

# Population genomics of the Viking world

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The maritime expansion of Scandinavian populations during the Viking Age (about AD 750–1050) was a far-flung transformation in world history<sup>1,2</sup>. Here we sequenced the genomes of 442 humans from archaeological sites across Europe and Greenland (to a median depth of about 1×) to understand the global influence of this expansion. We find the Viking period involved gene flow into Scandinavia from the south and east. We observe genetic structure within Scandinavia, with diversity hotspots in the south and restricted gene flow within Scandinavia. We find evidence for a major influx of Danish ancestry into England; a Swedish influx into the Baltic; and Norwegian influx into Ireland, Iceland and Greenland. Additionally, we see substantial ancestry from elsewhere in Europe entering Scandinavia during the Viking Age. Our ancient DNA analysis also revealed that a Viking expedition included close family members. By comparing with modern populations, we find that pigmentation-associated loci have undergone strong population differentiation during the past millennium, and trace positively selected loci—including the lactase-persistence allele of *LCT* and alleles of *ANKK1* that are associated with the immune response—in detail. We conclude that the Viking diaspora was characterized by substantial transregional engagement: distinct populations influenced the genomic makeup of different regions of Europe, and Scandinavia experienced increased contact with the rest of the continent.

The events of the Viking Age altered the political, cultural and demographic map of Europe in ways that are evident to this day. Scandinavian diasporas established trade and settlements that stretched from the American continent to the Asian steppe<sup>1</sup>. They exported ideas, technologies, language, beliefs and practices to these lands, developed new socio-political structures and assimilated cultural influences<sup>2</sup>.

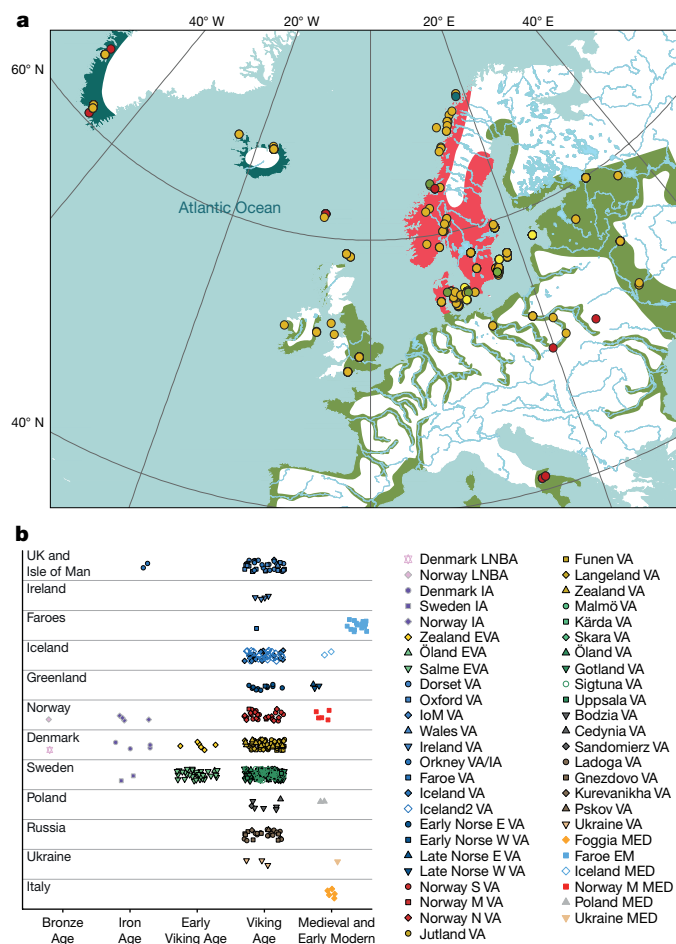
To explore the genomic history of the Viking Age, we shotgun-sequenced DNA extracted from 442 human remains from archaeological sites dating from the Bronze Age (about 2400 BC) to the Early Modern period (about AD 1600) (Fig. 1, Extended Data Fig. 1). The data from these ancient individuals were analysed together with published

data from 3,855 present-day individuals across two reference panels (Supplementary Note 6), and data from 1,118 ancient individuals (Supplementary Table 3).

## Scandinavian ancestry and Viking Age origins

Although Viking Age Scandinavian populations shared a common cultural background, there was no common word for Scandinavian identity at this time<sup>1</sup>. Rather than there being a single ‘Viking world’, a series of interlinked Viking worlds emerged from rapidly growing maritime exploration, trade, war and settlement, following the adoption

A list of affiliations appears at the end of the paper.



**Fig. 1 | Overview of the Viking Age genomic dataset. a**, Map of the Viking World from eighth to eleventh centuries AD, showing geographical location and broad age category of sites with ancient samples newly reported in this study. Age categories of sites (circles) are colour-coded as: dark green, LNBA (2400–500 BC); light green, Iron Age (500 BC–AD 700); yellow, Early Viking Age (AD 700–800); Viking Age (AD 800–1100); Medieval and Early Modern (AD 1100–1600). Red region, area of Viking origins; green region, area of Viking raids, settlement and trade; dark blue region, area of pioneer Viking colonization. **b**, All of the ancient individuals from this study ( $n = 442$ ), and previously published Viking Age samples from Sigtuna<sup>10</sup> and Iceland<sup>18</sup>, categorized on the basis of their spatiotemporal origin. The ancient samples are divided into the following five broad categories: Bronze Age (BA), Iron Age (IA), Early Viking Age (EVA), Viking Age (VA), Medieval (MED) and Early Modern (EM). Random jitter has been added along the x axis in each category to aid visualization. LNBA, Late Neolithic and Bronze Age; Norse W, Norse western settlement; Norse E, Norse eastern settlement; Norway S, southern Norway; Norway N, northern Norway; Norway M, middle Norway.

of deep-sea navigation among coastal populations of Scandinavia and the area around the Baltic Sea<sup>3,4</sup>. Thus, it is unclear to what extent the Viking phenomenon refers to people with a recently shared genetic background or how far population changes accompanied the transition from the Iron Age (500 BC–AD 700) to the Viking Age in Scandinavia.

The Viking Age Scandinavian individuals of our study fall broadly within the diversity of ancient European individuals from the Bronze Age and later (Fig. 2, Extended Data Figs. 2, 3, Supplementary Note 8), but with subtle differences among the groups that indicate complex fine-scale structure. For example, many Viking Age individuals from the island of Gotland cluster with Bronze Age individuals from the Baltic region, which indicates mobility across the Baltic Sea (Fig. 2, Extended Data Fig. 3). Using  $f_4$ -statistics to contrast genetic affinities with steppe pastoralists and Neolithic farmers, we find that Viking Age

individuals from Norway are distributed in a manner similar to that of earlier Iron Age individuals, whereas many Viking Age individuals from Sweden and Denmark show a greater affinity to Neolithic farmers from Anatolia (Extended Data Fig. 4a). Using the qpAdm program, we find that the majority of groups can be modelled as three-way mixtures of hunter-gatherer, farmer and steppe-related ancestry. The three-way model was rejected for some groups from Sweden, Norway and the Baltic region, which could be fit using four-way models that additionally included either Caucasus hunter-gatherer or East-Asian-related ancestry (Extended Data Figs. 4b, c)—the latter of which is consistent with previously documented gene flow from Siberia<sup>5–7</sup>.

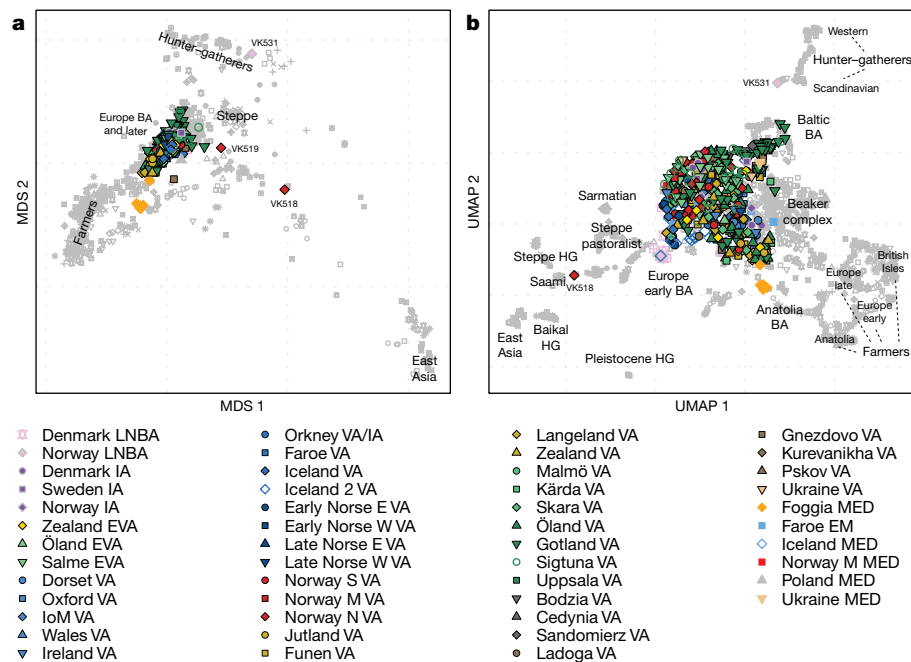
Investigating genetic continuity with Iron Age groups that are temporally more proximate to the Viking Age Scandinavian populations, we find that most Viking Age groups can be fit using a single Iron Age source and broadly fall into two categories: (i) English Iron Age sources (most of the Viking Age individuals from Denmark, as well as populations of the British Isles) and (ii) Scandinavian Iron Age sources (from Norway, Sweden and the Baltic region) (Extended Data Fig. 5a). Notable exceptions are individuals from Kärda in southern Sweden, for whom only early Medieval Longobard individuals from Hungary can be fit as a single source group ( $P > 0.01$ ) (Extended Data Fig. 5a). Groups with poor one-way fits can be modelled by including either additional north-eastern ancestry (for example, Viking Age individuals from Ladoga) or additional southeastern ancestry (for example, Viking Age individuals from Jutland) (Extended Data Fig. 5b). Overall, our analyses suggest that the genetic makeup of Viking Age Scandinavian populations largely derives from ancestry of the preceding Iron Age populations—but these analyses also reveal subtle differences in ancestry and gene flow from both the south and east. These observations are largely consistent with archaeological findings<sup>8,9</sup>.

## Viking Age genetic structure in Scandinavia

To elucidate the fine-scale population structure of Viking Age Scandinavia, we performed genotype imputation on a subset of 298 individuals with sufficient ( $>0.5\times$ ) coverage (289 from this study, along with 9 previously published individuals<sup>10</sup>) and inferred the genomic segments they shared via identity-by-descent with a reference panel of present-day European individuals ( $n = 1,464$ ) (Supplementary Notes 6, 10, 11). Genetic clustering using multidimensional scaling and uniform manifold approximation and projection (UMAP) shows that Viking Age Scandinavian individuals cluster into three groups by geographical origin, with close affinities to their respective present-day counterparts (Fig. 3a, Supplementary Fig. 10.1). Some individuals—particularly those from the island of Gotland in eastern Sweden—have strong affinities with Eastern Europeans; this probably reflects individuals with Baltic ancestry, as clustering with Bronze Age individuals from the Baltic region is evident in the identity-by-state UMAP analysis (Fig. 2b) and through  $f_4$ -statistics (Supplementary Fig. 9.1).

