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Upper Palaeolithic genomes reveal deep roots of modern Eurasians

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We extend the scope of European palaeogenomics by sequencing the genomes of Late Upper Palaeolithic (13,300 years old, 1.4-fold coverage) and Mesolithic (9,700 years old, 15.4-fold) males from western Georgia in the Caucasus and a Late Upper Palaeolithic (13,700 years old, 9.5-fold) male from Switzerland. While we detect Late Palaeolithic–Mesolithic genomic continuity in both regions, we find that Caucasus hunter-gatherers (CHG) belong to a distinct ancient clade that split from western hunter-gatherers ~45 kya, shortly after the expansion of anatomically modern humans into Europe and from the ancestors of Neolithic farmers ~25 kya, around the Last Glacial Maximum. CHG genomes significantly contributed to the Yamnaya steppe herders who migrated into Europe ~3,000 BC, supporting a formative Caucasus influence on this important Early Bronze age culture. CHG left their imprint on modern populations from the Caucasus and also central and south Asia possibly marking the arrival of Indo-Aryan languages.

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Ancient genomes from Eurasia have revealed three ancestral populations that contributed to contemporary Europeans in varying degrees¹. Mesolithic individuals, sampled from Spain all the way to Hungary^{1–3}, belong to a relatively homogenous group, termed western hunter-gatherers (WHG). The expansion of early farmers (EF) out of the Levant during the Neolithic transition led to major changes in the European gene pool, with almost complete replacement in the south and increased mixing with local WHG further north^{1–5}. Finally, a later wave originating with the Early Bronze Age Yamnaya from the Pontic steppe, carrying partial ancestry from ancient North Eurasians (ANE) and ancestry from a second, undetermined source, arrived from the east, profoundly changing populations and leaving a cline of admixture in Eastern and Central Europe^{1,3,6}. This view, which was initially based on a handful of genomes, was recently confirmed by extensive surveys of Eurasian samples from the Holocene^{5,7}.

Here, we extend our view of the genetic makeup of early Europeans by both looking further back in time and sampling from the crossroads between the European and Asian continents. We sequenced a Late Upper Palaeolithic ('Satsurblia' from Satsurblia cave, 1.4-fold coverage) and a Mesolithic genome ('Kotias' from Kotias Klde cave, 15.4-fold) from Western Georgia, at the very eastern boundary of Europe. We term these two individuals Caucasus hunter-gatherers (CHG). To extend our overview of WHG to a time depth similar to the one available for our samples from the Caucasus, we also sequenced a western European Late Upper Palaeolithic genome, 'Bichon' (9.5-fold) from Grotte du Bichon, Switzerland. These new genomes, together with already published data, provide us with a much-improved geographic and temporal coverage of genetic diversity across Europe after the Last Glacial Maximum (LGM)⁸. We show that CHG belong to a new, distinct ancient clade that split from WHG ~45 kya and from Neolithic farmer ancestors ~25 kya. This clade represents the previously undetermined source of ancestry to the Yamnaya, and contributed directly to modern populations from the Caucasus all the way to Central Asia.

Results

Samples, sequencing and authenticity. Recent excavations of Satsurblia cave in Western Georgia yielded a human right temporal bone, dated to the Late Upper Palaeolithic 13,132–13,380 cal. BP. Following the approach of Gamba *et al.*³, extractions from the dense part of the petrous bone yielded sequencing libraries comprising 13.8% alignable human sequence which were used to generate 1.4-fold genome coverage. A molar tooth sampled from a later Mesolithic (9,529–9,895 cal. BP) burial in Kotias Klde, a rockshelter also in Western Georgia showed excellent preservation, with endogenous human DNA content of 76.9%. This was sequenced to 15.4-fold genome coverage. Grotte du Bichon is a cave situated in the Swiss Jura Mountains where a skeleton of a young male of Cro-magnon type was found and dated to the late Upper Palaeolithic 13,560–13,770 cal. BP (for further details on the archaeological context see Supplementary Note 1). A petrous bone sample extraction from this also gave excellent endogenous content at 71.5% and was sequenced to 9.5-fold coverage. The sequence data from each genome showed sequence length and nucleotide misincorporation patterns which were indicative of post-mortem damage and contamination estimates, based on X chromosome and mitochondrial DNA tests (Supplementary Note 2), were <1%, comparable to those found in other ancient genomes^{2,3,8}.

