed when switching between the Maasai and the Luhya as the reference in Analysis A (Table S4), but is above the 0.05 significance threshold in Analysis B. However, again, it is striking to see that there is no significant difference in D-statistics between (YRI, East Asia) and (LWK, East Asia) (two-tailed p-value = {0.1302, 0.1235} for East Asians and {0.1234, 0.1243} for Europeans, Table S4).

The results for the *f*-statistics show no significant difference between the Maasai and the other two African populations for Test 2 (two-tailed p-value = 0.2021, Table S8). Comparisons of the Maasai separately to the Yoruba and the Luhya show that the main reason for the lack of significance is no significant difference in f when comparing the Yoruba and Maasai (two-tailed p-value = 0.4944 for Europe, 0.4284 for East Asia, Table S7). The estimates of *f* using the Luhya are, however, significantly different from the estimates using the Maasai for both Europeans (twotailed p-value = 0.0286, Table S7) and East Asians (two-tailed p-value = 0.0286, Table S7). Test 2 also shows that the Maasai have a significantly greater admixture rate relative to the Luhya (two-tailed p-value = 0.0666, Table S8), but not the Yoruba (two-tailed p-value = 0.4847, Table S8).

#### Consistency

All the D statistics, f statistics, and p-values of the randomization tests were calculated for two sets of slightly differently prepared data (see Materials and Methods). The results are presented in all tables under the columns *Analysis A* and *Analysis B*. For both analyses, the East Asian populations show a significantly higher estimate of *D* than the European populations. The two analyses also consistently show the South Asian Gujarati population exhibiting D-statistics closer to the European population than the East Asian populations. Both analyses also show results that suggest the MKK has more shared genetic variants with Neandertals compared to the other African populations. The differences in data preparation, while giving slightly different estimates of *D*, do not change our conclusions.

The *f*-statistics also suggest higher admixture into East Asians over Europeans and more similarity in admixture rates between the GIH and Europeans, as compared to GIH and East Asians, but the Maasai genetic similarity is not observed.

	Populations			Analysis A			Analysis B	
	Compa	red		D		f	[	)
	P2	P1	Mean	StDev	Mean	StDev	Mean	StDev
	CEU	YRI	0.0658	0.0059	0.0232	0.0069	0.0615	0.0064
SL	CEU	LWK	0.0702	0.0069	0.0308	0.0068	0.0659	0.0073
oear	CEU	МКК	0.0586	0.0060	0.0219	0.0069	0.0566	0.0063
lor	TSI	YRI	0.0635	0.0069	0.0207	0.0063	0.0583	0.0077
ш	TSI	LWK	0.0676	0.0079	0.0285	0.0065	0.0627	0.0086
	TSI	MKK	0.0561	0.0073	0.0191	0.0059	0.0530	0.0080
	СНВ	YRI	0.0717	0.0037	0.0307	0.0049	0.0695	0.0038
JS	СНВ	LWK	0.0762	0.0055	0.0376	0.0044	0.0738	0.0051
Asiaı	СНВ	МКК	0.0642	0.0036	0.0290	0.0048	0.0644	0.0031
ast ∕	JPT	YRI	0.0751	0.0045	0.0340	0.0042	0.0707	0.0043
Ш	JPT	LWK	0.0790	0.0061	0.0417	0.0038	0.0750	0.0055
	JPT	MKK	0.0679	0.0045	0.0324	0.0037	0.0656	0.0039
د s	GIH	YRI	0.0675	0.0037	0.0264	0.0038	0.0657	0.0047

0.0054

0.0034

0.0347

0.0245

0.0033

0.0037

0.0701

0.0606

0.0058

0.0041

Table S1 Average D and f found for CGDP populations (Afr, non-Afr)