We used ChromoPainter<sup>11</sup> and a reference panel enriched with Scandinavian individuals ( $n = 1,464$ ) (Supplementary Notes 6, 11) to identify long, shared haplotypes and detect subtle population structure (Supplementary Figs. 11.1–11.10). We find ancestry components in Scandinavia with (inexact and indicative) affinities with present-day populations (Supplementary Fig. 11.11), which we refer to as ‘Danish-like’, ‘Swedish-like’, ‘Norwegian-like’ and ‘North Atlantic-like’ (that is, possible individuals from the British Isles entering Scandinavia). The sampling is heavily structured, so these complex results (Supplementary Fig. 11.12) are visualized over time and space (Fig. 4) using spatial interpolation<sup>12</sup> to account for sampling locations and report significant trends (Supplementary Table 11.2) using linear regression (Supplementary Notes 11, 12).

Norwegian-like and Swedish-like components cluster in Norway and Sweden, respectively, whereas Danish-like and North-Atlantic-like components are widespread (Fig. 4, Supplementary Fig. 11.12,



**Fig. 2 | Genetic structure of Viking Age samples. a**, Multidimensional scaling (MDS) of  $n = 1,305$  ancient genomes, on the basis of a pairwise identity-by-state sharing matrix of the Viking Age and other ancient samples (Supplementary Table 3). Outlier individuals with hunter-gatherer (VK531) or Saami-related

ancestry (VK518 and VK519) are highlighted. **b**, UMAP analysis of the same dataset as in **a**, with fine-scale ancestry groups highlighted. HG, hunter-gatherer.

Supplementary Table 6). Unexpectedly, Viking Age individuals from Jutland (Denmark) lack Swedish-like and Norwegian-like genetic components (Supplementary Fig. 11.12). We also find that gene flow within Scandinavia was broadly from south to north, dominated by movement from Denmark into Norway and Sweden (Supplementary Table 11.2).

We identified two ancient individuals from northern Norway (designated VK518 and VK519) with affinities to present-day Saami populations in Norway and Sweden. The VK519 individual probably also had Norwegian-like ancestors, which indicates genetic contacts between Saami groups and other Scandinavian populations.

The genetic data are structured by topographical boundaries rather than by the borders of present-day countries. Thus, the southwestern part of Sweden in the Viking Age is genetically more similar to Viking Age populations of Denmark than to those of central mainland Sweden, probably owing to geographical barriers that prevented gene flow.

We quantified genetic diversity using two measures: conditional nucleotide diversity (Supplementary Note 9) and variation in inferred ancestry on the basis of ChromoPainter results (Extended Data Fig. 6, Supplementary Note 11, Supplementary Fig. 11.13). We also visualized this diversity as the spread of individuals on a multidimensional scaling plot based on a pairwise identity-by-state sharing matrix (Fig. 3b).

Diversity varies markedly from the more-homogeneous inland and northern parts of Scandinavia to the diverse Kattegat (eastern Denmark and western Sweden) and Baltic Sea regions, which suggests an important role for these maritime regions in interaction and trade during the Viking Age. On Gotland, there are many more Danish-like and North-Atlantic-like genetic components (as well as an additional 'Finnish-like' ancestry component) than Swedish-like components, which indicates extensive maritime contacts for Gotland during the Viking Age.

Our results for Gotland and Öland agree with archaeological indications that these were important maritime communities from the Roman period (AD 1–400) onwards<sup>13,14</sup>. A similar pattern is observed on the central Danish islands (such as Langeland) but at a lower level. The data indicate that genetic diversity on the islands increased from the early (about eighth century AD) to the late Viking Age (about tenth

to eleventh centuries AD), which suggests increasing interregional interaction over time. Evidence for genetic structure within Viking Age Scandinavia<sup>2,4,15–17</sup>—with diversity in cosmopolitan centres such as Skara, and trade-oriented islands such as Gotland—highlight the importance of sea routes during this period.

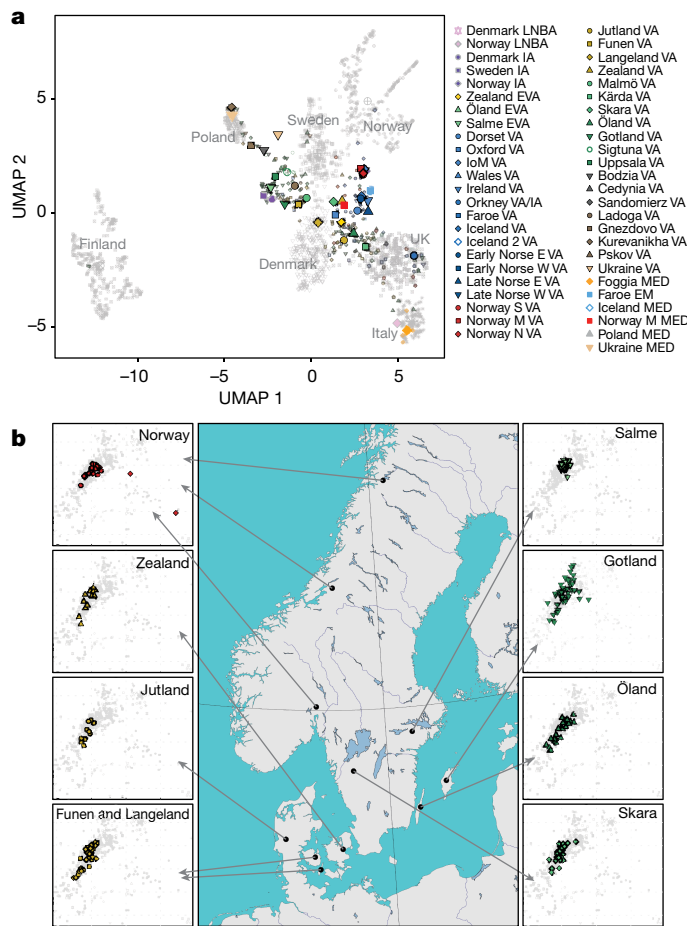
## Viking migrations

Our fine-scale ancestry analyses of genomic data are consistent with patterns documented by historians and archaeologists (Figs. 3, 4, Supplementary Fig. 11.12): eastward movements mainly involved Swedish-like ancestry, whereas individuals with Norwegian-like ancestry travelled to Iceland, Greenland, Ireland and the Isle of Man. The first settlement in Iceland and Greenland also included individuals with North-Atlantic-like ancestry<sup>18,19</sup>. A Danish-like ancestry is seen in present-day England, in accordance with historical records<sup>20</sup>, place names<sup>21</sup>, surnames<sup>22</sup> and modern genetics<sup>23,24</sup>, but Viking Age Danish-like ancestry in the British Isles cannot be distinguished from that of the Angles and Saxons, who migrated in the fifth to sixth centuries AD from Jutland and northern Germany.

Viking Age execution sites in Dorset and Oxford (England) contain North-Atlantic-like ancestry, as well as Danish-like and Norwegian-like ancestries. If these sites represent Viking raiding parties that were defeated and captured<sup>25,26</sup>, then these raids were composed of individuals of different origins. This pattern is also suggested by isotopic data from a warrior cemetery in Trelleborg (Denmark)<sup>27</sup>. Similarly, the presence of Danish-like ancestry in an ancient sample from Gnezdovo in present-day Russia indicates that eastern migrations were not entirely composed of Viking individuals from Sweden.

Our results show that 'Viking' identity was not limited to individuals of Scandinavian genetic ancestry. Two individuals from Orkney who were buried in Scandinavian fashion are genetically similar to present-day Irish and Scottish populations, and are probably the first Pictish genomes published (see 'Evidence for Pictish genomes' in Supplementary Note 11, Supplementary Figs. 11.3, 11.12, 11.14, Supplementary Table 6). Two other individuals from Orkney had 50% Scandinavian





**Fig. 3 | Genetic structure and diversity of ancient samples. a**, UMAP analysis of  $n = 1,624$  ancient and modern Scandinavian individuals, on the basis of the first 10 dimensions of MDS using identity-by-descent segments of imputed individuals. Large symbols indicate median coordinates for each group. **b**, Genetic diversity in major populations of the Scandinavian Viking Age. Plots next to the map show MDS analysis on the basis of a pairwise identity-by-state sharing matrix. Norway denotes all the sites from Norway. The scale is identical for all the plots.

ancestry, and five such individuals were found in Scandinavia. This suggests that Pictish populations may have been integrated into Scandinavian culture by the Viking Age.

### Viking Age gene flow into Scandinavia

Non-Scandinavian ancestry in samples from Denmark, Norway and Sweden agrees with known trading routes (Supplementary Notes 11, 12): for example, Finnish and Baltic ancestry reached modern Sweden (including Gotland), but is absent in most individuals from Denmark and Norway. By contrast, western regions of Scandinavia received ancestry from the British Isles (Supplementary Notes 11, 12). The first evidence of South European ancestry (>50%) in Scandinavia is during the Viking Age in Denmark (for example, individuals VK365 and VK286 from Bogøvej) and southern Sweden (for example, VK442 and VK350 from Öland, and VK265 from Kärda) (Fig. 4, Supplementary Table 6).

### Disappearance from Greenland

From around AD 980 to 1440, southwest Greenland was settled by people of Scandinavian ancestry (probably from Iceland)<sup>28,29</sup>. The fate of these populations in Greenland remains debated, but probable causes of their disappearance are social or economic processes in Europe

(for example, political relations within Scandinavia and changed trading systems) and natural processes, including climatic change<sup>29–31</sup>.

According to our data, the Greenland Norse populations were an admixture between Scandinavians (mostly from Norway) and individuals from the British Isles, similar to the first settlers of Iceland<sup>18</sup>. We see no evidence of long-term inbreeding in the genomes of Greenlandic Norse individuals, although we have only one high-coverage genome from the later period of occupation of the island (Supplementary Note 10, Supplementary Figs. 10.2, 10.3). This result could favour a relatively brief depopulation scenario, consistent with previous demographic models<sup>32</sup> and archaeological findings. We also find no evidence of ancestry from other populations (Palaeo-Eskimo, Inuit or Native American) in the Greenlandic Norse genomes (Supplementary Fig. 9.4), which accords with the skeletal remains<sup>32</sup>. This suggests that sexual interaction between the Greenland Norse populations and these other groups was absent, or occurred only on a very small scale.

### Genetic composition of earliest Viking voyage

Although maritime raiding has been a constant of seafaring cultures for millennia, the Viking Age is partly defined by this activity<sup>33</sup>. However, the exact nature and composition of Viking war parties is unknown<sup>5</sup>. One raiding or diplomatic expedition has left direct archaeological traces: at Salme in Estonia, 41 men from Sweden who died violently were buried in two boats, accompanied by high-status weaponry<sup>34,35</sup>. Importantly, the Salme boat burial predates the first textually documented raid (on Lindisfarne (England) in 793) by nearly half a century.