Continuity across the Palaeolithic–Mesolithic boundary. Kotias and Satsurblia, the two CHG, are genetically different from all

other early Holocene (that is, Mesolithic and Neolithic) ancient genomes^{1–6,8–10}, while Bichon is similar to other younger WHG. The distinctness of CHG can be clearly seen on a principal component analysis (PCA) plot¹¹ loaded on contemporary Eurasian populations¹, where they fall between modern Caucasian and South Central Asian populations in a region of the graph separated from both other hunter-gatherer and EF samples (Fig. 1a). Clustering using ADMIXTURE software¹² confirms this view, with CHG forming their own homogenous cluster (Fig. 1b). The close genetic proximity between Satsurblia and Kotias is also formally supported by *D*-statistics¹³, indicating the two CHG genomes form a clade to the exclusion of other pre-Bronze Age ancient genomes (Supplementary Table 2; Supplementary Note 3), suggesting continuity across the Late Upper Palaeolithic and Mesolithic periods. This result is mirrored in western Europe as Bichon is close to other WHG in PCA space (Fig. 1a) and outgroup f_3 analysis (Supplementary Fig. 1), belongs to the same cluster as other WHG in ADMIXTURE analysis (Fig. 1b), and forms a clade with other WHG to the exclusion of other ancient genomes based on *D*-statistics (Supplementary Table 3; Supplementary Note 3). Thus, these new data indicate genomic persistence between the Late Upper Palaeolithic and Mesolithic both within western Europe and, separately, within the Caucasus.

Deep coalescence of early Holocene lineages. The geographical proximity of the Southern Caucasus to the Levant begs the question of whether CHG might be related to early Neolithic farmers with Near Eastern heritage. To address this question formally we reconstructed the relationship among WHG, CHG and EF using available high-quality ancient genomes^{1,3}. We used outgroup f_3 -statistics¹⁴ to compare the three possible topologies, with the correct relationship being characterized by the largest amount of shared drift between the two groups that form a clade with respect to the outgroup (Fig. 2a; Supplementary Note 4). A scenario in which the population ancestral to both CHG and EF split from WHG receives the highest support, implying that CHG and EF form a clade with respect to WHG. We can reject a scenario in which CHG and WHG form a distinct clade with respect to EF. The known admixture of WHG with EF^{1,3–5} implies that some shared drift is found between WHG and EF with respect to CHG, but this is much smaller than the shared drift between CHG and EF. Thus, WHG split first, with CHG and EF separating only at a later stage.

We next dated the splits among WHG, CHG and EF using a coalescent model implemented with G-PhoCS¹⁵ based on the high-coverage genomes in our data set (Fig. 2b for a model using the German farmer Stuttgart¹ to represent EF; and Supplementary Table 5 for models using the Hungarian farmer NE1 (ref. 3)) and taking advantage of the mutation rate recently derived from Ust'-Ishim¹⁰. G-Phocs dates the split between WHG and the population ancestral to CHG and EF at ~40–50 kya (range of best estimates depending on which genomes are used; see Supplementary Table 5 and Supplementary Note 5 for details), implying that they diverged early on during the colonisation of Europe¹⁶, and well before the LGM. On the WHG branch, the split between Bichon and Loschbour¹ is dated to ~16–18 kya (just older than the age of Bichon), implying continuity in western Europe, which supports the conclusions from our previous analyses. The split between CHG and EF is dated at ~20–30 kya emerging from a common basal Eurasian lineage¹ (Supplementary Fig. 2) and suggesting a possible link with the LGM, although the broad confidence intervals require some caution with this interpretation. In any case, the sharp genomic distinctions between these post-LGM populations

