#### **ANTHROPOLOGY**

# **Genetic history of Cambridgeshire before and after the Black Death**

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**The extent of the devastation of the Black Death pandemic (1346–1353) on European populations is known from documentary sources and its bacterial source illuminated by studies of ancient pathogen DNA. What has remained less understood is the effect of the pandemic on human mobility and genetic diversity at the local scale. Here, we report 275 ancient genomes, including 109 with coverage >0.1×, from later medieval and postmedieval Cambridgeshire of individuals buried before and after the Black Death. Consistent with the function of the institutions, we found a lack of close relatives among the friars and the inmates of the hospital in contrast to their abundance in general urban and rural parish communities. While we detect long-term shifts in local genetic ancestry in Cambridgeshire, we find no evidence of major changes in genetic ancestry nor higher differentiation of immune loci between cohorts living before and after the Black Death.**

#### **INTRODUCTION**

<span id="page-0-4"></span><span id="page-0-3"></span><span id="page-0-2"></span><span id="page-0-0"></span>Evidence from ancient DNA (aDNA) continues to increase our understanding of the human past. By linking the genetic profiles to a place and time, it allows us to study population movements (*[1](#page-11-0)*, *[2](#page-11-1)*), genetic relatedness (*[3](#page-11-2)*–*[5](#page-11-3)*), infectious diseases (*[6](#page-11-4)*, *[7](#page-11-5)*), and natural selection (*[8](#page-11-6)*, *[9](#page-11-7)*) as they occurred. When combined with historical and archaeological contexts, such information offers a more detailed perspective on life in past societies.

aDNA studies centered around broad geographical regions and long time periods have been fundamental in establishing major migration events, population turnovers, or continuity in both prehistory and historic periods, while being less informative about everyday life experience within complex societies. Taking what we call "the whole town approach," we have studied hundreds of skeletal remains from later medieval (*c*. 1000–1550 CE) Cambridgeshire ([Table 1\)](#page-1-0). They were excavated from burial grounds connected with different social groups: urban and rural parish churchyards, urban charitable institutions, and religious institutions. For historical context, we also included postmedieval burial grounds (*c*. 1550–1855 CE). Apart from Clopton and Hemingford Grey, all the sites are within a few kilometers of each other [\(Fig. 1A\)](#page-2-0).

Cambridge during the later medieval period was a middle-sized market town where people from all sections of society crossed paths. After they died, most townspeople were buried in one of the parish

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<span id="page-0-7"></span><span id="page-0-6"></span><span id="page-0-5"></span><span id="page-0-1"></span>cemeteries, including All Saints by the Castle; however, other places of burial existed and increased over time. Toward the end of the 12th century, the Hospital of St John the Evangelist was founded by the townspeople as a charitable institute for the poor, the infirm and the sick. Most of the burials included in this study come from a cemetery for charitable inmates of the Hospital. The 13th century saw the founding of the university and houses of the mendicant orders, including an Augustinian Friary (from which some of our study population are drawn). Apart from the friars, some patrons of the friary were also buried in the cemetery and chapter house of the friary. The first wave of the Second Plague Pandemic (which we hereafter refer to with the commonly used term "Black Death," although the term was not in use until the 18th century) hit Cambridge in 1349; some of its victims were found in a mass burial of unknown size on Bene't Street (*[10](#page-11-8)*). Two parish cemeteries outside Cambridge, Cherry Hinton and Clopton, are within the rural hinterland of the town. [Table 1](#page-1-0) and table S1 list the archaeological sites covered in this study, the dating of the burials, and the function of the medieval and postmedieval sites.

<span id="page-0-10"></span><span id="page-0-9"></span><span id="page-0-8"></span>The Black Death and subsequent plague outbreaks had multiple effects on medieval society in England. Its death toll in Europe, estimated at 30 to 65% (*[11](#page-11-9)*, *[12](#page-11-10)*), could have posed selective pressure for better resistance to the plague. Genetic adaptations via the innate (*[13](#page-11-11)*) and adaptive immune system (*[14](#page-11-12)*) have been proposed. Although reference bias can pose challenges to detect allele frequency changes due to natural selection (*[15](#page-11-13)*), a study of 206 aDNA extracts from individuals buried in London and Denmark before, during, and after the Black Death revealed enrichment of immune genes among highly differentiated single-nucleotide variants, suggesting major impact of the pandemic in shaping the disease susceptibility of surviving population (*[16](#page-11-14)*).

<span id="page-0-14"></span><span id="page-0-13"></span><span id="page-0-12"></span><span id="page-0-11"></span>Besides genetic susceptibility, it is not clear to what extent social identity modified the morbid and mortal effects of the Black Death. Plague mortality appeared to be selective with respect to frailty (*[17](#page-11-15)*, *[18](#page-11-16)*) caused possibly by factors such as malnutrition, impaired immunocompetence, and others affected by social conditions. For example, the Great Famine of 1315–1322 could have severely affected people of low socioeconomic position. In this sense, health inequality between social groups sets the background for understanding the potential for

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different experiences through the pandemic. As longer-term consequences of the mortality, it has been argued that the Black Death pandemic initiated or accelerated profound socioeconomic changes, such as increased social mobility, improved quality of life of the laboring population, and technological innovations to increase productivity (*[19](#page-11-17)*, *[20](#page-11-18)*). Together with evidence from osteology, isotopes, and the rich context around the burial grounds, we aim to explore to what extent genetic data might aid the construction of a social history, both in relation to the pandemic and regarding the more stable aspects of later medieval life.

#### **RESULTS**

We extracted aDNA and generated whole-genome shotgun sequence data with a mean coverage of 0.228 from a total of 250 later medieval and 25 postmedieval skeletons, retrieving for further analyses 190 ge-nomes at coverage >0.01× [\(Table 1](#page-1-0) and table S1). They form the most extensive bioarchaeological sampling within a focused temporal and geographical range to date. The examined medieval sites represent burials of individuals from different social and cause of death backgrounds, including urban cemeteries of the charitable poor from the Hospital of St. John, All Saints parish cemetery, Augustinian Friary, Bene't Street plague burial, and rural cemeteries of Cherry Hinton and Clopton [\(Ta](#page-1-0)[ble 1,](#page-1-0) [Fig. 1A,](#page-2-0) and the Supplementary Materials). The analyses of the medieval genomes were performed in context of postmedieval genomes from four sites in Cambridgeshire ([Table 1](#page-1-0)) as well as published genomic data of the Late Iron Age/Roman (*c*. 100 BCE to 400 CE) and Early Saxon periods (*c*. 400–700 CE) from Cambridgeshire and

<span id="page-1-2"></span><span id="page-1-1"></span>elsewhere from England (*[21](#page-11-19)*, *[22](#page-11-20)*). Average endogenous human DNA content was 13% and average contamination rate 1.06%, with 209 individuals under 5%. Average damage in the first 5 base pairs (bp) was 8.02% (table S1). A subset of 143 genomes sequenced to >0.05× coverage were imputed to study the changes in phenotypes related to health and lifestyle. The imputed genomes include 109 individuals with coverage >0.1×, which were subsequently used to resolve genetic ancestry, kinship, recent inbreeding, and heterozygosity.