Kinship analysis of the genomes of 34 individuals from the Salme burial reveals 4 brothers buried side by side and a third-degree relative of 1 of the 4 brothers (Supplementary Note 4). The ancestry profiles of the Salme individuals were similar to one another when compared to the profiles of other burials of the Viking Age (Supplementary Notes 10, 11), which suggests a relatively genetically homogeneous group of people of high status (including close kin).

The five Salme relatives are not the only kin in our dataset; we also identified two pairs of kin in which the related individuals were excavated hundreds of kilometres apart from each other, which markedly illustrates the mobility of individuals during the Viking Age.

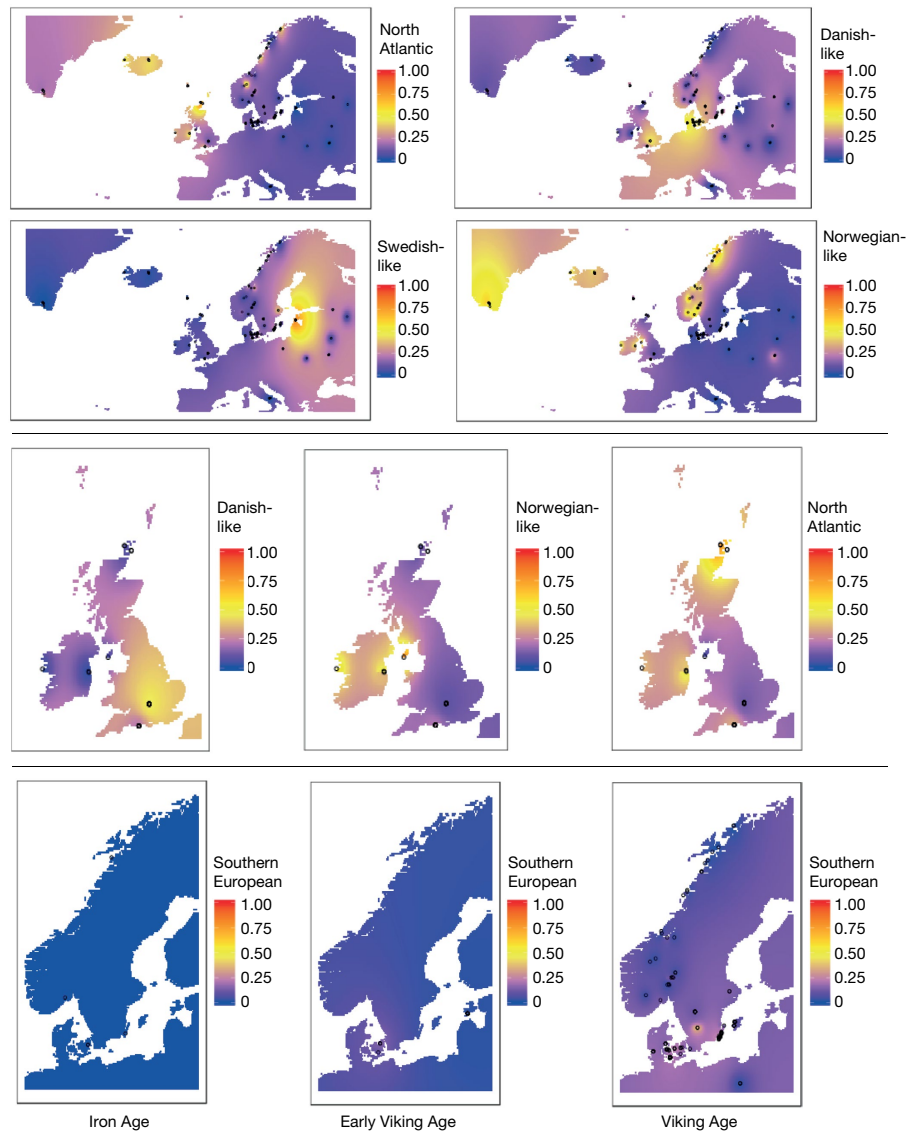
### Positive selection in northern Europe

We looked for single-nucleotide polymorphisms (SNPs) with allele frequencies that have changed significantly in the last 10,000 years<sup>36,37</sup>, beyond what can be explained by temporal changes in ancestry alone (Supplementary Note 14). Extended Data Figure 8a shows the likelihood ratio scores in favour of selection in the entire 10,000-year period (the general scan), the period up to 4,000 years before present (the ancient scan) and the period from 4,000 years before present up to today (the recent scan).

As expected<sup>38,39</sup>, the strongest candidates for selection are SNPs near *LCT*, the frequency of which increased after the Bronze Age<sup>40,41</sup>. Our dataset traces the frequency of the lactase-persistence allele (rs4988235) and its evolution since the Bronze Age. Extended Data Figure 8b shows that Viking Age groups had very similar allele frequencies at the *LCT* lactase-persistence SNP to those of present-day northern European populations. Conversely, Bronze Age Scandinavian individuals, as well as individuals from central Europe associated with Corded Ware and Bell Beaker assemblages, have a low frequency of this SNP despite evidence for milk consumption. Our Iron Age samples have intermediate frequencies, which suggests a rise in lactase persistence during this period. The frequency is higher in the Bronze Age of the Baltic Sea region than in Bronze Age Scandinavia, which is consistent with gene flow between the two regions explaining the increasing frequency of lactase persistence in Scandinavia.

Other candidates for selection include previously identified regions, including the one containing the *TLR1*, *TLR6* and *TLR10* genes, the HLA





**Fig. 4 | Spatiotemporal patterns of Viking and non-Viking ancestry in Europe during the Iron Age, Early Viking Age and Viking Age.** We performed inverse distance-weighting interpolation of the ancestry painting proportions of each individual genome on a dense grid of points covering the European continent, to better visualize the distribution of ancestry paintings at different periods (Supplementary Note 12). Top, distinct spheres of influence in the Viking world. Middle, Danish Viking ancestry in southern Britain, Norwegian

Viking ancestry in Ireland and Isle of Man and non-Scandinavian ('North Atlantic') ancestry in Orkney, Ireland and southern Britain. Bottom, Late southern European ancestry in southern Scandinavia. The Swedish-like ancestry is the highest in present-day Estonia owing to the ancient samples from the Salme ship burial, which originated from the Mälaren valley of Sweden (according to archaeological sources).  $n = 289$  genomes used for interpolation.

region, and the genes *SLC45A2* and *SLC22A4*<sup>41</sup>. We also find additional candidate regions for selection that have associated trajectories that start before the Viking Age, which suggests shared phenotypes between ancient Viking and present-day Scandinavian populations (Supplementary Note 14). These regions include one that overlaps *DCC* and that is implicated in colorectal cancer<sup>42</sup>, as well as one that overlaps *AKNA* and is involved in the secondary immune response<sup>43</sup>.

### Evolution of complex traits in Scandinavia

To search for signals of recent population differentiation at SNP markers associated with complex traits, we compared genotypes of Viking Age individuals with those of a panel of present-day Danish individuals<sup>44</sup>. We obtained summary statistics from 16 well-powered genome-wide association studies through the GWAS ATLAS<sup>45</sup> and tested for a difference in the distribution of polygenic scores between the two groups (Supplementary

Note 15). The polygenic scores of Viking Age individuals and present-day Danish individuals differed for three traits: black hair colour ( $P = 0.00089$ ), standing height ( $P = 0.019$ ) and schizophrenia ( $P = 0.0096$ ), although the latter two were not significant after accounting for the number of tests (Extended Data Fig. 7). Currently, we cannot conclude whether the observed differences in allele frequencies are due to selection acting on these alleles between the Viking Age and the present time or to some other factors (such as more ethnic diversity in the Viking Age sample). A binomial test of the number of black hair colour risk alleles at higher frequency in the Viking Age sample and the present-day sample was also significant (65/41;  $P = 0.025$ ), which suggests that the signal is not entirely driven by a few large-effect loci.

### Viking genetic legacy in populations today

To test whether present-day Scandinavian populations share increased ancestry with their respective counterparts in the Viking Age, we first

computed  $D$ -statistics of the form  $D(\text{Yoruba (YRI)}, \text{ancient}; \text{present-day population 1}, \text{present-day population 2})$ , which measure whether an ancient test individual shares more alleles with present-day population 1 or with present-day population 2. Viking Age individuals shift subtly from Scandinavia towards their present-day counterparts in the distributions of these statistics (Extended Data Fig. 5c, Supplementary Figs. 9.2, 9.3).

We further examined ancient ancestry in present-day populations using fineSTRUCTURE (Supplementary Note 11, Supplementary Fig. 11.14). Within Scandinavia, most present-day populations resemble their Viking Age counterparts. The exception is Swedish-like ancestry, which is present at only 15–30% within Sweden today: one cluster from Sweden is closer to ancient Finnish populations, and a second is more closely related to Danish and Norwegian populations. Danish-like ancestry is now high across the whole region.

Outside of Scandinavia, the genetic legacy of Viking Age populations is consistent—although limited. A small Scandinavian ancestry component is present in Poland (up to 5%). Within the British Isles, it is difficult to assess how much of the Danish-like ancestry is due to pre-existing Anglo-Saxon ancestry, but the Viking Age contribution does not exceed 6% in England (Supplementary Note 11). The genetic effects are stronger in the other direction. Although some North-Atlantic-like individuals in Orkney became culturally Scandinavian, others found themselves in Iceland, Norway and beyond, leaving a genetic legacy that persists today. Present-day Norwegian individuals vary between 12 and 25% in North-Atlantic-like ancestry; this ancestry is more uniformly 10% in Sweden.

## Discussion

Our genomic analyses shed light on long-standing questions raised by historical sources and archaeological evidence from the Viking Age. We largely confirm the long-argued movements of Vikings outside Scandinavia: Vikings from present-day Denmark, Norway, and Sweden going to Britain, the islands of the North Atlantic, and sailing east towards the Baltic region and beyond, respectively. However, we also see ancient Swedish-like and Finnish-like ancestry in the westernmost fringes of Europe, and Danish-like ancestry in the east, defying modern historical groupings. It is likely that many such individuals were from communities with mixed ancestries, thrown together by complex trading, raiding and settling networks that crossed cultures and the continent.

During the Viking Age, different parts of Scandinavia were not evenly connected, leading to clear genetic structure in the region. Scandinavia probably comprised a limited number of transport zones and maritime enclaves<sup>46</sup> with active external contacts, and limited external gene flow into the rest of the Scandinavian landmass. Some Viking Age Scandinavian locations are relatively homogeneous—particularly mid-Norway, Jutland and the Atlantic settlements. This contrasts with the strong genetic variation of populous coastal and southern trading communities such as in the islands of Gotland and Öland<sup>47–49</sup>. The high genetic heterogeneity in coastal communities implies increased population size, extending a previously proposed<sup>10</sup> urbanization model for the Late Viking Age city of Sigtuna (which suggested that more-cosmopolitan trading centres were already present at the end of the Viking Age in Northern Europe) both spatially and further back in time. The formation of large-scale trading and cultural networks that spread people, goods and warfare took time to affect the heartlands of Scandinavia, which retained pre-existing genetic differences into the Medieval period.