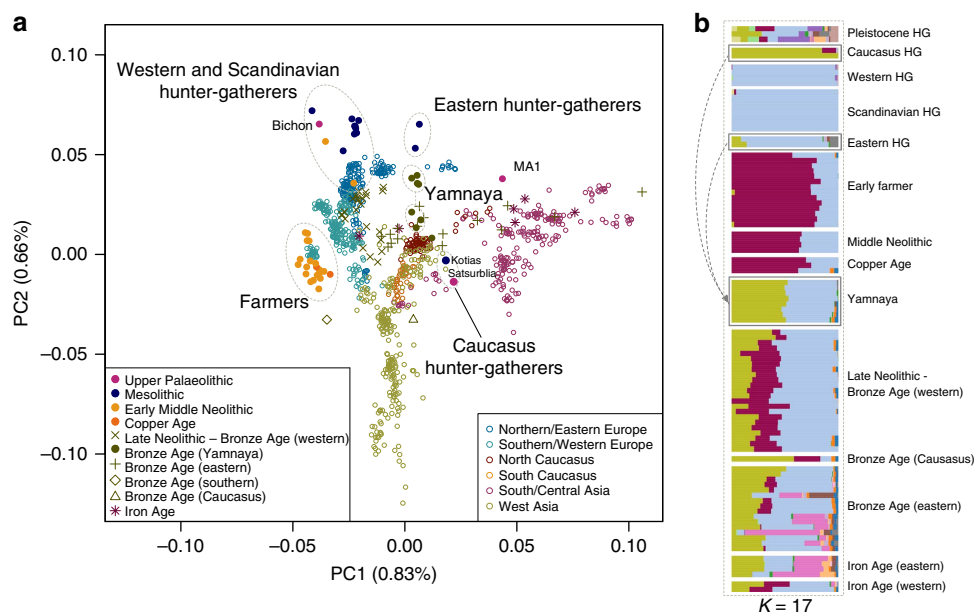


Figure 1 | Genetic structure of ancient Europe. (a). Principal component analysis. Ancient data from Bichon, Kotias and Satsurblia genomes were projected¹¹ onto the first two principal components defined by selected Eurasians from the Human Origins data set¹. The percentage of variance explained by each component accompanies the titles of the axes. For context we included data from published Eurasian ancient genomes sampled from the Late Pleistocene and Holocene where at least 200 000 SNPs were called^{1–7,9} (Supplementary Table 1). Among ancients, the early farmer and western hunter-gatherer (including Bichon) clusters are clearly identifiable, and the influence of ancient north Eurasians is discernible in the separation of eastern hunter-gatherers and the Upper Palaeolithic Siberian sample MA1. The two Caucasus hunter-gatherers occupy a distinct region of the plot suggesting a Eurasian lineage distinct from previously described ancestral components. The Yamnaya are located in an intermediate position between CHG and EHG. (b). ADMIXTURE ancestry components¹² for ancient genomes ($K=17$) showing a CHG component (Kotias, Satsurblia) which also segregates in the Yamnaya and later European populations.

contrasts with the comparative lack of differentiation between the earlier Eurasian genomes, for example, as visualized in the ADMIXTURE analysis (Fig. 1a), and it seems likely that this structure emerged as a result of ice age habitat restriction. Like EF, but in contrast to WHG, CHG carry a variant of the *SLC24A5* gene¹⁷ associated with light skin colour (rs1426654, see Supplementary Note 6). This trait, which is believed to have risen to high frequency during the Neolithic expansion¹⁸, may thus have a relatively long history in Eurasia, with its origin probably predating the LGM.

A partial genome from a 24,000-year-old individual (MA1) from Mal'ta, Siberia⁶ had been shown to be divergent from other ancient samples and was shown by Lazaridis *et al.*¹, using f_4 statistics, to have more shared alleles with nearly all modern Europeans than with an EF genome. This allowed inference of an ANE component in European ancestry, which was subsequently shown to have an influence in later eastern hunter-gatherers and to have spread into Europe via an incursion of Steppe herders beginning ~4,500 years ago^{5,7}. Several analyses indicate that CHG genomes are not a subset of this ANE lineage. First, MA1 and CHG plot in distinct regions of the PCA and also have very different profiles in the ADMIXTURE analysis (Fig. 1). Second, when we test if CHG shows any evidence of excess allele sharing with MA1 relative to WHG using tests of the form $D(\text{Yoruba}, \text{CHG}; \text{MA1}, \text{WHG})$ no combinations were significantly positive (Supplementary Table 6). Last, we also tested whether the ancestral component inferred in modern Europeans from MA1 was distinct from any that may have been donated from CHG using tests of the form $D(\text{Yoruba}, \text{MA1}; \text{CHG}, \text{modern North European population})$ (Supplementary Table 7). All northern Europeans showed a significant sharing of alleles with MA1 separate to any they shared with CHG.

WHG and CHG are the descendants of two ancient populations that appear to have persisted in Europe since the mid Upper Palaeolithic and survived the LGM separately. We looked at runs of homozygosity (ROH; Fig. 3) which inform on past population size^{3,19,20}. Both WHG and CHG have a high frequency of ROH and in particular, the older CHG, Satsurblia, shows signs of recent consanguinity, with a high frequency of longer (>4 Mb) ROH. In contrast, EF are characterised by lower frequency of ROH of all sizes, suggesting a less constricted population history^{20,21}, perhaps associated with a more benign passage through the LGM than the more northern populations (see Supplementary Note 7 for further details).

Caucasus hunter-gatherer contribution to subsequent populations.

We next explored the extent to which Bichon and CHG contributed to contemporary populations using outgroup $f_3(\text{African}; \text{modern}, \text{ancient})$ statistics, which measure the shared genetic history between an ancient genome and a modern population since they diverged from an African outgroup. Bichon, like younger WHG, shows strongest affinity to northern Europeans (Supplementary Fig. 3), while contemporary southern Caucasus populations are the closest to CHG (Fig. 4a and Supplementary Fig. 3), thus implying a degree of continuity in both regions stretching back at least 13,000 years to the late Upper Palaeolithic. Continuity in the Caucasus is also supported by the mitochondrial and Y chromosomal haplogroups of Kotias (H13c and J2a, respectively) and Satsurblia (K3 and J), which are all found at high frequencies in Georgia today^{22–24} (Supplementary Note 8).