South Asians

GIH

GIH

LWK

MKK

0.0719

0.0601

# Table S2: Average D and f (comparisons within Africans and non-Africans)

	Populations			Analysis A			Analy	Analysis B	
	Com	npared	D		f	f			
	P2	P1	Mean	StDev	Mean	StDev	Mean	StDev	
a a	YRI	LWK	0.0042	0.0063	0.0081	0.0039	0.0042	0.0069	
/ithi	МКК	YRI	0.0110	0.0050	0.0023	0.0047	0.0102	0.0051	
N A	МКК	LWK	0.0154	0.0062	0.0101	0.0037	0.0145	0.0063	
-u									
on r can ions	CEU	TSI	0.0034	0.0102	0.0026	0.0084	0.0049	0.0120	
ithiı Afri Reg	JPT	СНВ	0.0050	0.0049	0.0037	0.0049	0.0019	0.0043	
$\sim$									
JS	СНВ	CEU	0.0077	0.0062	0.0075	0.0071	0.0110	0.0069	
gior	СНВ	TSI	0.0110	0.0077	0.0096	0.0067	0.0152	0.0086	
n re	JPT	CEU	0.0124	0.0068	0.0111	0.0070	0.0125	0.0073	
rica	JPT	TSI	0.0152	0.0084	0.0136	0.0063	0.0168	0.0089	
η-Af	СНВ	GIH	0.0056	0.0042	0.0048	0.0045	0.0062	0.0042	
non	JPT	GIH	0.0100	0.0030	0.0089	0.0035	0.0079	0.0049	
een	GIH	CEU	0.0025	0.0057	0.0024	0.0068	0.0053	0.0081	
etw	GIH	TSI	0.0057	0.0066	0.0048	0.0059	0.0098	0.0097	
B	East Asia	Europe	0.0110	0.0074	0.0100	0.0071	0.0131	0.0078	

## Table S3 Average D and f found for merged populations (non-Afr, Afr)

Populations			Anal	Analysis B			
C	ompared	D		f		D	
P2	P1	Mean	StDev	Mean	StDev	Mean	StDev
5	YRI	0.0651	0.0063	0.0225	0.0068	0.0605	0.0069
pea	LWK	0.0694	0.0072	0.0301	0.0067	0.0649	0.0078
nro	МКК	0.0579	0.0065	0.0210	0.0067	0.0555	0.0070
ш	all Afr	0.0644	0.0077	0.0239	0.0076	0.0604	0.0078
Ē	YRI	0.0734	0.0045	0.0324	0.0048	0.0701	0.0041
Asia	LWK	0.0776	0.0059	0.0398	0.0045	0.0744	0.0053
East /	МКК	0.0660	0.0044	0.0307	0.0046	0.0650	0.0035
	all Afr	0.0727	0.0063	0.0337	0.0058	0.0699	0.0053
GIH	all Afr	0.0712	0.0059	0.0280	0.0053	0.0656	0.0058

Table S4 Randomization test (Test 1) p-values (Two-tailed: Group 1 different from Group 2, One Tail: Group 1 > Group 2, where the sets compared are (Group 1, Group Ref) versus (Group 2, Group Ref))

	Sets of			Analysis A		Analysis B		
	Populations	Compared	p (Two-tailed)	p (One Tail)	Difference	p (Two-tailed)	p (One Tail)	Difference
5	(East Asia, YRI)	(Europe , YRI)	0.0008	0.0001	0.0084	0.0004	0.0000	0.0096
ans	(East Asia, LWK)	(Europe , LWK)	0.0010	0.0002	0.0082	0.0005	0.0000	0.0095
Afric	(East Asia, MKK)	(Europe , MKK)	0.0017	0.0003	0.0082	0.0009	0.0000	0.0095
	(East Asia, Afr)	(Europe , Afr)	0.0010	0.0002	0.0083	0.0006	0.0000	0.0096
	(MKK, YRI)	(LWK, YRI)	0.0285	0.0142	0.0187	0.0286	0.0144	0.0144
sue	(MKK, LWK)	(YRI, LWK)	0.0014	0.0014	0.0111	0.0042	0.0042	0.0104
rica	(YRI, Europe)	(LWK, Europe)	0.1234	0.0674	0.0043	0.1243	0.0631	0.0044
g Af	(YRI, East Asia)	(LWK, East Asia)	0.1302	0.0728	0.0042	0.1235	0.0621	0.0043
arin	(MKK, Europe)	(LWK, Europe)	0.0284	0.0142	0.0115	0.0855	0.0429	0.0095
mp;	(MKK, East Asia)	(LWK, East Asia)	0.0285	0.0143	0.0116	0.0857	0.0430	0.0094
Co	(MKK, Europe)	(YRI, Europe)	0.0098	0.0070	0.0072	0.0404	0.0224	0.0050
	(MKK, East Asia)	(YRI, East Asia)	0.0054	0.0054	0.0074	0.0352	0.0225	0.0051

Table S5 Randomization test (Test 2) p-values. Two-tailed: D(P1,P2) not zero; One-tail: D(P1,P2)>0, ie P2 is more similar to Neanderthal than P1. The largest group was used as reference (see Methods); for (LWK,MKK) both groups were successively used as reference because they have the same number of individuals.