Finally, our findings show that Vikings were not simply a direct continuation of Scandinavian Iron Age groups. Instead, we observe gene flow from the south and east into Scandinavia, starting in the Iron Age and continuing throughout the duration of the Viking Age, from an increasing number of sources. Many Viking Age individuals—both within and outside Scandinavia—have high levels of non-Scandinavian ancestry, which suggests ongoing gene flow across Europe.

## Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41586-020-2688-8>.

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## Methods

No statistical methods were used to predetermine sample size. The experiments were not randomized and investigators were not blinded to allocation during experiments and outcome assessment.

### Laboratory work

Laboratory work was conducted in the dedicated ancient DNA clean-room facilities at the Globe Institute (University of Copenhagen), according to strict ancient DNA standards<sup>50,51</sup>. The overwhelming majority of ancient samples were petrous bones and teeth (Supplementary Table 1). The details of DNA extraction can be found in Supplementary Note 2. Double-stranded blunt-end DNA libraries were prepared using Illumina-specific adapters and NEBNext DNA Sample Pre Master Mix Set 2 (E6070) kit. We used an Agilent Bioanalyzer 2100 to quantify the amount of the purified DNA libraries. The libraries were sequenced 80-bp single-read chemistry on Illumina HiSeq 2500 machines at the Danish National High-throughput DNA Sequencing Centre.

### Bioinformatics analysis and quality assessment

We used AdapterRemoval v.2.1.3<sup>52</sup> for removing Illumina adaptor sequences, keeping only sequences with a minimum length of 30 bp. Adaptor-free sequences were mapped against the human reference genome build 37 using BWA v.0.7.10 aligner<sup>53</sup> with the seed (-l parameter) disabled for higher sensitivity of ancient DNA reads<sup>54</sup>. DNA sequences were processed with samtools v.1.3.1<sup>55</sup>, and only sequences with mapping quality  $\geq 30$  were kept. Picard v.1.127 (<http://broadinstitute.github.io/picard>) was used to sort the reads and remove duplicates. DNA libraries were combined at the sample level and realigned using GATK v.3.3.0<sup>55</sup> with Mills and 1000G gold-standard insertions and deletions (indels). At the end, realigned .bam files had the md-tag updated and extended base alignment qualities calculated using samtools calmd. Read depth and coverage were determined using pysam (<http://code.google.com/p/pysam/>) and BEDtools<sup>56</sup>. The mapping statistics for the ancient samples are summarized in Supplementary Table 2.

We used mapDamage v.2.0 to obtain approximate Bayesian estimates of damage parameters<sup>57</sup>. Data authenticity was assessed by estimating the rate of mismatches to the consensus mitochondrial sequence using contamMix<sup>58</sup> and Schmutzi<sup>59</sup>, as well as the excess of heterozygous positions in male haploid X chromosomes using ANGSD<sup>60</sup>. The sex of ancient individuals was determined by calculating the Ry parameter<sup>61</sup>.

### Uniparental haplogroup determination and kinship analysis

The mitochondrial haplogroups of the ancient individuals were assigned using haplogrep<sup>62</sup>. To get the mtDNA consensus sequences, we aligned the trimmed reads of ancient samples to the human mitochondrial reference genome: revised Cambridge Reference Genome (rCRS). Base quality  $\geq 20$  and mapping quality  $\geq 30$  filtering options were applied. Only SNPs at sites  $\geq 3\times$  coverage were considered for consensus calling using samtools mpileup/bcftools v.1.3.1<sup>55</sup>.

We identified male Y chromosome lineages using the pathPhynder workflow (<https://github.com/ruidlpm/pathPhynder>) and Yleaf v.2<sup>63</sup>. For the latter, the analysis was restricted to 26,083 biallelic SNPs from the International Society of Genetic Genealogy (ISOGG) 2019 database ([https://isogg.org/tree/ISOGG\\_YDNA\\_SNP\\_Index.html](https://isogg.org/tree/ISOGG_YDNA_SNP_Index.html)).

We used NgsRelate<sup>64</sup> to detect family relationships between all pairs of individuals. NgsRelate is a maximum-likelihood based program that—for a pair of individuals based on genotype likelihoods—estimates the three coefficients,  $k_0$ ,  $k_1$  and  $k_2$ , which denote the proportions of the genome in which the pair of analysed individuals share 0, 1 and 2 alleles identical-by-descent, respectively. We only included the 376 samples with sequencing depth above 0.1 $\times$  for the analysis. From these, we estimated genotype likelihoods and allele frequencies with ANGSD<sup>60</sup> using the SAMtools genotype likelihood model (-gl 1) including reads with mapping quality  $\geq 30$  and bases with base quality  $\geq 20$ . We estimated

genotype likelihoods and allele frequencies only for the autosomal transversion sites for which the 1000 Genomes CEU population (Utah residents with northern and western European ancestry) has a minor allele frequency of 0.05, resulting in 1,752,719 sites. READ<sup>65</sup> was used to confirm the degree of relatedness between pairs of individuals. The pedigree reconstructions on the basis of the kinship coefficients were conducted using Pedigree Reconstruction and Identification of a Maximum Unrelated Set (PRIMUS)<sup>66</sup>.

### Imputation

We imputed the genotypes of 298 ancient samples (289 from this study, and 9 from a previous study<sup>10</sup>) that had a sequencing depth greater than 0.5 $\times$ . We used Beagle v.4.1<sup>67</sup> for imputations based on the genotype likelihood data, which was first estimated by GATK v.3.7.0 UnifiedGenotyper. To generate the genotype data, we called only biallelic sites present in the 1000 Genomes dataset, and only the observed alleles (-genotyping\_mode GENOTYPE\_GIVEN\_ALLELES). The resulting .vcf files were filtered by setting genotype likelihoods to 0 for all three genotypes (for example, hom ref, het and hom alt) for sites with potential deamination (C>T and G>A), as described in a previous study<sup>68</sup>. Following this, the per-individual .vcf files were merged using bcftools v.1.3.1. The combined .vcf files were then split into 15,000 markers each and imputed separately using Beagle 4.0 using the 1000 Genomes phase-3 map included with Beagle (\*.phase3.v5a.snps.vcf.gz and plink.chr\*.GRCh37.map) with input through the genotype likelihood option. Run time for imputing using Beagle was approximately 280,000 core hours.

### Merge with existing panels

**Scandinavian panel.** To assess the genetic relationships of various Viking Age groups with their present-day counterparts, we constructed a reference panel enriched with Scandinavian populations on the basis of published datasets: the EGAD00010000632 data set from a previous publication<sup>23</sup> (UK dataset) and the EGAD00000000120 dataset from The International Multiple Sclerosis Genetics Consortium and The Wellcome Trust Case Control Consortium 2 (ref. <sup>69</sup>) (EU dataset) (see Supplementary Note 6 for details). The seven most relevant populations from Denmark, Sweden, Norway, Finland, Poland, UK and Italy were considered ( $n = 1,464$ ) with a total number of 414,264 SNPs. The Han Chinese (CHB) and Yoruba (YRI) populations from the 1000 Genomes project phase-3 database were merged to this panel as outgroups.

**The 1000 Genomes panel.** We used a set of 1,995 individuals from 20 populations (excluding individuals from the AMR super-population, as well as admixed ASW and ACB populations) of the 1000 Genomes project phase-3 release 5 (<ftp://1000genomes.ebi.ac.uk/vol1/ftp/release/20130502/>). We restricted the dataset to a set of 12,731,663 biallelic transversion SNPs located within the strict mappability mask regions ([ftp://1000genomes.ebi.ac.uk/vol1/ftp/release/20130502/supporting/accessible\\_genome\\_masks/](ftp://1000genomes.ebi.ac.uk/vol1/ftp/release/20130502/supporting/accessible_genome_masks/)).

Analyses of phenotype associated SNPs were carried out using five European-ancestry populations: Spanish (IBS), Tuscan (TSI), CEU, British (GBR) and Finnish (FIN), along with CHB and YRI as outliers. These were used to assess genome-wide allele frequencies for various SNPs associated with pigmentation phenotypes and lactose intolerance.

**Ancient panels.** We constructed datasets for population genetic analyses by merging the newly sequenced Viking Age individuals as well as other previously published ancient individuals<sup>40,41,68,70–96</sup> with the two modern reference panels. Ancient individuals were represented with pseudohaploid genotypes, by using mpileup command of samtools and randomly sampling an allele passing filters (mapping quality  $\geq 30$  and base quality  $\geq 30$ ), further requiring that it matched one of the two alleles observed in the reference panel (Supplementary Table 3).

## Clustering analyses

On the basis of the pseudohaploid individuals from the ancient panels, we ran ADMIXTURE<sup>97</sup> by thinning the dataset for linkage disequilibrium using plink with recommended settings (--indep-pairwise 50 10 0.1). This dataset contained 1,324 individuals for 151,235 markers for the autosomal chromosomes. Only samples with >20,000 SNPs overlapping with the Human Origins panel were kept in the analysis, resulting in 378 samples from this study. We did 50 replicates with different seeds for  $k = 2$  to  $k = 10$ . We used pong<sup>98</sup> to identify the best run for each  $k$  and similar components between different  $k$  values.

The large number of ancient individuals included in the analysis panels facilitates genetic clustering using the ancient individuals themselves, rather than projecting them on axes of variation inferred from modern populations. We carried this out using MDS on a distance matrix obtained from pairwise identity-by-state sharing between individuals, using the cmdscale function in R. We performed the main genetic clustering on a set of 1,306 ancient Eurasian individuals with >50,000 SNPs with genotype data, restricting to the batch-corrected SNP set described in Supplementary Note 8. Results from the batch-corrected MDS were combined with further dimensionality reduction using UMAP, implemented in the uwot package in R.

## Population genetics

We used  $f_4$  statistics to investigate allele-sharing between sets of test individuals and different modern and ancient groups (Supplementary Note 9). To characterize the deep ancestry relationship of the study individuals we calculated  $f_4$ (YRI, test individual; Barcin\_EN.SG, Yamnaya\_EBA.SG) for all ancient Europeans from the Bronze Age onwards (1000 Genomes panel merge). This statistic contrasts genetic affinities of the test individuals with two major ancestry groups that contributed to the gene pool of ancient Europeans from the Bronze Age onwards: Anatolian farmers and Steppe pastoralists. Genetic continuity with Scandinavian Iron Age groups was investigated using  $f_4$ (YRI, test group; test individual, Scandinavian Iron Age group) (1000 Genomes panel merge). This statistic measures whether a test individual is consistent with forming a clade with Scandinavian Iron Age groups to the exclusion of a test group from outside of Scandinavia. Genetic affinities between ancient groups and present-day populations were investigated using  $f_4$ (YRI, test individual; present-day test population, present-day reference population) (Scandinavian panel).

## Ancestry modelling using qpAdm

We estimated ancestry proportions of Viking Age groups using qpAdm<sup>70</sup>, which is based on  $f_4$ -statistics of the form  $f_4(X, O1; O2, O3)$ , in which  $X$  is either the source or target population, and  $O1$ ,  $O2$  and  $O3$  are triplets of outgroups to the source and target groups. To minimize batch effects and/or biases due to ancient DNA damage or SNP ascertainment, we used a set of 1,800,038 transversion-only sites that were found polymorphic with minor allele frequency  $\geq 0.5\%$  and missing genotype rate of  $\leq 15\%$  in the 1000 Genomes panel merge.