EF share greater genetic affinity to populations from southern Europe than to those from northern Europe with an inverted

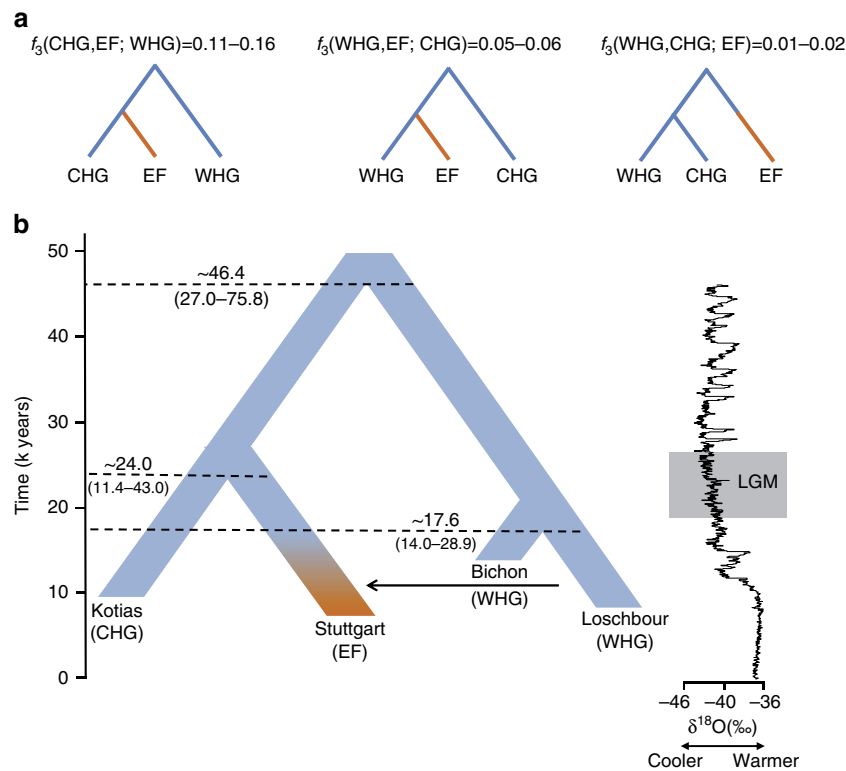


Figure 2 | The relationship between Caucasus hunter-gatherers (CHG), western hunter-gatherers and early farmers. (a). Alternative phylogenies relating western hunter-gatherers (WHG), CHG and early farmers (EF, highlighted in orange), with the appropriate outgroup f_3 -statistics. (b). The best supported relationship among CHG (Kotias), WHG (Bichon, Loschbour), and EF (Stuttgart), with split times estimates using G-Phocs¹⁵. Oxygen 18 values (per mile) from the NGRIP core provide the climatic context; the grey box shows the extent of the Last Glacial Maximum (LGM).

pattern for WHG^{1–5}. Surprisingly, we find that CHG influence is stronger in northern than Southern Europe (Fig. 4a and Supplementary Fig. 3A) despite the closer relationship between CHG and EF compared with WHG, suggesting an increase of CHG ancestry in Western Europeans subsequent to the early Neolithic period. We investigated this further using D -statistics of the form $D(\text{Yoruba}, \text{Kotias}; \text{EF}, \text{modern Western European population})$, which confirmed a significant introgression from CHG into modern northern European genomes after the early Neolithic period (Supplementary Fig. 4).

CHG origins of migrating Early Bronze Age herders. We investigated the temporal stratigraphy of CHG influence by comparing these data to previously published ancient genomes. We find that CHG, or a population close to them, contributed to the genetic makeup of individuals from the Yamnaya culture, which have been implicated as vectors for the profound influx of Pontic steppe ancestry that spread westwards into Europe and east into central Asia with metallurgy, horseriding and probably Indo-European languages in the third millennium BC^{5,7}. CHG ancestry in these groups is supported by ADMIXTURE analysis (Fig. 1b) and admixture f_3 -statistics^{14,25} (Fig. 5), which best describe the Yamnaya as a mix of CHG and Eastern European hunter-gatherers. The Yamnaya were semi-nomadic pastoralists, mainly dependent on stock-keeping but with some evidence for agriculture, including incorporation of a plow into one burial²⁶. As such it is interesting that they lack an ancestral coefficient of the EF genome (Fig. 1b), which permeates through western European Neolithic and subsequent agricultural populations. During the Early Bronze Age, the Caucasus was in communication with the steppe, particularly via the Maikop

culture²⁷, which emerged in the first-half of the fourth millennium BC. The Maikop culture predated and, possibly with earlier southern influences, contributed to the formation of the adjacent Yamnaya culture that emerged further to the north and may be a candidate for the transmission of CHG ancestry. In the ADMIXTURE analysis of later ancient genomes (Fig. 1b) the Caucasus component gives a marker for the extension of Yamnaya admixture, with substantial contribution to both western and eastern Bronze Age samples. However, this is not completely coincident with metallurgy; Copper Age genomes from Northern Italy and Hungary show no contribution; neither does the earlier of two Hungarian Bronze Age individuals.

