Infectious diseases are among the strongest selective pressures driving human evolution. This includes the single greatest mortality event in recorded history, the first outbreak of the second pandemic of plague, commonly called the Black Death, which was caused by the bacterium *Yersinia pestis*. This pandemic devastated Afro-Eurasia, killing up to 30–50% of the population. To identify loci that may have been under selection during the Black Death, we characterized genetic variation around immune-related genes from 206 ancient DNA extracts, stemming from two different European populations before, during, and after the Black Death. Immune loci are highly differentiated sites relative to a set of non-immune loci, suggesting positive selection. We identify 245 variants that are highly differentiated within the London dataset, four of which were replicated in an independent cohort from Denmark, and represent the strongest candidates for positive selection. The selected allele for one of these variants, rs2549794, is associated with the production of a full-length (versus truncated) ERAP2 transcript, variation in cytokine response to *Y. pestis* and increased ability to control intracellular *Y. pestis* in macrophages. Finally, we show that protective variants overlap with alleles that are today associated with increased susceptibility to autoimmune diseases, providing empirical evidence for the role played by past pandemics in shaping present-day susceptibility to disease.

Infectious diseases have presented one of the strongest selective pressures in the evolution of humans and other animals. Not surprisingly, many candidates for population-specific positive selection in humans involve immune response genes, consistent with the hypothesis that exposure to new and/or re-emerging pathogens has driven adaptation. However, it is challenging to connect signatures of natural selection with their causative pathogens unless the underlying loci are still associated with susceptibility to the same pathogen in modern populations. Clarifying the dynamics that have shaped the human immune system is key to understanding how historical diseases contributed to disease susceptibility today.

We sought to identify signatures of natural selection in Europeans imposed by *Yersinia pestis*, the bacterium responsible for bubonic plague. The first recorded plague pandemic began with the Plague of Justinian in AD 541 (refs. 1,2). Nearly 800 years later, the Black Death (1346–1350) marked the beginning of the second pandemic of plague, which spread throughout Europe, the Middle East and Northern Africa, reducing the population by up to 30–50%3,4. With no recent exposure to *Y. pestis* variation in cytokine response to *Y. pestis* and increased ability to control intracellular *Y. pestis* in macrophages. Finally, we show that protective variants overlap with alleles that are today associated with increased susceptibility to autoimmune diseases, providing empirical evidence for the role played by past pandemics in shaping present-day susceptibility to disease.
plague, Europeans living through the Black Death probably represented immunologically naive populations with little to no prior adaptation to *Y. pestis*. The high mortality rate suggests that genetic variants that conferred protection against *Y. pestis* infection might have been under strong selection during this time. Indeed, the nearly decadal plague outbreaks over the subsequent four hundred years of the second pandemic in Europe often (but not always) were associated with reduced mortality rates\(^{1,13}\), which could have been due to pathogen evolution or changing cultural practices, but potentially also linked to human genetic adaptation to *Y. pestis*.

**Positive selection on immune genes**

Genomic targets of selection imposed by *Y. pestis* during the Black Death, if present, have remained elusive\(^{13-15}\). To better identify such loci, we characterized genetic variation from ancient DNA extracts derived from individuals who died shortly before, during or soon after the Black Death in London and across Denmark. This unique sampling design differentiates, to the greatest extent possible, signatures due to *Y. pestis* from those associated with other infectious diseases or other selective processes (although we cannot exclude these entirely). From London, individuals were sampled from three cemeteries close to one another, tightly dated by radiocarbon, stratigraphy and historical records to before, during and after the Black Death (Fig. 1, Supplementary Table 1 and Supplementary Methods). From Denmark, individuals were sampled from five localities, geographically spread across the country, which were dated via archaeological means (such as burial arm positions), stratigraphy and historical records. We grouped all individuals into those that lived pre-Black Death (London: around AD 1000–1250, Denmark: around AD 850 to around AD 1350) and post-Black Death (London: AD 1350–1539, Denmark: around AD 1350 to around AD 1800). Within London, we also included individuals buried in the plague cemetery, East Smithfield, all of whom died during a two-year window of the Black Death between 1348 and 1349 (ref.\(^{16}\)). Analysis of the mitogenomic diversity from these individuals identifies solely European mitogenomic haplotypes, avoiding a possible confound between natural selection and population replacement from non-European sources\(^{17}\).

In total we screened 516 samples (\(n = 318\) from London; \(n = 198\) from across Denmark) for the presence of human DNA using a modified polymerase chain reaction (PCR) assay for the single copy nuclear c-myc gene\(^{17,18}\) and identified 360 with sufficient endogenous DNA content for downstream enrichment and sequencing of additional nuclear loci (Supplementary Methods). As many of our samples were poorly preserved and had low endogenous DNA content, we used hybridization capture to enrich for and sequence 356 immune-related genes, 496 genome-wide association study (GWAS) loci previously associated with immune disorders and 250 putatively neutral regions (1.5 kb each), as defined by Gronau and colleagues\(^{19}\) (Supplementary Table 2; Supplementary Methods). The targeted immune genes were manually curated on the basis of their role in immune-related processes, and
include innate immune receptors, key immune transcription factors, cytokines and chemokines, and other effector molecules (Supplementary Table 3). To ensure that deamination and other forms of ancient DNA damage did not lead to spurious genotype calls, we trimmed 4 base pairs (bp) from the start and end of each sequencing read (Supplementary Fig. 1) and excluded all singleton variants (n = 106,757). Our final dataset contained 33,110 biallelic variants within the targeted regions (2,669 near GWAS loci, 19,972 in immune genes and 10,469 in putatively neutral regions), with a mean coverage of 4.6± reads per site per individual (see Supplementary Table 1 for per-individual coverage). We further filtered our results by excluding samples with missing genotype calls at more than 50% of those sites (returning n = 206 individuals, Fig. 1c) and excluding variants with genotype calls for less than 10 individuals per time period and population. Using genotype likelihoods, we then calculated the minor allele frequency (MAF) per population at each time point. Finally, we retained only sites with a mean MAF (averaged across London and Denmark) greater than 5% (n = 22,868 sites), as our power to detect selection for variants below 5% is very low (Supplementary Fig. 2).