## Genetic diversity

The genetic diversity of ancient groups was assessed using conditional nucleotide diversity, as previously described<sup>73</sup>. For this analysis, pairwise differences between individuals were calculated using SNPs polymorphic in an outgroup population (YRI) and with a minor allele count  $\geq 5$  in the 1000 Genomes merge.

## Identity-by-descent analysis

The imputed genotypes of 298 individuals were used to infer genomic segments shared via identity-by-descent within the context of a reference panel of 1,464 present-day Europeans, using IBDseq<sup>99</sup> (version r1206) with default parameters. We conducted genetic clustering by MDS on a distance matrix obtained from pairwise identity-by-descent

sharing and UMAP to reveal fine-scale population structure among Viking Age individuals.

## Painting

To assess the fine-scale variation in genetic ancestry proportions of Viking Age individuals we used Chromosome Painting<sup>11</sup>. The following describes the general workflow of the Chromosome Painting analysis (see Supplementary Note 11 for details).

First, we created a modern reference panel using 1,675 modern individuals sampled from northern Europe, using the standard FineSTRUCTURE pipeline. We applied ChromoPainter to paint all modern individuals using the remaining individuals as donors using fs2.0.8. Related individuals were identified through increased haplotype similarity, and admixed individuals were identified by their FineSTRUCTURE clustering. These were removed, leading to 1,554 unrelated individuals who were re-painted. These individuals were then clustered using FineSTRUCTURE, resulting in 40 populations. After removal of small populations and merging of the CHB and YRI subpopulations, this resulted in 23 modern populations with geographical meaning. We named the resulting clustering the modern reference panel, which consists of 23 modern surrogate populations and 23 modern donor populations (Supplementary Fig. 11.2).

Second, we created an ancient reference panel using the modern reference panel, by applying ChromoPainter to paint all ancient individuals using the modern population palette (Supplementary Fig. 11.3). We then created a supervised ancient population palette consisting of 14 populations which either (a) represent a modern ancestry direction or (b) are best associated with a modern ancestry direction. The paintings consider the average per-individual donor rate to each of the seven modern populations, normalizing each donor label to have a mean of 1 (Supplementary Fig. 11.4). The individuals that contribute most to a population represent it (above a threshold amount chosen by identifying a change point). The remaining individuals are assigned to the population that they are best associated with. We create an ancient population surrogate for each modern population, consisting of the individuals that represent each modern population. For  $k = 7$  modern populations, this results in a matrix of  $k = 7$  rows (surrogate populations) and  $2K = 14$  columns (donor palette populations), which captures the ancient population structure (Supplementary Fig. 11.6).

Third, we inferred ancestry by learning about population structure in modern individuals or ancient individuals, painting them with respect to the ancient population panel and fitting them as a mixture using the ancient population surrogates, using the non-negative least squares implemented in GLOBETROTTER<sup>100</sup> (Supplementary Information section 11) with uncertainty estimated using 100 bootstrap replicates. All samples were analysed by leaving out one individual per donor population so that modern and ancient individuals are exchangeable (as the ancient individual is itself excluded from its own ancient donor population). We report this in a number of ways. The inferred ancestry results (Supplementary Table 6) are summarized by taking the mean across inferred populations in Supplementary Fig. 11.11; Supplementary Fig. 11.12 shows the means over sample information labels. We performed a spatiotemporal regression (Supplementary Table 11.2) using the model  $a_{ik} = \alpha_{jk}t_i + \beta_{jk}x_i + \gamma_{jk}y_i + \varepsilon_{ijk}$  in which  $a_{ik}$  is the amount of ancestry individual  $i$  possesses from population  $k$  in regional analysis  $j$ ,  $t_i$  is the age category of the individual (1 = Iron Age, 2 = Early Viking Age, 3 = Viking Age, 4 = Medieval) and  $x_i$  and  $y_i$  are the longitude and latitude of the burial location of the individual, respectively. The modern ancestry results are estimated using the spatial median instead of the mean, to account for ancestry being constrained in a  $k$ -dimensional simplex (Supplementary Fig. 11.14), with uncertainty quantified by bootstrap resampling of individuals (Supplementary Fig. 11.15).

Fourthly, we performed sensitivity analyses to ensure that the inference procedure performed as expected. We checked that sequence depth was not associated with cluster membership (Supplementary

Fig. 11.7), and that sequence depth did not significantly affect inferred ancestry (Supplementary Fig. 11.8) by downsampling individuals with high-depth data available, rephasing, re-imputing and repainting them, and assigning ancestry using the above procedure. Results 2× and above were extremely similar, whereas at 1× there was some loss of precision but the broad structure remained clear.

Finally, we ran a principal components analysis of the ancient + modern populations painted against our donor populations (Supplementary Fig. 11.9) as well as an all-versus-all ChromoPainter analysis including modern and ancient individuals (Supplementary Fig. 11.10).

### Ancestry diversity measure

We wish to quantify diversity in ancestry for a population of individuals, with diverse meaning a large deviation of individual ancestry estimates from the average ancestry in that population. We compute the average Kullback–Leibler divergence for each individual label from the average of that label:

$$D(A^{(l)}) = \frac{1}{n_l} \sum_{i=1}^{n_l} KL(A_i^{(l)} \parallel p^{(l)})$$

in which  $A^{(l)}$  is the  $n_l$  by  $K$  matrix of ancestry estimates in label  $l$ ,  $p^{(l)}$  is the length  $K$  vector of average ancestries in that label, and  $KL(Q \parallel P) = \sum_{k=1}^K q_k \log_2 \left( \frac{q_k}{p_k} \right)$ . We performed a simulation study to validate this measure (Supplementary Information section 11, Supplementary Fig. 11.13), which allowed us to calibrate the expected diversity as a function of sample size.

### Spatiotemporal patterns

To visualize the migration patterns of the Vikings, we used inverse distance weighting interpolation—implemented in the function `idw` of the R package `gstat`—to interpolate the proportion of each ancient genome that was attributed by our fineSTRUCTURE analysis (Supplementary Table 6) to one of the predefined ancestry groups: UK, Denmark, Norway, Sweden, Italy, Poland and Finland. We used the Shepard method of interpolation<sup>12,101</sup> with the weight for a given interpolation location  $x$  equal to  $1/(d(x,v)^2)$ , in which  $v$  is the location of an observed sample and  $d(a,b)$  is the distance between two points  $a$  and  $b$ . For plotting maps, we used a Mercator projection and downloaded coastal contours at 1:50-m scale from Natural Earth (<https://www.naturalearthdata.com/>).

### Lactase persistence and pigmentation SNPs

For ancient populations we estimated the derived A allele frequency of the SNP rs4988235, known to affect expression of the lactase (*LCT*) gene. The ancestral G allele is responsible for lactase intolerance in adult Europeans<sup>39</sup>. We used ANGSD<sup>60</sup> to estimate the allele frequencies of the ancient population on the basis of the genotype likelihood data. We used the five European populations (CEU, FIN, GBR, TSI and IBS) and two outgroups (YRI and CHB) from the 1000 Genomes Project as comparative groups. We also included the present-day Danish population from the IPSYCH case-cohort study<sup>44</sup> and geographically proximate Iron and Bronze Age populations to trace frequency shifts of SNP rs4988235 through time. We also used ANGSD<sup>60</sup> to estimate the frequencies of 22 SNPs (HLIRISplex<sup>102</sup>) with strongest influence on human pigmentation phenotypes in the Viking Age and Early Viking Age Scandinavian population.

### Signatures of selection

We aimed to find SNPs with allele frequencies that changed significantly in the last 10,000 years, using our ancient human genomes to look at the frequencies of alleles in the past. We combined our Viking Age and Iron Age genomes with previously published present-day, Bronze Age, Neolithic and Mesolithic sequence data typed at the Human Origins array (Supplementary Note 6). We filtered for genomes that were younger than 8000 BC and that were located within a bounding box

encompassing the European continent:  $30^\circ < \text{latitude} < 75^\circ$  and  $-15^\circ < \text{longitude} < 45^\circ$ . We then used neoscan in Ohana<sup>36,103</sup> to scan for variants with allele frequencies that were strongly associated with time, after controlling for genome-wide changes in ancestry that might have also occurred over time. We analysed only sites with a minor allele frequency  $>1\%$  (Supplementary Note 14).

### Tracking the evolution of complex traits in Scandinavia

We wanted to examine whether we could identify signals of recent population differentiation of complex traits by comparing genotypes of Viking Age samples excavated in Scandinavia (that is, Denmark, Sweden and Norway) with those of a present-day Scandinavian population. For the latter, we used imputed genotypes from subjects born in Denmark between 1981 and 2011 from the IPSYCH case-cohort study<sup>44</sup>. We downloaded summary statistics from the genome-wide association study ATLAS webpage (<https://atlas.ctglab.nl/>)<sup>45</sup>, from studies of 16 disease- and anthropometric traits (excluding those related to cognition) published in 2017 or later with SNP heritability estimated at  $>0.1$ , sample size of  $>100,000$  and  $>100$  identified genome-wide significant loci. We calculated polygenic risk scores based on independent ( $R^2 < 0.1$  within 10-Mb range) genome-wide significant allelic effects and standardized them to a unit representing the standard deviation of the mean of their distribution. We then removed outliers (anyone with a value for any of the 25 principal components falling more than 4 standard deviations away from the group mean) reiteratively from within each ancestry group (treating the Scandinavian Viking age samples as one ancestry group), and subsequently tested for difference in polygenic risk score distribution between Viking Age samples and Danish-ancestry IPSYCH random population samples using a linear regression model correcting for sex and the 25 principal components.

### Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

### Data availability

Sequence data are available at the European Nucleotide Archive under accession number PRJEB37976.

### Code availability

Functions for calculating  $f$ -statistics are available as an R package at GitHub (<https://github.com/martinsikora/admixr>).

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**Author contributions** E.W. initiated and led the study. E.W., A.M., D.J.L., Martin Sikora, F.R., R.N., K.K., L.H., S.M.S., J.B., N.P., T.W., A.I., M.E.A., M.W.P., N.L., J.A., I. Moltke and A.A. designed the study. A.M., P.d.B.D., L.M.C., M.M.B., A.K.F., I.L. and J.S. produced the data. A.M., D.J.L., Martin Sikora, F.R., S.R., I. Moltke, R.N., T.W., L.M.C., E.J., A.I., M.W.P., T.K., R.M., G.R., C.B., J.V.M.-M., H.M., A.A., J.C., K.H.I. and M.E.A. analysed or assisted in analysis of data. E.W., A.M., D.J.L., Martin Sikora, F.R., S.M.S., K.K., L.H., R.N., M.C. and A.I. interpreted the results with considerable input from I. Moltke, M.E.A., M.W.P., T.K., H.W., R.M., G.R., T.W., C.H.J., J.A., N.L., N.P., J.B., A.A., M.T.P.G., L.O. and other authors. E.W., A.M., D.J.L., Martin Sikora, F.R., S.M.S., K.K. and L.H. wrote the manuscript with considerable input from M.C., J.B., N.P., I. Moltke, N.L., A.I., R.M., E.J., J.A., M.L.J., C.H.J., M.W.P., M.E.A., G.R. and M.M., with contributions from all authors. A.M., L.M.C., M.W.P., H.W., M.M.B., P.d.B.D., A.K.F., M.A., R.A., M.M., E.C., G.S., A.B., A.F., B. Schütz, B. Skar, C.A., C.F., D.B., D.P., G.T.-W., H.G., I.L., I.G., I. Mainland, I.P., I.M.M., J.M., J. Gibson, J.P., J. Gustafsson, L. Simpson, L. Strand, L.L., Maeve Sikora, M.F., M.V., M.R., M.B., T.P., M. Søvsø, N.G., T.C., O.K., O.U., P.F., P.H., S.S., S.V.A., S.E., V.M., W.B., Y.M., P.P., M.D.J., A.P., D.G.B., M.L.J., J.A., N.L., N.P., M.T.P.G., M.E.A., J.B. and E.W. excavated, curated, sampled and/or described analysed skeletons.