Modern impact of CHG ancestry. In modern populations, the impact of CHG also stretches beyond Europe to the east. Central and South Asian populations received genetic influx from CHG (or a population close to them), as shown by a prominent CHG component in ADMIXTURE (Supplementary Fig. 5; Supplementary Note 9) and admixture f_3 -statistics, which show many samples as a mix of CHG and another South Asian population (Fig. 4b; Supplementary Table 9). It has been proposed that modern Indians are a mixture of two ancestral components, an Ancestral North Indian component related to modern West Eurasians and an Ancestral South Indian component related more distantly to the Onge²⁵; here Kotias proves the majority best surrogate for the former^{28,29} (Supplementary Table 10). It is estimated that this admixture in the ancestors of Indian populations occurred relatively recently, 1,900–4,200 years BP, and is possibly linked with migrations introducing Indo-European languages and Vedic religion to the region²⁸.

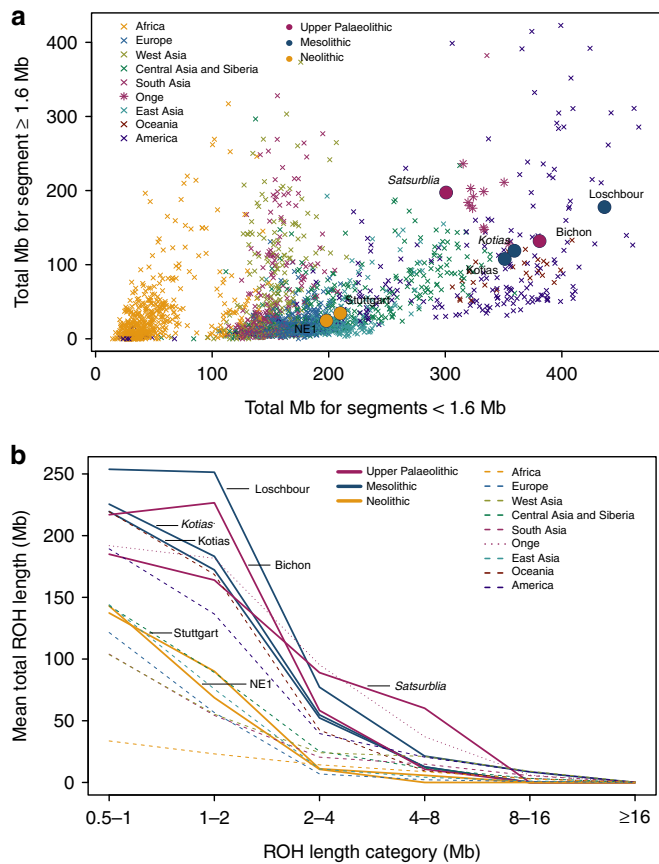


Figure 3 | Distribution of ROH. (a) The total length of short ROH (<1.6 Mb) plotted against the total length of long ROH (≥1.6 Mb) and (b) mean total ROH length for a range of length categories. ROH were calculated using a panel of 199,868 autosomal SNPs. For Kotias we analysed both high-coverage genotypes and genotypes imputed from downsampled data (marked in italics; see Supplementary Information). Diploid genotypes imputed from low-coverage variant calls were used for Satsurbilia and high-coverage genotypes were used for all other samples. A clear distinction is visible between either WHG and CHG who display an excess of shorter ROH, akin to modern Oceanic and Onge populations, and EF who resemble other populations with sustained larger ancestral population sizes.

Discussion

Given their geographic origin, it seems likely that CHG and EF are the descendants of early colonists from Africa who stopped south of the Caucasus, in an area stretching south to the Levant and possibly east towards Central and South Asia. WHG, on the other hand, are likely the descendants of a wave that expanded further into Europe. The separation of these populations is one that stretches back before the Holocene, as indicated by local continuity through the Late Palaeolithic/Mesolithic boundary and deep coalescence estimates, which date to around the LGM and earlier. Several analyses show that CHG are distinct from another inferred minor ancestral population, ANE, making them a divergent fourth strand of European ancestry that expands the model of the human colonization of that continent.

The separation between CHG and both EF and WHG ended during the Early Bronze Age when a major ancestral component linked to CHG was carried west by migrating herders from the Eurasian Steppe. The foundation group for this seismic change was the Yamnaya, who we estimate to owe half of their ancestry to CHG-linked sources. These sources may be linked to the