#### **Genetic ancestry**

The frequencies of mitochondrial DNA (mtDNA) haplogroups in England have remained relatively stable since the Neolithic (table S2). Similarly, principal components analysis (PCA) reveals that all 109 individuals with >0.1× coverage from later medieval Cambridgeshire [\(Fig. 1, A and B\)](#page-2-0) share their autosomal ancestry with modern northern and western European populations without evidence of migration from more distant regions [\(Fig. 1C](#page-2-0) and fig. S1); the same conclusion is supported when projecting pseudohaploid genomes without imputation onto PC space established by modern genomes (figs. S2 to S11). In contrast to genomes from the Roman or Early Saxon periods (*[21](#page-11-19)*, *[22](#page-11-20)*), most later medieval genomes cluster with those from the modern English genomes from the UK Biobank data [\(Fig. 1C](#page-2-0)). Individual outliers who, similarly to most Early Saxon period individuals, are placed among modern Dutch and Danish populations, include a few from Cherry Hinton and the Hospital of St John. Two of them (PSN332 from the Hospital and PSN930 from Cherry Hinton) are also outliers in terms of dental enamel  ${}^{87}Sr/{}^{86}Sr$  values (PSN332 = 0.7122, PSN930 = 0.7108) (*[23](#page-11-21)*). These values, particularly for PSN332, are

<span id="page-1-0"></span>Table 1. Ancient genomes from later medieval and postmedieval Cambridgeshire. Note: N, total number of genomes sequenced per site and by coverage, followed by the numbers assigned to pre– and post–Black Death if the site was in use around the mid-14th century. The pre– and post–Black Death labels are estimated according to whether the burial dates to before or after the year 1348. Some medieval individuals straddle this division and cannot be assigned.



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<span id="page-2-0"></span>**Fig. 1. Sampling locations and genetic ancestry.** (**A**) Sampling locations in Cambridgeshire. (**B**) Zoomed in map of Cambridge. (**C**) PC plot of 109 later medieval and postmedieval genomes with coverage >0.1× in context of ancient (Roman and Saxon period) and modern (UK Biobank) references, with PC1 and PC2 accounting for 0.165 and 0.07% of total variance explained, respectively. (**D**) Supervised Uniform Manifold Approximation and Projection (UMAP) cluster analysis using PiC with modern references based on 5-cM LSAI sharing among modern and 108 ancient genomes, including 80 from later medieval period, with coverage >0.2×. (**E**) Intensity of maximum PiC scores. (F) Transect of time of correlations between the regional PiC vectors. The "Modern" correlation for East Anglia is shown as the correlation between PiC vectors of East Anglia and Bedfordshire/Hertfordshire.

higher than the estimated biosphere  ${}^{87}Sr/{}^{86}Sr$  values for the East of England (*[24](#page-11-22)*), indicating that they did not spend their childhoods in the area local to where they were buried.

<span id="page-2-2"></span><span id="page-2-1"></span>To study the genetic affinity changes across time at finer geographic resolution, we defined interindividual connections by identifying long [>5 centimorgan (cM)] shared allele intervals (LSAIs) with IBIS (*[25](#page-11-23)*) and explored the modularity of individual connectedness (PiC) (*[26](#page-12-0)*) among the historical and modern genomes. Similarly to PCA results, we find that the majority of historical genomes from Cambridgeshire cluster by their connectedness with modern UK Biobank genomes from East England [\(Fig. 1D](#page-2-0) and table S3) whereas a small fraction of later medieval and Roman period genomes, which display low LSAI sharing with any population [\(Fig. 1E\)](#page-2-0), cluster with the UK Biobank donors born in France who also display low levels of LSAI sharing. The Early Saxon period genomes show higher connectedness with Scandinavian genomes, which is also reflected in individual PCA outliers from Cherry Hinton. Overall, we observe regional shifts in

individual connectedness over time ([Fig. 1F\)](#page-2-0). We observe increasing Danish connectedness in the transition from Roman to Early Saxon period; later, during and after the later medieval period, there is an increase of LSAI sharing with both modern Dutch genomes [mirroring documentary evidence showing the Dutch as the most common late medieval immigrants locally (*[27](#page-12-1)*, *[28](#page-12-2)*)] and genomes from a broader zone of England. Last, we identify a major shift in modern East England toward higher LSAI sharing with Wales and Scotland, clearly reflecting the political and economic integration of recent Britain.

<span id="page-2-3"></span>Our analyses of individual connectedness in the People of the British Isles (*[29](#page-12-3)*) data suggest that all later medieval genomes from Cambridgeshire likely draw most of their genetic ancestry broadly from the same sources as present-day central/eastern England population [\(Fig. 2\)](#page-3-0). Although we are able to distinguish certain regional differences in the modern data with our approach, such as between Cornwall and Devon or between North and South Yorkshire, we observe less resolution in a broad area between Lincolnshire and Surrey where our ancient genomes come from [\(Fig. 2\)](#page-3-0). This means that even if some of the individuals had come from Kent or Lincolnshire, for example, we would not be able to detect such fine-scale migration patterns among regions within that area.

Changes in genetic ancestry or selective pressure could cause phenotypic changes over time. We analyzed 214 genomes with >0.05× coverage from the Roman period to the 19th century, including previously published data (*[21](#page-11-19)*, *[22](#page-11-20)*, *[30](#page-12-4)*), for changes in allele frequency of 113 phenotype informative single-nucleotide polymorphisms (SNPs) related to diet, health, and pigmentation (tables S4 to S6). Of 74 SNPs related to health and diet, only 2 involved in autoimmune diseases reached the adjusted significance threshold in the analysis of variance (ANOVA) statistical tests, showing differences between the medieval and postmedieval periods. One SNP, i.e., rs6822844, is an intronic variant in the <span id="page-3-4"></span><span id="page-3-2"></span>*KIAA1109/Tenr/IL2/IL21* block and has been identified as a risk factor in several autoimmune diseases, including celiac disease, rheumatoid arthritis and type 1 diabetes (*[31](#page-12-5)*, *[32](#page-12-6)*). The other variant, i.e., the intronic rs1891467 in the *TGFB2* gene, has been associated with sarcoidosis (*[33](#page-12-7)*). We did not find significant allele frequency changes during and after the later medieval period for the 39 SNPs affecting eye, hair, and skin color included in the HIrisPlex-S set (*[34](#page-12-8)*), which is expected considering the proximity of the later medieval and present-day English in genetic ancestry [\(Fig. 2\)](#page-3-0).

## <span id="page-3-5"></span><span id="page-3-1"></span>**Social landscape**

#### *Kinship and relatedness*

Although the "kinship" bonds that tie together social groups often go beyond or replace "blood-relationships," the types and intensity of



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<span id="page-3-0"></span>**Fig. 2. UMAP plot of individual connectedness among modern and ancient genomes from Britain.** (**A**) Density of maximum PiC score values per individual in one of the extracted communities. (B) UMAP coordinates of the medieval and postmedieval genomes (> 0.2× coverage) from Cambridgeshire. Archaeological site codes as shown in [Fig. 1.](#page-2-0) (**C**) Individual connectedness among modern genomes of the "People of the British Isles" project based on PiC scores of 20 significant communities with more than 10 members extracted from the combined data with the Louvain method (unsupervised cluster analysis). (**D**) Map showing the color codes by counties for the modern genomes used in the UMAP plot A. (**E**) UMAP coordinates of the Iron Age/Roman and Saxon period genomes.