**Competing interests** The authors declare no competing interests.

## Additional information

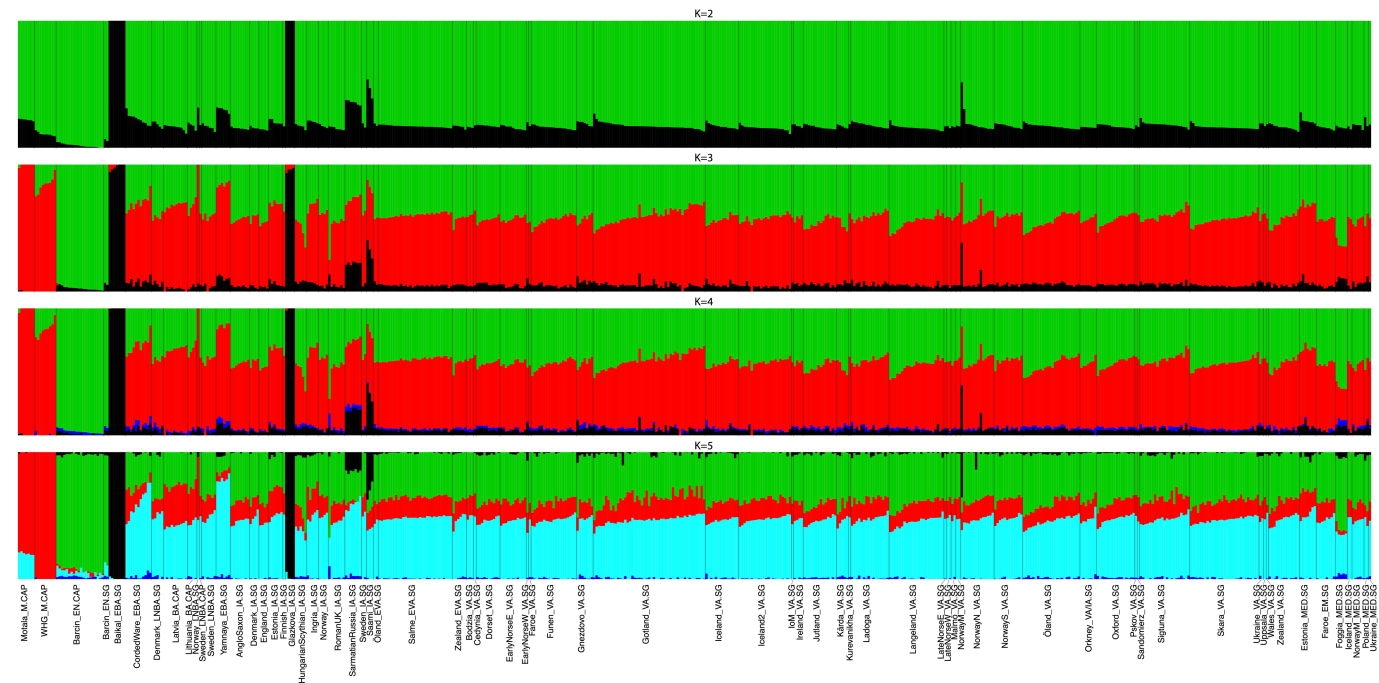
**Supplementary information** is available for this paper at <https://doi.org/10.1038/s41586-020-2688-8>.

**Correspondence and requests for materials** should be addressed to R.N., T.W. or E.W.

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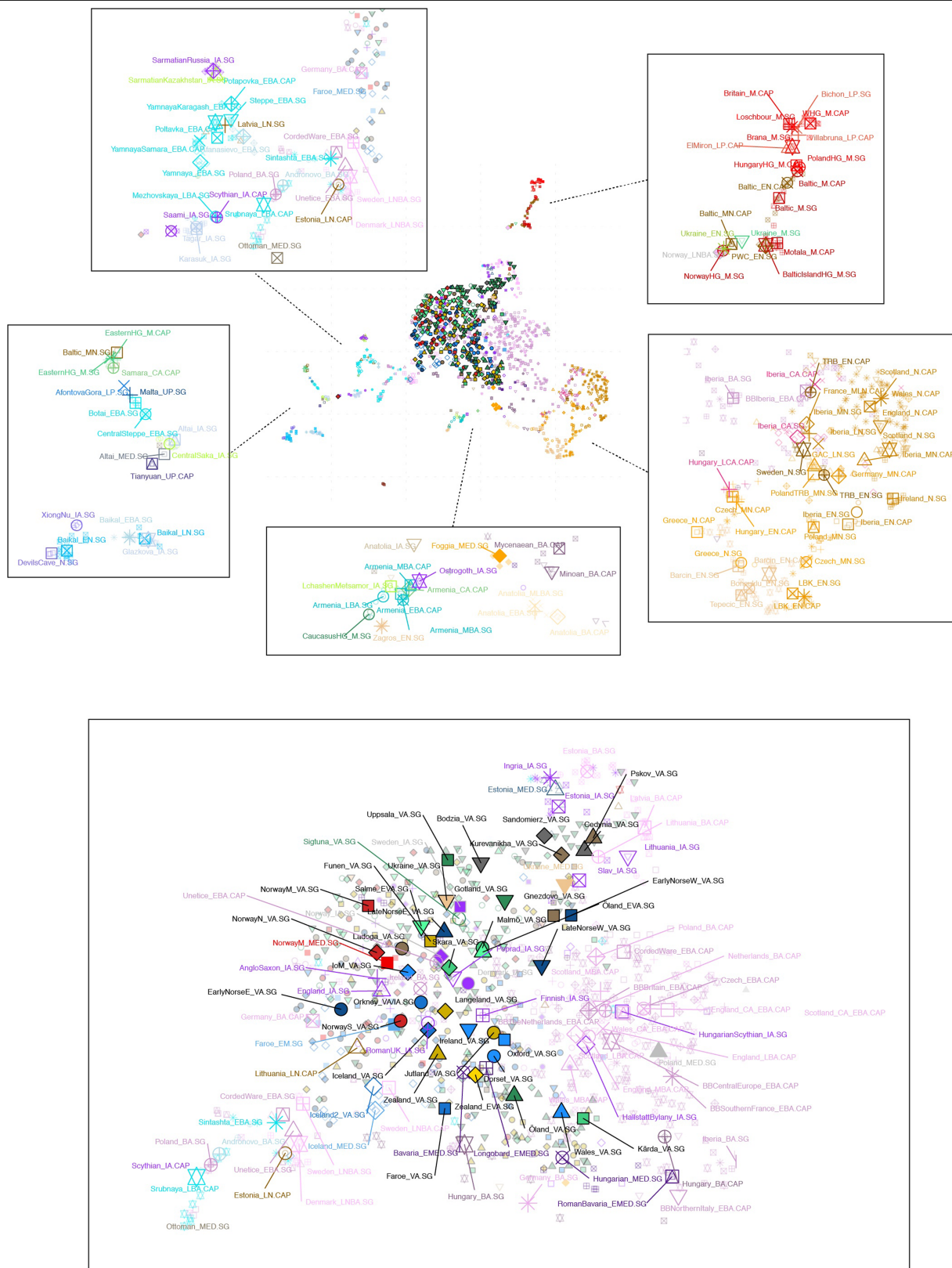
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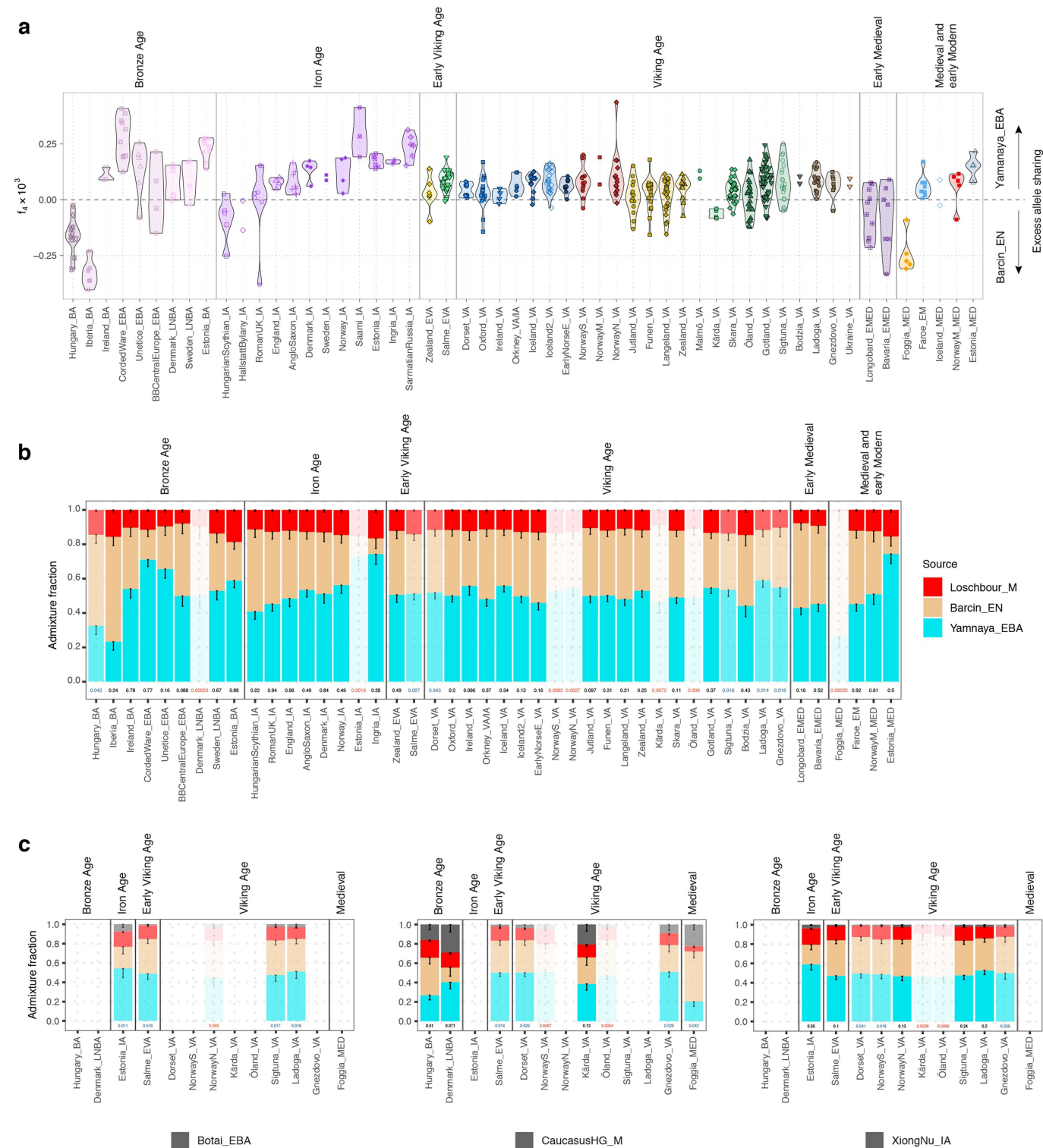
**Extended Data Fig. 2 | Model-based clustering analysis.** Admixture plot ( $K = 2$  to  $K = 5$ ) for 378 ancient individuals, spanning 71 populations. This figure is a subset of the most relevant individuals and populations from Supplementary Fig. 7.2; see Supplementary Note 7 for further details. This plot consists of 378 ancient samples from this study; Viking Age samples from Sigtuna (Sweden)<sup>10</sup> ( $n = 21$ ), Iceland<sup>18</sup> ( $n = 22$ ) and other ancient comparative groups ( $n = 146$ ).