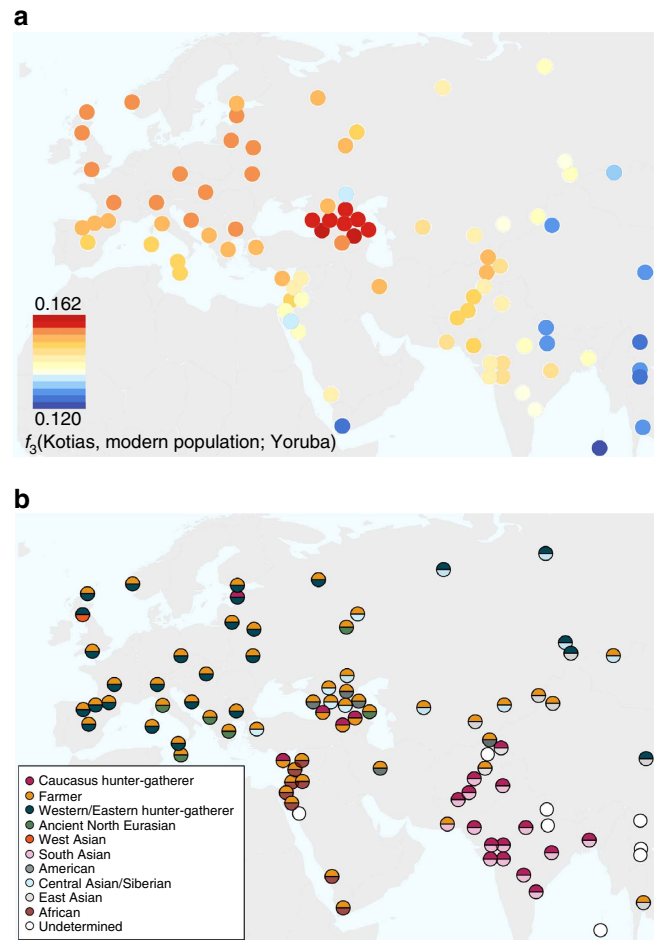


Figure 4 | The relationship of Caucasus hunter-gatherers to modern populations. (a). Genomic affinity of modern populations¹ to Kotias, quantified by the outgroup f_3 -statistics of the form $f_3(\text{Kotias, modern population; Yoruba})$. Kotias shares the most genetic drift with populations from the Caucasus with high values also found for northern Europe and central Asia. (b). Sources of admixture into modern populations: semicircles indicate those that provide the most negative outgroup f_3 statistic for that population. Populations for which a significantly negative statistic could not be determined are marked in white. Populations for which the ancient Caucasus genomes are best ancestral approximations include those of the Southern Caucasus and interestingly, South and Central Asia. Western Europe tends to be a mix of early farmers and western/eastern hunter-gatherers while Middle Eastern genomes are described as a mix of early farmers and Africans.

Maikop culture, which predated the Yamnaya and was located further south, closer to the Southern Caucasus. Through the Yamnaya, the CHG ancestral strand contributed to most modern European populations, especially in the northern part of the continent.

Finally, we found that CHG ancestry was also carried east to become a major contributor to the Ancestral North Indian component found in the Indian subcontinent. Exactly when the eastwards movement occurred is unknown, but it likely included migration around the same time as their contribution to the western European gene pool and may be linked with the spread of Indo-European languages. However, earlier movements associated with other developments such as that of cereal farming and herding are also plausible.

The discovery of CHG as a fourth ancestral component of the European gene pool underscores the importance of a dense

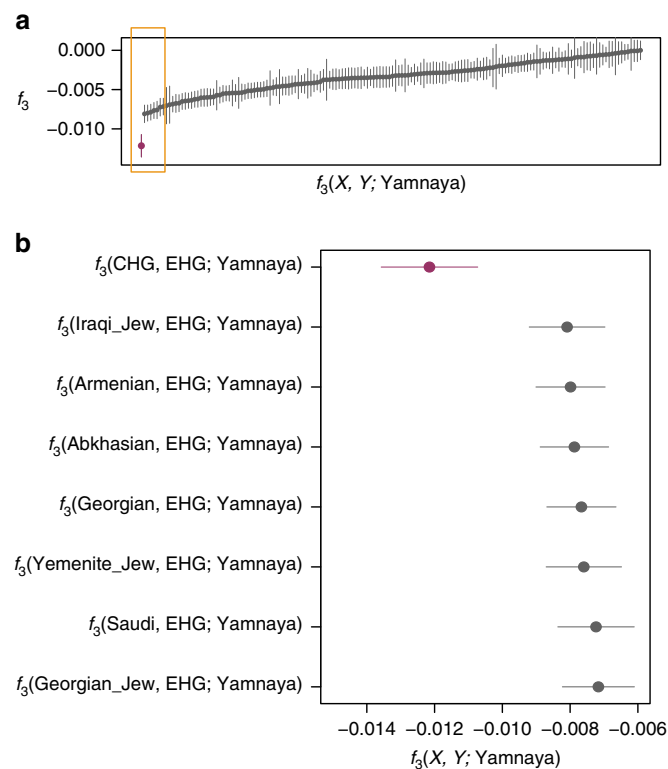


Figure 5 | Lowest admixture f_3 -statistics of the form $f_3(X, Y; \text{Yamnaya})$.

These statistics represent the Yamnaya as a mix of two populations with a more negative result signifying the more likely admixture event. (a) All negative statistics found for the test $f_3(X, Y; \text{Yamnaya})$ with the most negative result $f_3(\text{CHG, EHG; Yamnaya})$ highlighted in purple. Lines bisecting the points show the standard error. (b) The most significantly negative statistics which are highlighted by the yellow box in a. Greatest support is found for Yamnaya being a mix of Caucasus hunter-gatherers (CHG) and Russian hunter-gatherers who belong to an eastern extension of the WHG clade (EHG).

geographical sampling of human palaeogenomes, especially among diverse geographical regions. Its separation from other European ancestral strands ended dramatically with the extensive population, linguistic and technological upheavals of the Early Bronze Age resulting in a wide impact of this ancestral strand on contemporary populations, stretching from the Atlantic to Central and South Asia.