genetic relatedness among individuals buried in the same locality can be informative of the social structure of the population. To study the probability of genetic relatedness among burials of different social backgrounds we used READ-based estimates (*[3](#page-11-2)*) of pairwise differences in autosomes and the X chromosome in 171 later medieval genomes with >0.01× coverage. Among individual pairs with >10,000 overlapping SNPs, 21 cases of first- to third-degree relatedness were detected [\(Fig. 3A](#page-4-0) and table S7). All kinship pairs detected by READ that involved individuals with >0.1× coverage were confirmed in our analyses with IBIS (*[25](#page-11-23)*) to share multiple > 7-cM LSAI segments and kinship coefficient > 0.005 (table S7). Expectedly, considering the time gaps, none of the 97 later medieval individuals showed kinship with 463,855 modern UK Biobank individuals by the same threshold. Nine of the 12 tested postmedieval individuals were found to form a total of 20 fourth- to sixth-degree relationships with modern individuals who identify themselves as British, including one born outside of the United Kingdom. Among the later medieval individuals, we detect 12 cases of more distant form of relatedness within the same archaeological site beyond those identified with READ—10 at Cherry Hinton and 2 at All Saints—while none were found at the Hospital or Friary. We found multiple cases (more than 1% of all pairs considered) in the rural Cherry Hinton and urban All Saints parish cemeteries. In contrast, we detected only one pair of relatives, a middle-aged (46 to 59 years old) friar and a female child (second degree). No kinship relations were found at the Hospital, despite the large sample size analyzed. On average, pairwise differences between individuals from the Hospital were found to be greater than those between individuals from other sites ([Fig. 3B](#page-4-0)), highlighting the heterogeneity of the ancestry of individuals entering the Hospital.

We also searched for runs of homozygosity (ROH) tracks in the imputed genomes using hapROH (*[5](#page-11-3)*) and found that most individuals have no or very few ROH tracks that are longer than 4 cM (table S8), while three individuals (PSN357 from the Hospital, PSN870 from the Baptist Chapel, and PSN412 from Holy Trinity) have up to 150 to 175 cM in ROH, which is compatible with the parents being fourth- to fifth-degree relatives, including second- to third-cousin marriage ([Fig. 3C\)](#page-4-0).



<span id="page-4-0"></span>**Fig. 3. Kinship, genetic diversity and inbreeding.** (**A**) Normalized pairwise differences (P0) in autosomal data and X chromosome for later medieval sites with more than five burials. Each individual data point represents a pair of individuals (from a total of 171 individuals with >0.01× coverage tested), the aggregate coverage of which is reflected by the opacity of the color. Boundaries for the first- to third-degree of relatedness for autosomal data were defined as in (*[3](#page-11-2)*). Error bars with two SEMs are shown for the pairs of first- to third-degree of relatedness only. \*\* and \* correspond to significant differences at *P* < 0.01 and *P* < 0.05 by two-tailed *t* test, respectively. (**B**) Boxplot of normalized autosomal pairwise differences in four Cambridge medieval sites represented by the largest sample size in this study. Each rectangular data point represents a pairwise comparison of individuals sampled from the same site, normalized (with READ) by the average of all pairwise comparisons made in the pool of all later medieval individuals from Cambridge. The opacity of the rectangular color reflects the aggregate of the coverages of the two individuals. The results of significant (*P* < 0.01) two-tailed Wilcoxon rank sum test are shown with \*\*. (**C**) The total lengths of ROH tracks longer than 4 cM in individuals grouped by site, showing highly inbred outliers.

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<span id="page-5-0"></span>**Fig. 4. Heterozygote density and allele frequency differentiation in the HLA locus.** (**A**) Distribution of average heterozygote density in 1-Mbp windows of chromosome 6. Gray line shows the density of heterozygous sites at common variant positions with minor allele frequency higher than 0.05 in the HRC imputation panel in chromosome 6 for 50 imputed (>0.1× coverage) pre– and post–Black Death genomes from Cambridge. The orange line highlights windows containing genes in the HLA locus. (B) Scatter plot of Max(*F<sub>ST</sub>*) - maximum *F<sub>ST</sub>* between before and after the Black Death cohorts - and Het density values by 1-Mbp windows of chromosome 6. (C) Het density in the before ( $n = 31$ ) and after ( $n = 19$ ) the Black Death cohorts.

#### **Before/after the Black Death**

Because of the broad error range of the radiocarbon dates as well as the limited accuracy of the assignments to the pre/post–Black Death groups based on associated finds and position in the sequence of dated stratigraphic context, some individuals in the post–Black Death group might have been born before 1348, although two obvious cases from Bene't Street have been excluded from analysis. This might limit our power to detect changes after the Black Death. None of the individuals in the pre– or post–Black Death groups are among those tested positive for *Yersinia pestis* previously (*[10](#page-11-8)*, *[35](#page-12-9)*, *[36](#page-12-10)*). Our analyses of genetic ancestry [\(Figs. 1](#page-2-0) and [2,](#page-3-0) figs. S1 to S11, and table S2) were unable to detect changes in rates of long-distance migration associated with the Black Death comparable to those recently shown in case of Trondheim population (*[15](#page-11-13)*). However, besides the potential effect on broader regional ancestry, the Black Death pandemic could have left other detectable signatures on genetic diversity of the population at the genome scale, or, it could have affected specific genes and variants associated with infectious disease vulnerability. To

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examine its effect on the genetic diversity of the Cambridge medieval population further, we estimated heterozygosity and nucleotide diversity genome-wide and in the human leukocyte antigen (HLA) locus in imputed genomes.

<span id="page-5-1"></span>Genome-wide heterozygosity and nucleotide diversity are sensitive to demographic events such as bottlenecks, founder events, or admixture. High mortality during the pandemic could be detectable in extreme cases in a small isolated population as a reduction of diversity across all loci. Changes in the HLA region might capture possible signals of selection specifically at immunity loci. If any one or a few variants in this locus responded to selection, they would have been expected to affect the whole region because of linkage. Consistent with the long-term effect of balancing selection on HLA locus, we find this region has higher density of heterozygous positions at common variants and nucleotide diversity in our pool of imputed genomes ([Fig. 4A](#page-5-0) and fig. S13). However, the "before" and "after" the Black Death cohorts do not show higher than average allele frequency differentiation within the HLA region [\(Fig. 4B](#page-5-0)) nor notable differences in the heterozygote density [\(Fig. 4C\)](#page-5-0). Within a subset of 50 imputed genomes assigned to either before or after the Black Death and coverage >0.1×, we observed no significant differences in genomewide (two-tailed *t* test,  $P = 0.205$ ) nor HLA locus ( $P = 0.700$ ) heterozygosities (fig. S12). Similarly, we did not detect changes in nucleotide diversity in the HLA region or genome-wide after the Black Death (fig. S13).

Our analyses of 70 pre– and post–Black Death imputed genomes for changes in allele frequency in 25 variants previously identified as potential targets of selection in humans against viral and bacterial pathogens as well as four variants recently highlighted as selection targets specifically against *Y. pestis* (*[16](#page-11-14)*) revealed (tables S4 and S6) one significant ( $P = 0.003$ ) difference at individual test level at the rs42490 SNP in the *RIPK2* gene. The *RIPK2* allele previously shown to be protective against leprosy (*[37](#page-12-11)*) showed increased allele frequency after the Black Death. This result would not remain, however, significant after applying multiple test corrections and considering the limited sample size of our cohort it requires further validation in an independent dataset.