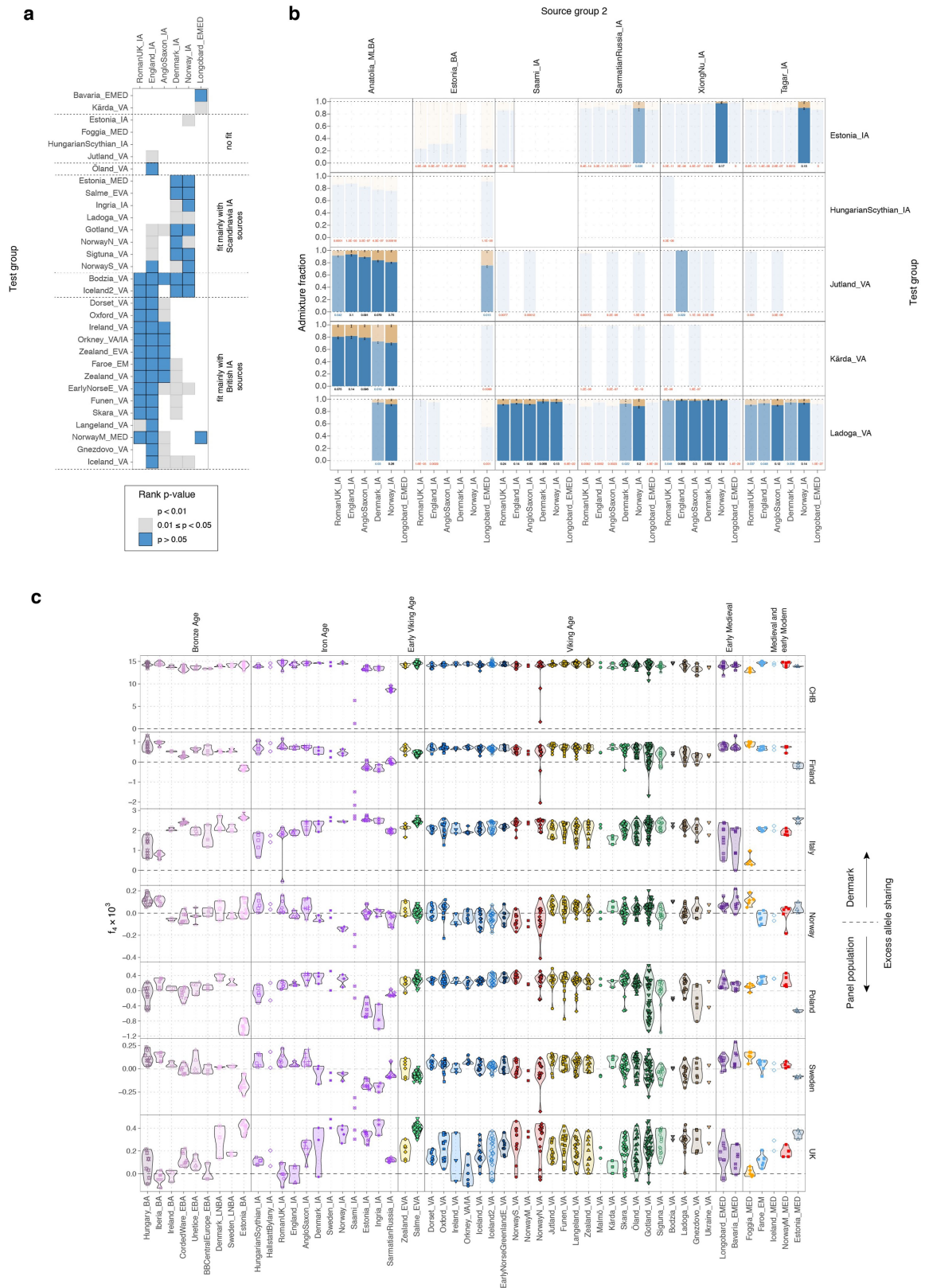
**Extended Data Fig. 3 | Fine-scale population structure.** The point cloud at the top centre shows an alternative view of the UMAP result from Fig. 2b, with all ancient individuals coloured on the basis of analysis group. The framed panels surrounding the point cloud highlight particular ancestry clusters

(as indicated), with labels and larger symbols corresponding to the median coordinates for the respective group. Similarly, the larger bottom panel shows median group coordinates for the large central point cloud, which includes the vast majority of European individuals from the Bronze Age onwards.



**Extended Data Fig. 4 | Ancestry modelling for distal sources. a**, Contrasting allele-sharing between Anatolian farmers (Barcin\_EN) and Steppe pastoralists (Yamnaya\_EBA) for European individuals from the Bronze Age and later. Violin plots showing distributions of statistics  $f_4$  (YRI, test individual; Barcin\_EN, Yamnaya\_EBA) for  $n = 515$  individuals with a minimum of 1,000,000 SNPs with genotypes and groups with at least 2 such individuals. **b**, Ancestry proportions of analysis groups from the Bronze Age and later inferred using qpAdm. Target groups were modelled using three distal sources representing

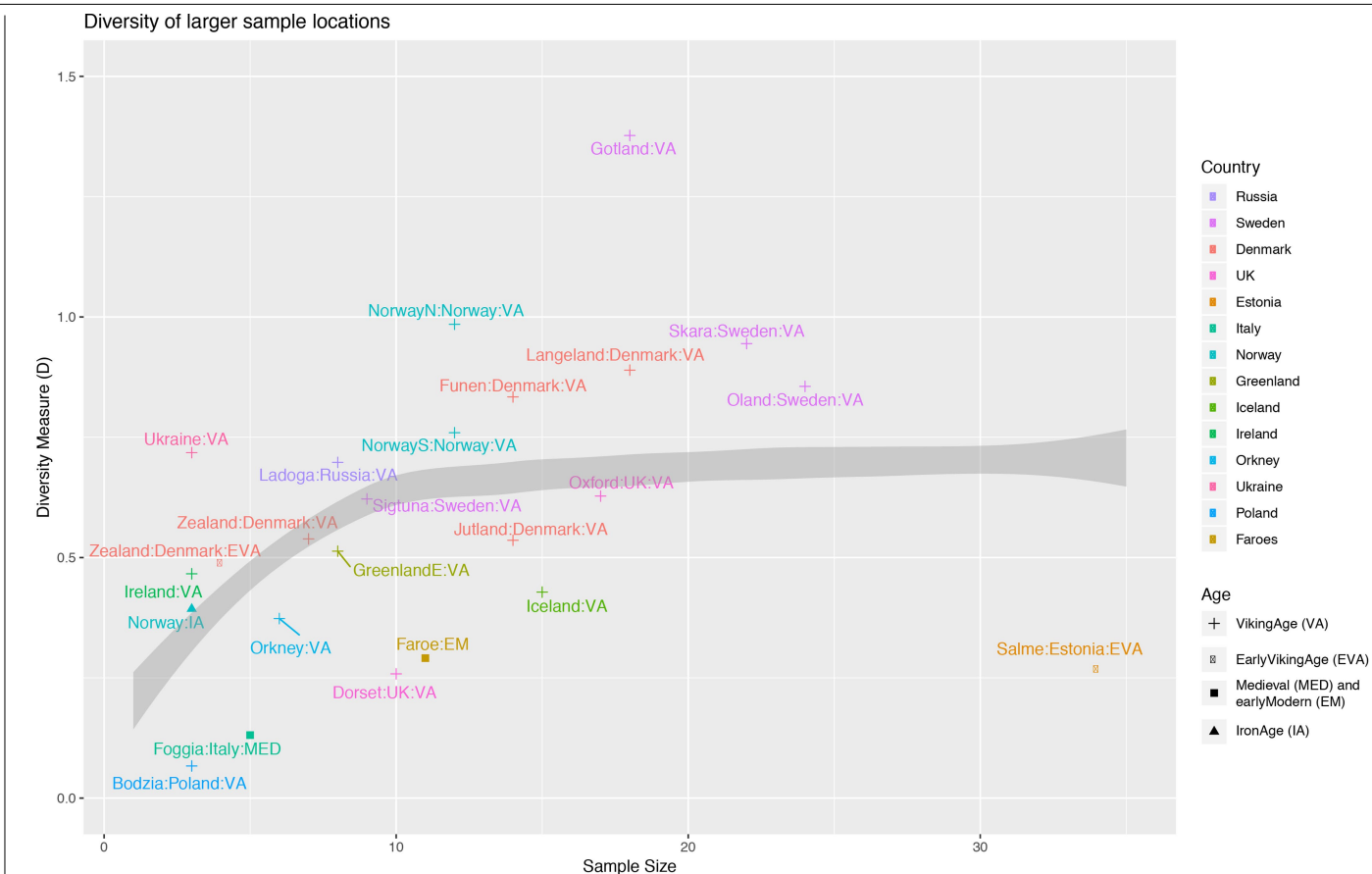
European hunter-gatherer (Loschbour\_M), Anatolian farmer (Barcin\_EN) and Steppe pastoralist (Yamnaya\_EBA) ancestry. Sample sizes for target groups can be found in Supplementary Table 10. Error bars indicate standard error obtained from qpAdm. **c**, Ancestry proportions of analysis groups for which the three-source model was rejected using qpAdm ( $P < 0.05$ ). Target groups were modelled including one additional distal source representing either Steppe hunter-gatherer (Botai\_EBA), Caucasus hunter-gatherer (CaucasusHG\_M) or East-Asian-related (XiongNu\_IA) ancestry.



Extended Data Fig. 5 | See next page for caption.