Methods

Sample preparation and DNA sequencing. DNA was extracted from three samples; two from Georgia (Kotias and Satsurblia) and one from Switzerland (Bichon; Supplementary Fig. 6). Sample preparation, DNA extraction and library construction were carried out in dedicated ancient DNA facilities at Trinity College Dublin (Kotias and Satsurblia) and the University of York, England (Bichon). DNA was extracted from Kotias and Satsurblia following a silica column based protocol³ based on ref. 30 and libraries were prepared and amplified with AccuPrime Pfx Supermix (Life Technology), using a modified version of ref. 31 as outlined in ref. 3. For the ancient Swiss sample Bichon, DNA was extracted following³² and libraries were built as described above with the exception that enzymatic end-repair was arrested using heat inactivation rather than a silica-column purification step^{33,34}. Libraries were first screened to assess their human DNA content on an Illumina MiSeq platform at TrinSeq, Dublin using 50 base pair (bp) single-end sequencing (Supplementary Table 11). Selected libraries were further sequenced on a HiSeq 2000 platform using 100 bp single-end sequencing (Supplementary Table 12).

Sequence processing and alignment. To reduce the effects of post-indexing contamination, raw reads were retained if the Hamming distance for the observed index was within 1 base of the expected index. Adapter sequences were trimmed from the 3' ends of reads using cutadapt version 1.3 (ref. 35), requiring an overlap of

1 bp between the adapter and the read. As ancient DNA damage is more apparent at the ends of sequences^{36,37}, the first and last two bp of all reads from the deep sequencing phase of analysis (Supplementary Table 12) were removed using SeqTK (<https://github.com/lh3/seqtk>). A minimum read length of 30 bp was imposed.

Sequences were aligned using Burrows-Wheeler Aligner (BWA) version 0.7 (ref. 38), with the seed region disabled, to the GRCh37 build of the human genome with the mitochondrial sequence replaced by the revised Cambridge reference sequence (NCBI accession number NC_012920.1). Sequences from the same sample were merged using Picard MergeSamFiles (<http://picard.sourceforge.net/>) and duplicate reads were removed using SAMtools version 0.1.19 (ref. 39). Average depth of coverage was calculated using genome analysis toolkit (GATK) Depth of Coverage and indels were realigned using RealignerTargetCreator and IndelRealigner from the same suite of tools⁴⁰. Reads with a mapping quality of at least 30 were retained using SAMtools³⁹, and mapDamage 2.0 (ref. 41) was used with default parameters to downscale the quality scores of likely damaged bases, reducing the influence of nucleotide misincorporation on results. Only data from the deep sequencing phase of the project (100 bp single-end sequencing on a HiSeq 2,000) were used in the subsequent analyses.

Authenticity of results. Rigorous measures were taken during laboratory work in an effort to minimize DNA contamination³ and negative controls were processed in parallel with samples. The authenticity of the data was further assessed *in silico* in a number of ways. The data were examined for the presence of short average sequence length and nucleotide misincorporation patterns which are characteristic of aDNA^{36,37} (Supplementary Figs 7 and 8). The degree of mitochondrial DNA contamination^{3,42} (Supplementary Table 13) and X chromosome contamination in male samples³ (Supplementary Tables 14–16) was also assessed (for further details see Supplementary Information).

Molecular sex and uniparental haplogroups. Genetic sex was determined by examining the ratio of Y chromosome reads to reads aligning to both sex chromosomes⁴³ (Supplementary Table 17). Mitochondrial haplogroups were assigned following⁴ with coverage determined using GATK Depth of Coverage⁴⁰ (Supplementary Table 18). YFitter⁴⁴, which employs a maximum likelihood based approach, was used to determine Y chromosome haplogroups for our ancient male samples (Supplementary Table 19).

Merging ancient data with modern reference data set. Genotype calls from Kotias, Satsurblia, Bichon and selected Eurasian samples (Supplementary Table 1; Supplementary Note 10) were merged with modern genotype calls from the Human Origins data set¹ using PLINK⁴⁵. This data set was first filtered to exclude genotypes which had a minor allele frequency of zero in the modern populations, non-autosomal sites and modern populations with less than four individuals. Genotypes where neither allele was consistent with the GRCh37 orientation of the human genome were also removed.

For ancient samples with $> 8 \times$ genome-wide coverage (namely Kotias, Bichon, Ust'-Ishim, Loschbour, Stuttgart, NE1 and BR2 (Supplementary Table 1)) genotypes were determined using GATK Unified Genotyper⁴⁰. Genotypes were called at single-nucleotide polymorphism (SNP) positions observed in the Human Origins data set using sequencing data with a base quality ≥ 30 , depth ≥ 8 and genotype quality ≥ 20 . The resulting VCF files were converted to PLINK format using VCFtools⁴⁶.