<span id="page-6-1"></span><span id="page-6-0"></span>None of the four immunity variants identified by Klunk *et al.* (*[16](#page-11-14)*) with significant allele frequency changes both in their London and Danish cohorts were replicated between Cambridge before and after Black Death cohorts (table S9). Using simulations, we demonstrated that this is unlikely to be caused by a lack of power due to our sample size (fig. S14). We did observe a similar enrichment (1.4-fold,  $P = 0.0001$ ) of variants related to immunity among highly differentiated variants ( $F_{ST}$  > 95th percentile) when using the same list of immunity-related and neutrally evolving variants as the authors (tables S9 and S10). Of the 245 highly differentiated immunity-related variants identified in their London cohort, 22 were replicated, significantly more than expected by chance  $(P = 0.0001)$ ; However, 10 of the 22 overlapping variants that are above the 95th threshold in the Cambridge cohort and 2 of the 3 variants above the 99th threshold show opposite directionality of allele frequency change in time in London and Cambridge cohorts (table S9). While the minor allele frequencies of the immunity variants appear to be highly correlated between our studies ( $r = 0.90$ ,  $P < 1 \times 10^{-15}$ ), the  $F_{ST}$  between the pre- and postpandemic cohorts are not ( $r = 0.019$ ,  $P = 0.26$ ). The significant enrichment of immunity genes cannot be reproduced with our data when using the full list of 37,574 neutral regions defined in (*[38](#page-12-12)*) instead of the relatively small number of variants ascertained by Klunk *et al.* (*[16](#page-11-14)*) in its subset of 250 regions (table S10). We observe a reduction (1.14-fold,  $P = 0.29$ ) of high  $F_{ST}$  values among the Klunk *et al.* (*[16](#page-11-14)*) immunity variants when we define the neutral 95th threshold using 55,965 variants from the full range of the 37,574 neutral regions, which becomes significant (1.73-fold,  $P < 1 \times 10^{-10}$ ) when also using an expanded set of immunity variants from InnateDB (table S10). Notably, within the pool of highly differentiated immune locus variants identified by Klunk et al. ([16](#page-11-14)), we observe significant excess of "gwas" variants, i.e., positions that had previously been confirmed to be polymorphic (table S10) over immunity variants ascertained in the exonic regions, suggesting that ascertainment of previously unidentified variants from low-coverage data (as by Klunk *et al.* (*[16](#page-11-14)*) in their "exon" and "neutral" categories) is one possible cause for the disappearance of the signal when we used the full set of neutral regions defined in (*[38](#page-12-12)*) that overlap with positions confirmed to be polymorphic in the Haplotype Reference Consortium (HRC) panel (*[39](#page-12-13)*) to define the threshold.

## **DISCUSSION**

<span id="page-6-3"></span>Our analyses of genetic ancestry in Cambridge through the transect of time revealed, first, notable differences between individuals from different time periods in their placement on PCA and a corresponding increase of LSAIs with continental North Europe between the Roman and more recent periods. These findings partly reflect the cumulatively massive changes in local genetic ancestry in East England that occurred over many centuries in the first millennium CE (*[22](#page-11-20)*, *[40](#page-12-14)*). Most of the individuals that fall outside of the local range of variation come from the rural cemetery at Cherry Hinton, possibly reflecting a diversity of Late Anglo-Saxon, Scandinavian, and Norman ancestries in the process of intermixing. Despite the scale of these ancestry changes, it is difficult to convincingly detect any first-generation migrants from other parts of western Europe using genetic information alone, partially becuse of an absence of comparable reference material from medieval Europe. In case of the Friary, a context where individuals with nonlocal ancestry would have been more likely to be found, it is also possible that some migrating friars living in Cambridge during their life moved back and were buried elsewhere. Second, with the analyses of individual connectedness patterns, we were able to distinguish between modern Dutch, Danish, and Norwegian ancestries and observed a major increase of population-scale connectedness in Cambridge with the Dutch dating to the later medieval period. This pattern could, in principle, reflect gene flow either from the Low Countries to Cambridge or gene flow from East England to the Low Countries, although the former is better supported by documented sources. Last, the major change we observed between medieval and modern periods in connectedness with modern Welsh and Scottish genomes is likely to reflect recent and ongoing migrations and mobility. In contrast to the highly region-specific LSAI sharing between modern and medieval genomes in Estonia (*[26](#page-12-0)*), later medieval Cambridge genomes do not show increased affinity to modern East Anglia compared to other regions. These recent changes in genetic ancestry suggest that modern Biobank sources may not be ideal references for the study of population histories, highlighting the need for aDNA sampling of historic populations from different time points.

<span id="page-6-4"></span>Although we observe major long-term changes in genetic ancestry, we were unable to detect a change in ancestry or genetic diversity between our pre– and post–Black Death cohorts to account for the hypothesized increased mobility after the Black Death caused by labor shortages. The short time span of a few decades, small sample size and relatively low genetic differentiation in central and eastern England [\[Fig. 2;](#page-3-0) (*[29](#page-12-3)*)] would all make detecting short- to mediumrange migration within England challenging. The lack of evidence for long-distance migration is in agreement with earlier mtDNA evidence in medieval British and Danish cohorts (*[41](#page-12-15)*) but in contrast to genome-wide evidence of increased long-distance migration from medieval Trondheim (*[15](#page-11-13)*). A recent isotope study also found that fewer adolescents moved into York from long distance following the Black Death, possibly because of the economic decline of York and the enforcement of labor laws (*[42](#page-12-16)*). These results suggest that the pandemic had a variable impact on human mobility in European cities depending on their regional, socioeconomic, and political context and circumstances.

<span id="page-6-2"></span>We found that all later medieval social groups we examined—the ordinary urban population buried at All Saints, rural population buried at Cherry Hinton, charitable inmates buried at the Hospital of St John, and friars and benefactors buried at Augustinian Friary shared largely the same local genetic ancestry. Although historical record suggests that some friars might have traveled from as far as the Mediterranean to attend the Augustinian studium generale (regional study center) they were either not sampled or buried elsewhere. In contrast to the lack of differentiation by ancestry, we found clear differences in kinship probabilities between the social groups. The lack of close genetic kinship is consistent with the function of the Hospital as a safety net for those without family support, although we note that genetic relatedness does not capture the whole range of social relationships viewed as kinship. Relatedly, the hospital inmates also show elevated background genetic diversity compared to other later medieval sites. Some of them could have arrived from farther away than the average townsfolk, and the lack of local social support might be related to their hardship. Given that friaries maintained long-lasting ties with local benefactor families, it is not unexpected that a female child who was a second-degree relative to one of the friars was allowed to be buried within the Friary, but no other close genetic relatives were found among the Friary population.

<span id="page-7-0"></span>We found a small number of cases of extended ROH tracks in the genome consistent with the parents being second to third cousins. These cases can be considered unexpected because marriage within four degrees of consanguinity was forbidden by the Church (*[43](#page-12-17)*). We speculate that this could result from incomplete church records, especially if the two parents attended different parish churches, or extrapair paternity events in the family history.

The Black Death has been hypothesized to exert selective pressure on genes related to health and immunity (*[14](#page-11-12)*, *[44](#page-12-18)*). If existing genetic variations differ in their resistance against the plague, and if the plague was a major cause of mortality, we would expect a drop in genetic diversity immediately after the pandemic. However, among individuals who lived shortly before or after the Black Death, we did not find changes in heterozygosity in the HLA region. Although the mortality rate (30 to 65%) was devastating, followed by subsequent plague outbreaks until the second half of the 17th century in Britain, it may not have exerted a strong enough uniform selective pressure for a long enough period to leave a detectable signature in the low-coverage genomes. It should be cautioned, though, that conclusions drawn from HLA analyses at SNP and haplotype level may differ (*[45](#page-12-19)*). Immel *et al.* (*[14](#page-11-12)*) identified frequency differences in three HLA haplotypes between 16th-century plague victims in Ellwangen, Germany and the modern local population that were significant at the individual test level (*P* < 0.05) while failing to pass the significance test after multiple test correction. While the signal of heterozygosity change has been recently used to map an ancient selective sweep in the HLA region (*[46](#page-12-20)*), the fact that we were unable to detect significant changes in HLA heterozygosity in our data may be a reflection of the short effect time of the pandemic.