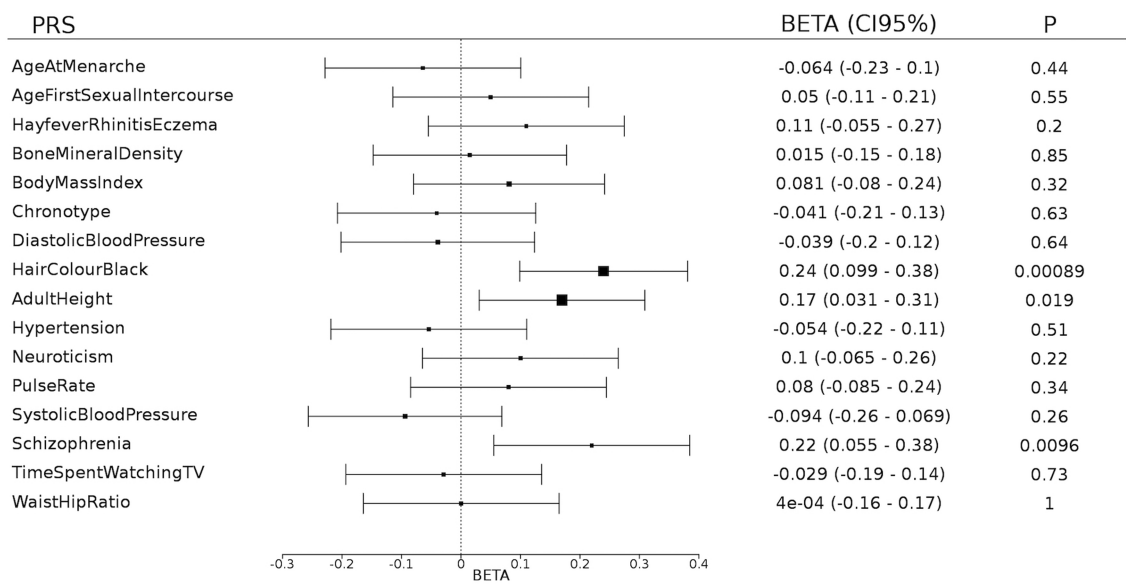
**Extended Data Fig. 5 | Ancestry modelling for proximate sources. a,** Testing for continuity between European Iron Age and later Viking Age and Medieval groups. Coloured squares depict whether a particular target group (row) can be modelled using a single source group (column).  $P$  values for  $f_4$  rank of 0 (corresponding to a single source group) were obtained using qpAdm with a set of 15 outgroups, which included European Bronze Age groups that preceded the source groups. Sample sizes for target groups can be found in Supplementary Table 12. **b,** Two-way admixture ancestry proportions of target groups for which a single source was rejected ( $P \leq 0.05$ ). Target groups were modelled using additional proximate Bronze and Iron Age sources. Sample sizes for target groups can be found in Supplementary Table 13. For both **a** and **b**, only ancient groups containing at least 3 individuals with a minimum of 1,000,000 SNPs with genotypes are plotted. **c,** Contrasting allele-sharing between populations of present-day Denmark and other populations. Violin plots showing distributions of statistics  $f_4$  (YRI, test individual; panel population, Denmark) for  $n = 489$  individuals with a minimum of 50,000 SNPs with genotypes and groups with at least 2 such individuals. Median values for distributions are indicated with horizontal lines.



**Extended Data Fig. 6 | Ancestry diversity of different population groups.** Diversity of different labels (that is, sample locations combined with historical age) are shown as a function of their sample size. The diversity measure is the Kullback–Leibler divergence from the label means, capturing the diversity of a

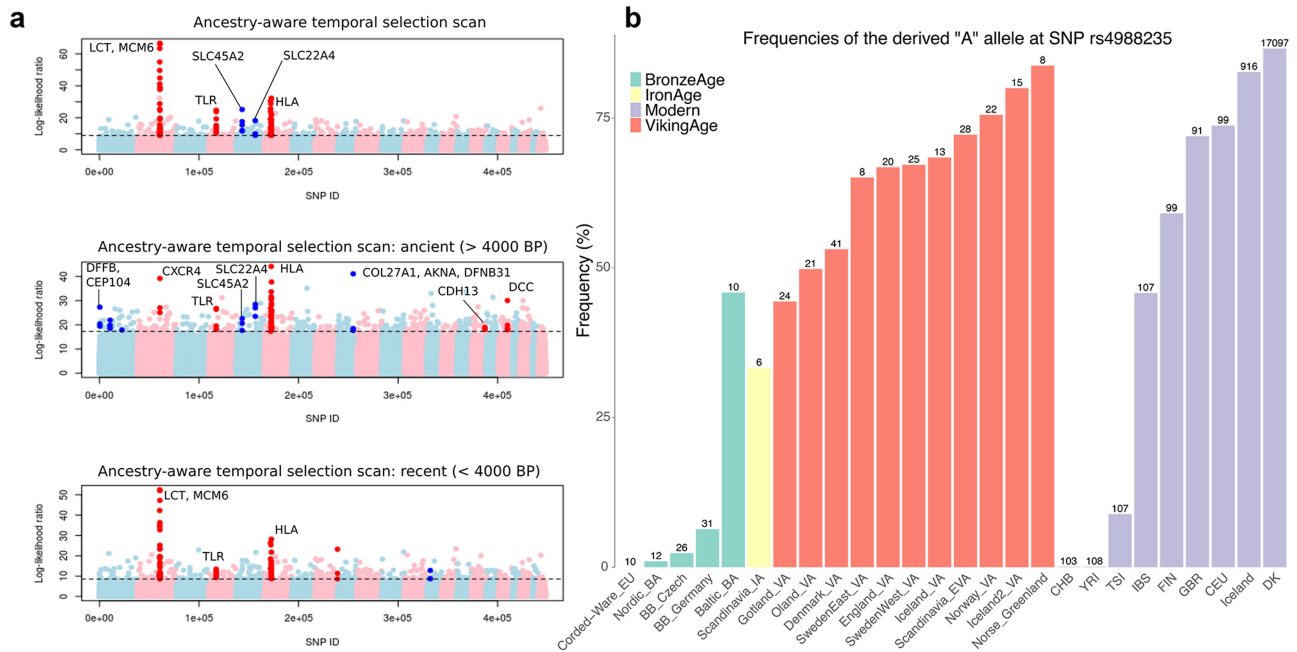
group with respect to the average of that group (see Supplementary Note 11 for details). Larger values are more diverse, although a dependence on sample size is expected. The simulation expectation for the best fit to the data ( $D = 0.2$ ) is shown.

Viking age sample compared against a present-day Danish random sample



**Extended Data Fig. 7 | Polygenic risk scores.** Polygenic risk scores (PRS) for 16 complex human traits in 148 Viking Age samples from Denmark, Sweden and Norway, compared against a reference sample of 20,551 Danish-ancestry individuals randomly drawn from all individuals born in Denmark in 1981–2005. The PRS is in each case based on allelic effects for >100 independent genome-wide significant SNPs from recent genome-wide association studies of the respective traits and standardised to a mean of 0 and standard deviation of 1 in the entire sample. Difference in PRS was estimated in a linear regression

correcting for sex and 25 principal components of overall genetic structure. The plotted BETA indicates the coefficient for the test-group (Viking Age sample) PRS compared to that of the Danish comparison sample, with error bars indicating the 95% confidence interval of BETA, and *P* indicating the two-tailed *P* value of the corresponding *t*-test (not corrected for number of tests). Only PRS for black hair colour is significantly different between the groups after taking account of multiple testing.



**Extended Data Fig. 8 | Positive selection in Europe. a**, Manhattan plots of the likelihood ratio scores in favour of selection looking at the entire 10,000-year period (top, general scan), the period up to 4,000 years before present (middle, ancient scan) and the period from 4,000 years before present up to the present day (bottom, recent scan). The highlighted SNPs have a score larger than the 99.9% quantile of the empirical distribution of log-likelihood ratios, and have at least two neighbouring SNPs ( $\pm 500$  kb) with a score larger than the same

quantile.  $n = 1,185$  genomes are used in the selection scan. **b**, Frequencies of the derived A allele rs4988235 SNP responsible for lactase persistence in humans for different Viking Age groups, present-day populations from the 1000 Genomes Project as well as relevant Bronze Age population panels. The numbers at the top of the bars denote the sample size on which the allele frequency estimates are based.



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|-----------------|---|
| Data collection | No specific software was used for data collection. All software used in this study is listed below.   |
| Data analysis   | <p>All software used in this work is publicly available. Corresponding publications are cited in the main text and supplementary material. List of software and respective versions:</p> <p>CASAVA v1.8.2<br/>           AdapterRemoval v2.1.3<br/>           bwa v0.7.10<br/>           bwa mem 0.7.10<br/>           picard tools v1.127<br/>           bamUtil v1.0.14<br/>           samtools v1.3.1<br/>           GATK v3.3.0<br/>           pysam 0.7.4 (python module)<br/>           bedtools 2.27.1<br/>           mapDamage2.0<br/>           contamMix v1.0-5<br/>           SHRIMP 2.2.3<br/>           YFitter v0.2<br/>           Haplogrep 2.0<br/>           FineSTRUCTURE<br/>           ANGSD v0.915<br/>           IBDseq v.r1206<br/>           PRANK v.150803<br/>           pathPhynder<br/>           BEASTv1.8.2</p> |

```

schmutzi v.1.5.4
Yleaf v2
admixtools v4.1
NGSrelate v.1
GLOBETROTTER
READ
PRIMUS v1.9
plink and v1.9
ADMIXTURE v1.3
RAxML-8.1.15
SnEff
SPAdes-3.9.0
R 3.2.3
Ohana
Beagle v4.1
python 2.7.12
perl v5.22.1
CALIB
FigTree v.1.4.4

```

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research [guidelines for submitting code & software](#) for further information.

## Data

Policy information about [availability of data](#)

All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

Sequence data are available at the European Nucleotide Archive under accession number PRJEB37976.

## Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

☒ Life sciences ☐ Behavioural & social sciences ☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://www.nature.com/documents/nr-reporting-summary-flat.pdf)

## Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

|                 |   |
|-----------------|---|
| Sample size     | We did not rely on statistical methods to predetermine sample sizes. Sample sizes in ancient population genetic studies are limited by the number of samples yielding endogenous DNA proportions amenable to whole genome sequencing.   |
| Data exclusions | We selected 442 samples for whole-genome sequencing, out of all (n=528) screened samples, based on their endogenous content and low contamination estimates. These criteria are described in detail in Supplementary Notes 2 and 3. Furthermore, closely related individuals were excluded from analyses requiring population allele frequencies. |
| Replication     | We did not attempt to specifically replicate experimental findings. But we note that samples from the same population carry similar genetic signatures. Moreover, genome-wide data allows for the analysis of multiple realisations of the sample history, by studying hundreds of thousands of SNP sites.  |
| Randomization   | We did not implement any randomization as no experimental groups or effect sizes were measured in this study.   |
| Blinding        | No blinding techniques were implemented, as experimental group assignment is not relevant for population genetic studies of this kind.  |

## Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

- |                                     |  |
|-------------------------------------|--|
| n/a                                 | Involvement in the study                             |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> Antibodies                  |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> Eukaryotic cell lines       |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> Palaeontology               |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> Animals and other organisms |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> Human research participants |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> Clinical data               |

Methods

- |                                     |   |
|-------------------------------------|---|
| n/a                                 | Involvement in the study                        |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> ChIP-seq               |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> Flow cytometry         |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> MRI-based neuroimaging |