For lower coverage samples genotypes were called at positions that overlapped with the Human Origins data set using GATK Pileup⁴⁰. Bases were required to have a minimum quality of 30 and all triallelic SNPs were discarded. For SNP positions with more than one base call, one allele was randomly chosen with a probability equal to the frequency of the base at that position. This allele was duplicated to form a homozygous diploid genotype which was used to represent the individual at that SNP position⁴⁷. This merged data set was used for PCA, ADMIXTURE, f_3 -statistics, D -statistics and ROH analysis.

Population genetic analyses. PCA was performed by projecting selected ancient Eurasian data onto the first two principal components defined by a subset of the filtered Human Origins data set (Fig. 1a). This analysis was carried out using EIGENSOFT 5.0.1 smartpca¹¹ with the lsqproject option on and the outlier removal option off. One SNP from each pair in linkage disequilibrium with $r^2 > 0.2$ was removed⁴⁷.

A clustering analysis was performed using ADMIXTURE version 1.23 (ref. 12). Genotypes were restricted to those that overlapped with the SNP capture panel described in ref. 5. Single-nucleotide polymorphisms in linkage disequilibrium were thinned using PLINK (v1.07)⁴⁵ with parameters—*indep-pairwise* 200 25 0.5 (ref. 5) resulting in a set of 229,695 SNPs for analysis. Clusters (K) (2–20) were explored using 10 runs with fivefold cross-validation at each K with different random seeds (Supplementary Fig. 5). The minimal cross-validation error was found at $K = 17$, but the error already starts plateauing from roughly $K = 10$, implying little improvement from this point onwards. (Supplementary Fig. 9).

D -statistics¹³ and f_3 -statistics^{14,25} were used to formally assess the relationships between samples. Statistics were computed using the qpDstat (D -statistics) and 3PopTest (f_3 -statistics) programs from the ADMIXTOOLS package¹⁴. Significance

was assessed using a block jackknife over 5 cM chunks of the genome¹⁴ and statistics were considered significant if their Z-score was of magnitude greater than 3 (ref. 25) corresponding approximately to a P value < 0.001. For f_3 -statistics where the test population was ancient the inbreed:YES option was used.

Dating split times using G-PhoCS. A coalescent model implemented with G-PhoCS¹⁵ was used to date the split among WHG, CHG and EF. Both a tree with only ancient genomes and a tree with an African San Pygmy⁴⁸ as the outgroup were considered (see Supplementary Information for further details).

Runs of homozygosity. Examination of ROH requires dense diploid genotypes. Imputation was used to maximise the information content of our most ancient sample, Satsurblia, following the procedure described in ref. 3. Implementing a genotype probability threshold of 0.99 (Supplementary Fig. 10), imputed data for Satsurblia and downsampled-Kotias (see Supplementary Information) was merged with high confidence diploid calls for selected ancient samples (namely Bichon, Loschbour, NE1, Stuttgart and Kotias) as well as with SNP data from modern samples using PLINK⁴⁵. This resulted in 199,868 overlapping high-quality diploid loci for ROH analysis which was carried out using PLINK⁴⁵ as described in ref. 3.

Phenotypes of interest. Genes which have been associated with particular phenotypes in modern populations were examined, including some loci which have been subject to selection in European populations (Supplementary Tables 20–23). We called genotypes in Bichon, Kotias and Satsurblia using GATK Unified Genotyper⁴⁰. For each position under investigation we only called alleles which were present in the 1,000 Genomes data set⁴⁹, using bases with a quality ≥ 30 in positions with a depth ≥ 4 . Because of the low average coverage of Satsurblia (1.44 \times) we also used imputed genotypes for this sample (see above) imposing a genotype probability cut-off of 0.85 (ref. 3). We used the 8-plex⁵⁰ and Hirisplex⁵¹ prediction models to predict hair, eye and skin colour for our samples. Other loci investigated are discussed in the Supplementary Information.

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Author contributions

D.G.B., A.M. and R.P. supervised the study. E.R.J., D.G.B. and A.M. designed the study and wrote the manuscript with contributions from all authors. E.R.J. prepared sequencing libraries and processed and analysed ancient DNA with support from S.C., R.M., R.L.M., G.G.-F., L.C. and C.G. G.G.-F. and M.H. prepared sequencing libraries for Bichon. E.R.J. performed population genetics analyses with help from D.G.B., A.M. and M.C. Coalescent modelling was performed by V.S., A.E., M.G.-L. and A.M. V.S. carried out ADMIXTURE analysis. R.P., T.M., O.B.-Y., W.M., A.B., Z.M., N.J., T.F.G.H. and D.L. provided archaeological samples and input about the archaeological samples.

Additional information

Accession codes: Alignment data are available through the European Nucleotide Archive (ENA) under the project accession number PRJEB11364. <http://www.ebi.ac.uk/ena/data/view/PRJEB11364>.

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