<span id="page-7-4"></span><span id="page-7-3"></span><span id="page-7-2"></span>Similarly, we did not find an excess of variants in genes related to health and immunity to be highly differentiated between the before and after the Black Death cohorts. The most highly differentiated variant in the *RIPK2* gene shows an increase in frequency after the Black Death of an allele protective against leprosy. Notably, this gene has been suggested to also have a role in the recognition of *Y. pestis* by the innate immune response (*[47](#page-12-21)*). Some scholars have suggested that the Second Plague Pandemic contributed to the decline of leprosy starting from the mid-13th century, as many people vulnerable to leprosy fell victim to the plague (*[48](#page-12-22)*, *[49](#page-12-23)*). On the other hand, we could not replicate in our Cambridge cohort the findings by Klunk *et al.* (*[16](#page-11-14)*) of significantly higher differentiation of immune genes, and, more specifically, differentiation of the four SNPs identified in their London

and Danish cohorts (table S4). The signal of enrichment disappears or even reverses direction when we use an extended set of neutral regions as our reference. The high correlation in allele frequency of common variants [minor allele frequency (MAF) > 0.1 in Klunk *et al.*  (*[16](#page-11-14)*) data] between our Cambridge cohort and the London/Danish cohort and the fact that we got qualitatively similar results when using the same variant lists suggest that differences in genotyping, either with or without imputation, are not likely the key factors; our results show that the subset of 250 regions used by Klunk *et al.* (*[16](#page-11-14)*) are likely to contain too few variants to be robustly representative of variation in the full list of 37,574 neutral regions defined by Gronau *et al.* (*[38](#page-12-12)*). Furthermore, Barton *et al.* (*[50](#page-12-24)*) have shown that the enrichment of high *FST* variants at immune loci in Klunk *et al.* (*[16](#page-11-14)*) results is attributable to a statistical artifact caused by different coverage of loci. Although not supporting the results of Klunk *et al. (*[16](#page-11-14)*)* study, our results are in line with other research showing relatively low or moderate differentiation of immunity genes by *FST* (*[51](#page-12-25)*, *[52](#page-12-26)*). Low differentiation is consistent with the expectations of the theory of balancing selection (*[45](#page-12-19)*) as well as poison/antidote model (*[53](#page-12-27)*) whereby infectious disease transmission between communities often co-occurs with mobility and gene flow that locally increase the effective population size of genes involved in host defense. Because immunity genes appear to show generally lower  $F_{ST}$  values than the rest of the genome, our observation of lower differentiation between the pre– and post–Black Death cohorts does not by itself reflect negative selection at the given time point.

<span id="page-7-7"></span><span id="page-7-6"></span><span id="page-7-5"></span><span id="page-7-1"></span>Our limited sample size sets restrictions to identification of selection, yet it is comparable to the one used by Klunk *et al.* (*[16](#page-11-14)*), and although low sequencing coverage could prevent us from identifying individual loci under selection, the polygenic approach taken by Klunk *et al.* (*[16](#page-11-14)*), which we have followed, should in principle be able to detect selection at many variants contributing to a trait. Although our results revealed no enrichment of highly differentiated variants in immunity genes between the before and after the Black Death cohorts, these results do not mean that plague had no selective impact on genetic variation in Cambridge. The immune response to *Y. pestis* might involve multiple pathways which are yet to be fully understood and by our "blind" approach of including all possible immune variants the signal of selection can be buried under the background of allele frequency changes of many loci that are not responsive to plague.

In sum, we have shown that even at very low coverage, wholegenome sequencing of historical genomes when combined with rich historical context and other archaeological evidence can help to reconstruct many aspects of medieval life: the occasional incomers in a rural community; a young child from a benefactor family who was buried in the Friary; the Hospital full of unrelated inmates who might have traveled from afar; the social groups that had little difference in genetic background; a pandemic whose genetic consequence in the Cambridge population remained evasive, despite the devastating effect on individual lives and medieval society. The range will surely expand with advancement in aDNA sequencing technology as well as framework to combine various sources of evidence.

#### **MATERIALS AND METHODS**

#### **Sample information and ethical statement**

A description of the archaeological sites included in this study can be found in the Supplementary Materials. All skeletal elements were sampled with permissions from the representative bodies/host institutions. Samples were taken and processed to maximize research value and minimize destructive sampling. Teeth were sampled from skeletons using gloves. Molars were preferred because of having more roots and larger mass, but premolars were also sampled.

### **Dating**

<span id="page-8-2"></span><span id="page-8-1"></span><span id="page-8-0"></span>The radiocarbon determinations were undertaken at the Scottish Universities Environmental Research Centre radiocarbon laboratory and followed their standard procedures (*[54](#page-12-28)*). Analysis was undertaken using OxCal v.4.3 (*[55](#page-13-0)*, *[56](#page-13-1)*) using the IntCal13 calibration curve (*[57](#page-13-2)*). A small number of additional radiocarbon determinations (marked by \* in table S1) were undertaken at a late stage, to date specific individuals that produced evidence for specific pathogen aDNA. These were undertaken at the 14Chrono Centre, Queen's University Belfast. These additional determinations were calibrated using IntCal 2020 (*[58](#page-13-3)*).

<span id="page-8-4"></span>At the Hospital of St John, Augustinian Friary and All Saints by the Castle, detailed Bayesian modeling of these results combined with expert archaeological judgment has helped to inform whether these individuals (along with others in the stratigraphic sequence) died before or after the Black Death pandemic of 1348–1349. For skeletons for which direct radiometric dates were not available, dating was based on associated finds and relative position within the sequence of dated stratigraphic contexts within a site.

#### **Sampling, aDNA extraction, and library preparation**

Inside a class IIB hood in the dedicated aDNA facility of the University of Cambridge Department of Archaeology or University of Tartu Institute of Genomics, root portions of teeth were removed with a sterile drill wheel. Petrous was sampled with a 10-mm core drill sterilized with bleach followed by distilled water and then ethanol rinse. In some cases, postcranial remains were also sampled (see table S1); in these cases, a small portion was cut off with a sterilized drill wheel and treated like a tooth root/petrous portion except that proteinase K was not added to the extraction buffer. Root and petrous portions were briefly brushed to remove surface dirt, any varnish or lacquer, and microbial film with full-strength household bleach (6% w/v NaOCl) using a disposable toothbrush that was soaked in 6% (w/v) bleach before use. They were then soaked in 6% (w/v) bleach for 5 min. Samples were rinsed twice with 18.2-megohm-cm  $H_2O$  and soaked in 70% (v/v) ethanol for 2 min, transferred to a clean paper towel on a rack inside a class IIB hood with the ultraviolet light on, and allowed to dry. They were weighed and transferred to polymerase chain reaction (PCR)–clean 5- or 15-ml conical tubes (Eppendorf) for chemical extraction.