(01 02)		Analysis A		Analysis B			
(「」,「∠)	p (Two-tailed)	p (One Tail)	D estimate	p (Two-tailed)	p (One Tail)	D estimate	
(LWK , YRI)	0.3457	0.181	0.0042	0.3611	0.1800	0.0042	
(YRI <i>,</i> MKK)	0.0418	0.0208	0.011	0.0524	0.0262	0.0102	
(LWK <i>,</i> MKK)	0.0669 / 0.1208	0.0669 / 0.094	0.0154	0.0670 / 0.0763	0.0667 / 0.0662	0.0145	
(YRI+LWK, MKK)	0.0101	0.007	0.0123	0.0135	0.0088	0.0116	
(Europe , East Asia)	0.0037	0.0028	0.011	0.0009	0.0008	0.0131	
(GIH , East Asia)	0.0346	0.0174	0.0078	0.0867	0.0435	0.0070	
(Europe , GIH)	0.4386	0.2358	0.0035	0.2345	0.1095	0.0067	

Table S6 Randomization test (Test 1) p-values for GIH (Two-tailed: Group 1 different from Group 2, One Tail: Group 1 > Group 2, where the sets compared are (Group 1, Group Ref) versus (Group 2, Group Ref))

	Sets of			Analysis A			Analysis B		
	Popula	tions Compared	p (Two-tailed)	p (One Tail)	Difference	p (Two-tailed)	p (One Tail)	Difference	
c	(GIH, Afr)	(Europe, Afr)	0.4232	0.2212	0.0024	0.1343	0.0560	0.0052	
l vs pea	(GIH, YRI)	(Europe, YRI)	0.4114	0.2124	0.0024	0.1234	0.0495	0.0052	
GIH	(GIH, LWK)	(Europe, LWK)	0.4062	0.2137	0.0025	0.1338	0.0572	0.0051	
ш	(GIH, MKK)	(Europe, MKK)	0.4728	0.2487	0.0023	0.1565	0.0681	0.0052	
st	(GIH, Afr)	(East Asia, Afr)	0.0101	0.0042	0.0059	0.0259	0.0160	0.0044	
s Ea an	(GIH, YRI)	(East Asia, YRI)	0.0101	0.0041	0.0059	0.0221	0.0160	0.0044	
H v Asi	(GIH, LWK)	(East Asia, LWK)	0.0101	0.0041	0.0057	0.0301	0.0181	0.0044	
B ال	(GIH <i>,</i> MKK)	(East Asia, MKK)	0.0101	0.0041	0.0059	0.0260	0.0159	0.0043	

Table S7: Randomization test (Test 1) p-values for f-statistics (Two-tailed: Group 1 different from Group 2, One Tail:Group 1 > Group 2, where the sets compared are (Group 1, Group Ref) versus (Group 2, Group Ref)))

	Sets	s of		Analysis A			
	Populations	Compared	p (Two-tailed)	p (One Tail)	Difference		
-uo	(East Asia, YRI)	(Europe, YRI)	0.0009	0.0004	0.0099		
ng n ans	(East Asia, LWK)	(Europe, LWK)	0.0016	0.0006	0.0096		
ıpari Afric	(East Asia, MKK)	(Europe, MKK)	0.0011	0.0004	0.0097		
Com	(East Asia, Afr)	(Europe, Afr)	0.0011	0.0004	0.0098		
	(MKK, YRI)	(LWK, YRI)	0.0283	0.0141	0.0104		
ากร	(MKK, LWK)	(YRI, LWK)	0.3449	0.1716	0.0021		
rica	(YRI, Europe)	(LWK, Europe)	0.0043	0.0043	0.0076		
g Af	(YRI, East Asia)	(LWK, East Asia)	0.0056	0.0056	0.0073		
arin	(MKK, Europe)	(LWK, Europe)	0.0286	0.0143	0.0091		
mpa	(MKK, East Asia)	(LWK, East Asia)	0.0286	0.0143	0.0090		
Col	(MKK, Europe)	(YRI, Europe)	0.4944	0.2522	0.0014		
	(MKK, East Asia)	(YRI, East Asia)	0.4284	0.2138	0.0017		

Table S8: Randomization test (Test 2) p-values for f-statistics. Two-tailed: D(P1,P2) not zero; One-tail: D(P1,P2)>0, ie P2 is more similar to Neanderthal than P1. The largest group was used as reference (see Methods); for (LWK,MKK) both groups were successively used as reference because they have the same number of individuals.