Inside a class IIB hood, per 100 mg of each sample, 2 ml of 0.5 M EDTA buffer (pH 8.0) (Fluka) and 50 μl of proteinase K (10 mg/ml; Roche) was added. Tubes were rocked in an incubator for 72 hours at room temperature. Extracts were concentrated to 250 μl using Amplicon Ultra-15 concentrators with a 30-kDa filter (Millipore). Samples were purified according to the manufacturer's instructions using buffers from the Minelute PCR Purification Kit (Qiagen) with the following changes: (i) the use of High-Volume spin columns (Roche); (ii) 10× PB buffer instead of 5×; and (iii) samples incubated with EB buffer (Qiagen) at 37°C for 10 min before elution. The columns were transferred to clean, labeled, 1.5-ml Eppendorf tubes. One hundred microliters EB buffer is added to the membrane and centrifuged at 13,000 rpm for 2 min after the 10-min incubation and stored at −20°C. Only one extraction was performed per sample for screening and 30 μl used for libraries.

Library preparation was conducted using a protocol modified from the manufacturer's instructions included in the NEBNext Library <span id="page-8-5"></span><span id="page-8-3"></span>Preparation Kit for 454 (E6070S, New England Biolabs, Ipswich, MA) as detailed in (*[59](#page-13-4)*). DNA was not fragmented and reactions were scaled to half volume; adaptors were made as described in (*[59](#page-13-4)*) and used in a final concentration of 2.5 μM each. DNA was purified on MinElute columns (Qiagen). Libraries were amplified using the following PCR setup: 50 μl of DNA library, 1× PCR buffer, 2.5 mM MgCl<sub>2</sub>, BSA (1 mg/ ml), 0.2 μM inPE1.0, 0.2 mM deoxynucleotide triphosphate (dNTP) each, HGS Taq Diamond (0.1 U/ $\mu$ l), and 0.2  $\mu$ M indexing primer. Cycling conditions were as follows: 5′ at 94°C, followed by 18 cycles of 30 s each at 94°, 60°, and 68°C, with a final extension of 7 min at 72°C. Amplified products were purified using MinElute columns and eluted in 35 μl of EB (Qiagen). Three verification steps were implemented to make sure library preparation was successful and to measure the concentration of double-stranded DNA/sequencing libraries—fluorometric quantitation (Qubit, Thermo Fisher Scientific), parallel capillary electrophoresis (Fragment Analyser, Advanced Analytical), and quantitative PCR.

### **DNA sequencing**

DNA was sequenced using the Illumina NextSeq 500/550 High-Output single-end 75 cycle kit at the University of Cambridge Department of Biochemistry DNA Sequencing Facility. As a norm, 20 samples were sequenced together on one flow cell; additional data were generated for samples over 5% endogenous human content to increase coverage.

### **Mapping**

Before mapping, the sequences of the adapters, indexes, and poly-G tails were removed from read ends and reads shorter than 30 bp were removed using cutadapt-1.11 (*[60](#page-13-5)*).

<span id="page-8-7"></span><span id="page-8-6"></span>The sequences were aligned to the reference sequence GRCh37 (hg19) using Burrows-Wheeler Aligner (BWA 0.7.12) (*[61](#page-13-6)*) and the command *aln* with reseeding disabled.

<span id="page-8-8"></span>After alignment, the sequences were converted to BAM format and only sequences that mapped to the human genome were kept with samtools 1.3 (*[62](#page-13-7)*). Data from different flow cell lanes were merged and duplicates were removed using picard 2.12.0 ([http://broadinsti](http://broadinstitute.github.io/picard/index.html)[tute.github.io/picard/index.html](http://broadinstitute.github.io/picard/index.html), accessed in September 2017). Sequencing coverage was estimated using Qualimap 2.2.1 (*[63](#page-13-8)*) after applying a custom accessibility mask that includes the 1000 genomes accessibility mask and a composite mappability track (*[64](#page-13-9)*).

#### <span id="page-8-10"></span><span id="page-8-9"></span>**aDNA authentication**

<span id="page-8-11"></span>As a result of degradation over time, aDNA can be distinguished from modern DNA by certain characteristics: short fragments and a high frequency of  $C > T$  substitutions at the 5' ends of sequences due to cytosine deamination. The program mapDamage2.0 (*[65](#page-13-10)*) was used to estimate the frequency of  $5^{\circ}$  C  $>$  T transitions. Rates of contamination were estimated from mtDNA using ContamMix (*[66](#page-13-11)*) and from X chromosome using two methods implemented in ANGSD (*[67](#page-13-12)*).

<span id="page-8-13"></span><span id="page-8-12"></span>Samtools 1.3 (*[62](#page-13-7)*) option stats was used to determine the number of final reads, average read length, average coverage, etc. The average endogenous DNA content (proportion of reads mapping to the human genome) was 13.22% (0.00 to 81.27%).

## **Calculating genetic sex estimation**

<span id="page-8-14"></span>Genetic sex was calculated using the methods and script described in (*[68](#page-13-13)*), estimating the fraction of reads mapping to Y chromosome out of all reads mapping to either X or Y chromosome. Genetic sex was calculated for genomess with a coverage >0.01×, and only reads with a mapping quality >30 were counted for the autosomal, X, and Y chromosomes.

### **Determining mtDNA haplogroups**

<span id="page-9-0"></span>mtDNA haplogroups were determined using Haplogrep2 (*[69](#page-13-14)*). Subsequently, the identical results between the individuals were checked visually by aligning mapped reads to the reference sequence using samtools-1.3 (*[62](#page-13-7)*) command tview and confirming the haplogroup assignment in PhyloTree (accessed at: [www.phylotree.org\)](http://www.phylotree.org). In addition, private mutations were noted for further kinship analysis.

### **Y chromosome variant calling and haplotyping**

<span id="page-9-2"></span><span id="page-9-1"></span>A total of 256,463 binary Y chromosome SNPs that have been detected as polymorphic in previous high-coverage whole Y chromosome sequencing studies (*[70](#page-13-15)*–*[72](#page-13-16)*) and by YFull [\(www.yfull.com/](http://www.yfull.com/snp-list/) [snp-list/](http://www.yfull.com/snp-list/)) were called in male individuals with more than  $0.001 \times$  autosomal coverage using ANGSD-0.916 (*[67](#page-13-12)*) "-doHaploCall" option. Basal haplogroup affiliations (table S1) could be determined for 128 individuals by assessing the proportion of derived allele calls (pD) in a set of primary (A, B, C...T) haplogroup defining internal branches, as defined in (*[71](#page-13-17)*), using 4081 informative sites. Haplogroup assignments were confirmed and specified using pathPhynder (*[73](#page-13-18)*). Further detailed subhaplogroup assignments within the phylogeny of the primary haplogroup were determined on the basis of mapping the derived allele calls to the internal branches of the YFull tee [\(www.yfull.](http://www.yfull.com/tree/) [com/tree/](http://www.yfull.com/tree/)), requiring support of at least two variants for the terminal branch assignment.

## <span id="page-9-4"></span>**Pseudohaploid genotype calling**

Autosomal variants were called with the ANGSD 0.917 (*[67](#page-13-12)*) command --doHaploCall, which randomly selects one base at each specified position in the genome. The pseudohaploid genotypes were used in genetic kinship analysis with READ and for PC analyses of individual sites with smartPCA (figs. S2 to S11).

#### **Imputation**

<span id="page-9-8"></span><span id="page-9-7"></span><span id="page-9-6"></span><span id="page-9-5"></span>Following (*[74](#page-13-19)*), genotype likelihoods were first updated with Beagle 4.1 (*[75](#page-13-20)*) from genotype likelihoods produced by ANGSD 0.917 (−doMajorMinor 3 -GL 1 -doPost 1 -doVcf 1 -sites …) (*[67](#page-13-12)*) in Beagle -gl mode, followed by imputation in Beagle -gt mode with Beagle 5 (*[76](#page-13-21)*) from sites where the genotype probability (GP) of the most likely genotype reaches 0.99. To balance between imputation time and accuracy, we used 503 Europeans genomes in 1000 Genomes Project Phase 3 (*[77](#page-13-22)*) as the reference panel in Beagle -gl step, and 27,165 genomes (except for chromosome 1, where the sample size is reduced to 22,691 because of a processing issue in the release) from the HRC (*[39](#page-12-13)*) in the Beagle -gt step. After filtering out rare variants with MAC  $<$  5, we retained a total of  $\sim$  37 million variants genomewide. Because Beagle treats "./." in the variant call format (VCF) input as sporadically missing and imputes them during haplotype phasing, which damages the accuracy when such missing genotypes are common, we imputed each genome individually so that missing genotypes were not included in the VCF input to Beagle 5. The output single-individual VCFs were then merged for downstream analysis. Apart from newly generated genomes in this study, we also imputed published genomes from (*[21](#page-11-19)*, *[22](#page-11-20)*, *[30](#page-12-4)*).

<span id="page-9-9"></span>We down-sampled a 10.83 $\times$  genome that has been previously reported in  $(78)$  $(78)$  $(78)$ , PSN31, to  $0.05 \times$  to  $1 \times$  to evaluate our imputation <span id="page-9-10"></span>pipeline (table S11). The accuracy was estimated by comparing the imputed genotypes to genotypes called using GATK HaplotypeCaller (*[79](#page-13-24)*) (with the flags --min-pruning 1 --min-dangling-branchlength 1).

For some analysis, we filtered the imputed genotypes further by GP and/or minor allele frequencies at the sites as described in the respective sections. In general we applied a coverage cutoff of at least  $0.1 \times$  for analyses requiring individual genotypes and lower cutoffs where allele frequency is of more concern. We also sought to address the questions using both imputed and genotyped data whenever possible.

### **Principal components analysis**

<span id="page-9-11"></span><span id="page-9-3"></span>We used FlashPCA2 (*[80](#page-13-25)*) for PCA of imputed genomes (without projection) together with modern reference genomes from selected groups in UK Biobank after excluding variants in linkage disequilibrium with the PLINK --indep-pairwise 1000 50 0.5 option and exclusion of the likely non-neutral regions exclusion\_regions\_hg19.txt [\(Fig. 1C](#page-2-0) and fig. S1). To check for possible artifacts introduced by imputation, we also performed PCA using modern reference genomes only, before projecting pseudo-haploid genomes (coverage >0.05×) onto the PC space following ([https://github.com/chrchang/eigensoft/](https://github.com/chrchang/eigensoft/blob/master/POPGEN/lsqproject.pdf) [blob/master/POPGEN/lsqproject.pdf](https://github.com/chrchang/eigensoft/blob/master/POPGEN/lsqproject.pdf), accessed in April 2020) (figs. S2 to S11). The error bars of the projected genomes represent 1 SD obtained from a 20-fold block jackknife. The results using imputed genomes and pseudohaploid genotypes are highly consistent with largely overlapping outliers (apart from those only included in PCA with lsq projection due to low coverage).

## **Detecting IBD and LSAI segments**

Identical by descent (IBD)/LSAI segments (*[26](#page-12-0)*) and kinship coefficients were estimated from merged plink files of 80 (coverage >0.2×) imputed ancient genomes, 503 Europeans from the 1000 Genome Project and UK Biobank data with IBIS version 1.20.9 (*[25](#page-11-23)*) using minimum shared segment length (−min\_L) thresholds 5 and/or 7 cM together with -maxDist 0.1 and -mt 300 parameters. In total, 269,319 binary SNPs with MAF > 0.05 were used.

Although IBIS has the highest IBD inference accuracy for >7-cM segments, which we use in kinship analyses, we use >5-cM threshold in our diachronic inferences of population affinities because our focus is on relationships at generational distances >15 at which longer IBD sharing expectations become relatively low, particularly in combination with the loss of sensitivity to detect long IBD segments from imputed aDNA sequences. Because true IBD segments of this length are not expected to be common at these generational distances, we need to consider the detected segments as "long shared allele intervals" (LSAIs) rather than IBD segments sensu stricto. Because they are inferred from unphased data after removal of rare variants (which cannot be imputed with sufficient accuracy), the LSAIs are likely to include undetected recombination points and smaller IBD segments residing on different haplotypes.

The PiC score (*[26](#page-12-0)*) for individual x in group Z was estimated as the proportion of individuals from group Z with whom individual x shared IBD above the given threshold. In practice, we estimated the count of connected individuals from group Z from sorted IBIS .coef output files by using the linux "join" function to add group codes to individual identifiers and by using the "crosstab" function of datamash (*[81](#page-13-26)*) to generate the table of counts, each of which we divided by the total number of individuals in group Z to obtain the individual connectedness proportions by groups (the PiC scores).

<span id="page-10-0"></span>Unsupervised community extraction analyses on IBIS .coef files calculated from merged plink files including imputed ancient genomes with coverage >0.2× and modern UK Biobank data were run with the Louvain algorithm (*[82](#page-13-27)*) implemented in the R library "igraph" (*[83](#page-13-28)*) with additional significance tests as described in (*[26](#page-12-0)*) using scripts available at <https://github.com/SABiagini/Louvain> (accessed in October 2021).

#### **Genetic kinship analysis**

Pseudohaploid genotypes of 171 (coverage >0.01×, not including two individuals from the plague pit at Bene't Street) later medieval genomes at 5,494,912 positions with MAF > 0.05 in the UK10K subset of the HRC panel were called with ANGSD version 0.917. For the comparison with published studies, the merged PLINK file from the PCA analysis was used and select populations retained using plink --keep and converted to .tped. The ANGSD output files were converted to .tped format, which was used as an input for kinship analyses with READ (*[3](#page-11-2)*). Given highly homogenous ancestry in all study sites, individuals from all sites were analyzed together for [Fig. 3](#page-4-0) as well as separately to test for potential biases in normalized P0 calculation. In addition to first- and second-degree relationships, we also estimated P0 cutoffs  $[15/16 = 0.9375$  $[15/16 = 0.9375$  $[15/16 = 0.9375$  as per  $(3)$ ] for the detection of thirddegree relatives while acknowledging that because of lack of relevant empirical data the false-positive and false-negative error rates for this class of relationship remain unknown.

#### **Runs of homozygosity**

We used hapROH (*[5](#page-11-3)*) to detect ROH in 109 ancient genomes with coverage >0.1×. Using information from a reference panel, hapROH has been shown to work for genomes with more than 400K of the 1240K-SNP panel covered at an error rate lower than 3% in pseudohaploid genotypes (*[5](#page-11-3)*). We note that the requirement is broadly in line with the imputation accuracy we get from coverages as low as 0.1×, where ~60% of common variants (MAF  $\geq$  0.05) in the HRC panel are recovered with an overall accuracy around 97% for diploid genotypes (table S11). Among common variants in the HRC panel, 853,159 overlap with the 1240K-SNP panel.

1000 Genomes Project data were used to construct the reference haplotypes. We kept the standard parameters in hapROH, which had been optimized for 1240K aDNA genotype data

e\_model='haploid', post\_model='Standard', random allele=True, roh in=1, roh out=20, roh jump=300, e rate=0.01, e rate ref=0.0, cutoff post=0.999, max\_gap=0, roh\_min\_l=0.01

We also tested the effect of imputation using down-sampled genomes of PSN31. There is an excess of inferred short  $(< 4 \text{ cM})$ ROH tracks at 0.05×, but the results are similar at 0.1× and above (table S8).