(D1 D7)		Analysis A	
(「⊥,「∠)	p (Two-tailed)	p (One Tail)	f estimate
(LWK , YRI)	0.0425	0.0212	0.0081
(YRI , MKK)	0.4847	0.2467	0.0023
(LWK , MKK)	0.0666	0.0666	0.0101
(YRI+LWK, MKK)	0.2021	0.1049	0.0047
(Europe , East Asia)	0.0072	0.0046	0.010
(GIH , East Asia)	0.0596	0.0296	0.0068
(Europe , GIH)	0.4696	0.2437	0.0032

Table S9: Randomization test (Test 1) p-values for f-statistics for GIH (Two-tailed: Group 1 different from Group 2, One Tail: Group 1 > Group 2, where the sets compared are (Group 1, Group Ref) versus (Group 2, Group Ref)), using f-statistics

		Sets of		Analysis A			
	Popula	tions Compared	p (Two-tailed)	p (One Tail)	Difference		
c	(GIH, Afr)	(Europe, Afr)	0.2311	0.1137	0.0040		
H vs pea	(GIH, YRI)	(Europe, YRI)	0.2363	0.1149	0.0040		
GIF	(GIH, LWK)	(Europe, LWK)	0.1839	0.0894	0.0046		
ш	(GIH <i>,</i> MKK)	(Europe, MKK)	0.2927	0.1464	0.0035		
st	(GIH, Afr)	(East Asia, Afr)	0.0263	0.0102	-0.0057		
s Ea ian	(GIH, YRI)	(East Asia, YRI)	0.0247	0.0102	-0.0059		
IH v Asi	(GIH, LWK)	(East Asia, LWK)	0.0505	0.0222	-0.0049		
ŋ	(GIH <i>,</i> MKK)	(East Asia, MKK)	0.0182	0.0082	-0.0062		



Figure S1 Box plot of the D-statistics for Analyses A and B for the set (Afr, X), where X was any of the non-African populations, CEU or TSI (Europeans, green), CHB or JPT (East Asians, blue), or GIH (South Asian, pink). The red line indicates D = 0.

Analysis A



**Figure S2**: Box plot of the *f*-statistics for Analysis A for the set (Afr, X), where X was any of the non-African populations, CEU or TSI (Europeans, green), CHB or JPT (East Asians, blue), or GIH (South Asian, pink). The red line indicates *f* = 0.

Analysis A



**Figure S3** Box plot of the D-statistics for Analyses A and B comparing East Asians and Europeans. The left partition shows the D-statistics comparing to African individuals (blue and green), while the right partition shows comparisons between non-Africans within (yellow) and between (purple) regional groups. The red line indicates D = 0. *Afr* denotes Africans, *Eur* Europeans, and *E Asn* East Asians.

Analysis A



**Figure S4** Box plot of the *f*-statistics for Analysis A comparing East Asians and Europeans. The left partition shows the *f*-statistics comparing to African individuals (blue and green), while the right partition shows comparisons between non-Africans within (yellow) and between (purple) regional groups. The red line indicates *f* = 0. *Afr* denotes Africans, *Eur* Europeans, and *E Asin* East Asians.



Analysis B



**Figure S5** Box plot of the D-statistics for Analyses A and B showing how the South Asian population GIH compares to other non-African populations. The left partition shows the D-statistics comparing each of the three non-African regional groups to African individuals (blue, green, and pink), while the right partition shows comparisons of East Asians (blue) or Europeans (green) to the GIH individuals. The red line indicates D = 0.

Analysis A



**Figure S6** Box plot of the *f*-statistics for Analysis A showing how the South Asian population GIH compares to other non-African populations. The left partition shows the *f*-statistics comparing each of the three non-African regional groups to African individuals (blue, green, and pink), while the right partition shows comparisons of East Asians (blue) or Europeans (green) to the GIH individuals. The red line indicates f = 0.



**Figure S7** Box plot of the D-statistics for Analyses A and B showing the differences between the three African populations, YRI, LWK, and MKK. The left partition shows the D-statistics calculated for East Asians compared to each of the African populations separately. The right partition shows the D-statistics calculated when comparing the different African populations directly to each other. The red line indicates D = 0. The blue-green color shows the comparisons with MKK. The tan colors show the comparisons without MKK.



**Figure S8** Box plot of the *f*-statistics for Analysis A showing the differences between the three African populations, YRI, LWK, and MKK. The left partition shows the *f*-statistics calculated for East Asians compared to each of the African populations separately. The right partition shows the *f*-statistics calculated when comparing the different African populations directly to each other. The red line indicates f = 0. The blue-green color shows the comparisons with MKK. The tan colors show the comparisons without MKK.