## **Heterozygosity**

<span id="page-10-1"></span>Genome-wide heterozygosity and heterozygosity in the HLA region (--chr 6 --from-kb 28477 --to-kb 33448) were estimated with the --het function in PLINK 1.9 (*[84](#page-13-29)*) for imputed genomes with >0.05× coverage and for 5,448,740 sites that had MAF > 0.05 in the HRC reference panel. To assess the impact of imputation accuracy on heterozygosity estimation we plotted the heterozygosity estimates of imputed genomes against their coverage (fig. S15) and observed weakly negative but nonsignificant correlation ( $r = -0.05$ ,  $P = 0.58$ ) and the highest variance in the group of genomes imputed from the lowest (0.05× to

0.1×) coverage. We retained for further analyses only genomes with >0.1× coverage.

#### **Nucleotide diversity**

<span id="page-10-2"></span>Nucleotide diversity in the HLA region and throughout the autosomes was estimated using vcftools --window-pi (*[85](#page-13-30)*), which by default outputs result in 10-kb windows, in imputed genomes with >0.1× coverage. Sites where the highest genotype probability is less than 0.99 were set to missing. The results are similar whether we filtered the variants by MAF > 0.05 in the HRC reference panel or not (fig. S13).

### **Phenotype prediction**

<span id="page-10-6"></span><span id="page-10-5"></span><span id="page-10-4"></span><span id="page-10-3"></span>Using the imputed genomes generated in this study (143 individuals with a coverage higher than 0.05×) and previously published (71 individuals) (*[21](#page-11-19)*, *[22](#page-11-20)*, *[30](#page-12-4)*), we extracted genotype calls for 39 of the 41 HIrisPlex-S variants and 74 SNPs involved in diet and diseases, coding the allele information as the number of the effective allele (0, 1, 2) using plink (tables S4 to S6). The diet and disease set of 74 SNPs was selected starting from lists of variants previously analyzed in aDNA studies (*[86](#page-13-31)*–*[88](#page-13-32)*), prioritizing those with a role in the response to pathogenic infection. In addition to the 70 variants that we have previously used for phenotype prediction (*[89](#page-13-33)*, *[90](#page-13-34)*), we also analyzed the four SNPs in immune loci recently detected to be highly differentiated before or after the Second Pandemic (*[16](#page-11-14)*) (table S6). A table with the HIrisPlex-S SNP alleles per individual was uploaded on the HIrisPlex-S webtool (<https://hirisplex.erasmusmc.nl/>) to obtain probabilities values for each eye, hair, and skin color category per individual. This output was then interpreted following the manual to obtain the final pigmentation prediction for each individual (table S5). We then grouped the individuals by time period. The groups were compared performing an ANOVA statistical test, applying a Bonferroni's correction on each group of variants (carbohydrate metabolism, lipid metabolism, vitamin metabolism, response to pathogens, autoimmune diseases, other diseases, and pigmentation) (table S4) to set the significance threshold. For only the significant SNPs, we also performed a post hoc Tukey test to identify the significantly different group pairs. For our medieval genomes, we also grouped them by relative time to the Second Pandemic (before or after) and performed a statistical *t* test as reported in table S4.

#### **Enrichment of immune genes**

<span id="page-10-7"></span>To test the enrichment for higher allele frequency differences at vari-ant positions of immune genes we used Weir and Cockerham ([91](#page-13-35))  $F_{ST}$ as implemented in PLINK (--fst case-control) on lists of variants derived from the Klunk *et al.* study (*[16](#page-11-14)*), as well as an expanded list of 54,931 variants polymorphic in the HRC panel from the 37,574 putatively neutral regions defined by Gronau *et al.* (*[38](#page-12-12)*) that had MAF > 0.1 in 70 imputed genomes from Cambridge dating to before and after the Black Death. As an expanded set of immunity regions, we extracted 19,940 variants with  $MAF > 0.1$  in the Cambridge cohort from 189,173 exonic regions of the 4723 innate immunity genes curated by [InnateDB](https://www.innatedb.com/moleculeSearch.do) [\(www.innatedb.com/](http://www.innatedb.com/)).

To explore how our sample size (before Black Death: *n* = 44; after Black Death:  $n = 26$ ) might limit our power to detect extreme  $F_{ST}$  values, we postulated a population of 10,000 individuals from before the Black Death and another of the same size from after the Black Death. For each combination of allele frequencies before and after the Black Death (on a grid of  $100 \times 100$  points), assuming Hardy-Weinberg equilibrium, we sampled from these large populations according to our sample sizes and calculated the observed *F<sub>ST</sub>* values. The sampling was repeated 10,000 times. The power of the analysis was calculated as the proportion of observed *FST* values that exceed a predefined threshold, which was taken from the 95th percentile of the distribution of *F<sub>ST</sub>* from a set of neutral variants according to Klunk *et al.* (*[16](#page-11-14)*).

We note that we already deviate from the authors if we choose the threshold using our own dataset, regardless of considerations about sample size. The variants within the same neutral region used by Klunk *et al.* (*[16](#page-11-14)*) produced a much higher 95th percentile threshold, whether we filter out variants with a minor allele frequency lower than 0.1 (0.027; 95% confidence interval: [0.022, 0.031]) or not (0.030; 95% confidence interval: [0.028, 0.038]). The threshold is even higher when we include variants ( $MAF > 0.1$ ) in the full list of neutral regions described by Gronau *et al.* (*[38](#page-12-12)*) (0.035; 95% confidence interval: [0.035, 0.036]). None of the  $F_{ST}$  values of the four variants reported by Klunk *et al.* ([16](#page-11-14)) in their London dataset would pass these thresholds. We believe this is related to the ascertainment of previously unidentified variants from low-coverage data in their study as mentioned in our main text.

Nevertheless we went on to adopt their threshold (0.0089) for our power analysis. We ask "what is the probability of obtaining an *FST* value greater than 0.0089 in our samples given each combination of true population allele frequencies before and after the Black Death." The results are shown in a heatmap (fig. S14). The gray zone along the diagonal line is the region where the true  $F_{ST}$  is below 0.0089. Expectedly, the closer the true allele frequencies, the lower the power of the analysis. The four squares in the plot correspond to the allele frequencies of the four highly differentiated variants estimated in their London cohort according to Klunk *et al.* (*[16](#page-11-14)*), which are replicated in their Danish cohort. If we assume that our Cambridge cohorts derive from the same meta-population as London, and that the allele frequencies estimated from the pre- and postplague London cohorts reported by Klunk *et al.* (*[16](#page-11-14)*) accurately reflect the allele frequencies in this meta-population, the power to obtain *F<sub>ST</sub>* scores higher than 0.0089 given our sample size ranges between 0.50 and 0.71 for those variants highlighted by Klunk *et al.*  (*[16](#page-11-14)*). Under these assumptions, however, the probabilities of obtaining the small *FST* values as we observe in the Cambridge cohort (table S9) range from low (rs2549794: 0.16; rs17473484: 0.19) to very low (rs1052025: 0.013; rs11571319: 0.016), suggesting that our failure to replicate high differentiation at these variants is unlikely due to our sample size.

#### **Supplementary Materials**

**This PDF file includes:** Supplementary Text Figs. S1 to S15 Legends for tables S1 to S11 References

**Other Supplementary Material for this manuscript includes the following:** Tables S1 to S11

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