To detect alleles that may have conferred protection from, or susceptibility to, *Y. pestis*, we searched within candidate regions (immune genes and GWAS loci) for variants that exhibit unexpectedly large changes in allele frequency between pre- and post-Black Death samples. Specifically, we identified alleles for which the degree of differentiation ( examines whether sequence diversity increases or decreases in frequency among individuals sampled post-Black Death in comparison with frequency among people who died during the Black Death and should oppose frequency patterns before, during and after the Black Death. 

Using this reasoning, we narrowed down our list of putatively selected loci from 245 to 35 (Supplementary Table 4). Finally, we asked if these loci were also highly differentiated before and after the Black Death in our Danish replication cohort (that is, among the top 10% most highly differentiated sites, and in the same direction as seen in London). Four loci met these criteria, representing the strongest candidates for selection (Fig. 2c–g). We calculated the selection coefficient (s) for each of these variants using a Hidden Markov Model (HMM) framework (based on ref. 20, Supplementary Methods). Statistical support for non-neutral evolution (s > 0) among our four candidate loci is strong when compared to that of neutral loci (P < 0.001 for each locus; Supplementary Table 5). Despite the large confidence intervals—a inherent limitation when trying to estimate s over a few generations—the absolute values for the point estimate of s range from 0.26 to 0.4, which are among the strongest selective coefficients yet reported in humans, to our knowledge (Extended Data Fig. 2 and Supplementary Table 5).

Functional dissection of candidate loci

None of our top candidate variants overlaps with (nor is in strong linkage disequilibrium with) coding variants, although one, near the ERAP2 gene, is strongly linked to a variant that affects splicing24,25. Their selective advantage may stem from an impact on gene expression levels, particularly in immune cell types that participate in the host response to *Y. pestis* infection. Macrophages in particular are recruited to sites of infection, where they interact with bacteria and contribute to plague resistance26,27. Macrophages phagocyze *Y. pestis*, but some bacteria survive and spread to the lymph node, where they replicate uncontrollably26,27. To test whether the four candidate loci we identified, or genes near them, are involved in the transcriptional response to *Y. pestis*, we incubated monocyte-derived macrophages from 33 individuals with heat-killed *Y. pestis* and compared their expression profiles to unstimulated control samples using RNA sequencing (Supplementary Methods). As expected, macrophages responded robustly to *Y. pestis*, such that the principal component 1 of the gene expression data, which separates baseline versus *Y. pestis* stimulated conditions, explains 56% of the total variance (Extended Data Fig. 3).

Seven genes within 100 kb of our four candidate loci were expressed in this dataset: locus 1 (rs2549794): ERAP1, ERAP2, LNPEP; locus 3 (rs11571319): CTLA4, ICOS; and locus 4 (rs17473484): TICAM2, TMED7, NFATC1, the only gene within 100 kb of locus 2 (rs1052025), was not expressed in this dataset. With the sole exception of LNPEP, all of these genes were differentially expressed in response to *Y. pestis* stimulation (Fig. 3a), supporting their putative role in the host response. Macrophages from an additional panel of eight individuals infected with live and fully virulent *Y. pestis* showed similar directional changes in gene expression to those observed in response to heat-killed bacteria. This was true genome-wide (r = 0.88, P < 1 × 10−4, Extended Data Fig. 4a) and for genes near our four candidate loci (ERAP1 is an exception; it was upregulated in response to live bacteria but downregulated in response to heat-killed bacteria, Extended Data Fig. 4b). To investigate whether changes in gene expression were specific to *Y. pestis* or shared with other infectious agents, we analysed gene expression data from macrophages infected with live *Listeria monocytogenes* (a Gram-positive bacterium) and *Salmonella typhimurium* (a Gram-negative bacterium, like *Y. pestis*),22 as well as monocytes activated with bacterial and viral ligands targeting the Toll-like receptor (TLR) pathways (TLR1/2, TLR4 and TLR7/8) and live influenza virus28. These data show that all genes near our candidate loci (with the exception of CTLA4) respond to other pathogenic agents but that the direction of change in expression differs depending on the stimulus. For example, ERAP2 is downregulated in response to all live bacteria or bacterial stimuli, including *Y. pestis*, but is upregulated in response to viral ones (Extended Data Fig. 5).

Having established that genes near our candidate loci show a transcriptional response to *Y. pestis* in macrophages, we next asked whether genetic variation at each of the four candidate loci is associated with gene expression levels at nearby genes (Fig. 3b,c). We identified an association between rs17473484 genotype and TICAM2 expression in which...
comparing the pre-Black Death (BD) population to the post-BD population in differentiated sites in functional regions relative to neutral regions when both London and Denmark after the Black Death, and the opposite direction for individuals who died during the Black Death (Supplementary Table 4)).

**Fig. 2** | Positive selection at immune loci. a,b. Enrichment of highly differentiated sites in functional regions relative to neutral regions when comparing the pre-Black Death (BD) population to the post-BD population in London (a) and Denmark (b). c, $F_{ST}$, between London before and after the Black Death, limited to the 535 sites that show qualitative patterns consistent with natural selection (namely allele frequency changes in the same direction in both London and Denmark after the Black Death, and the opposite direction for individuals who died during the Black Death (Supplementary Table 4)). Manhattan plot showing loci with patterns indicative of positive selection. Point size and colour intensity (which alternates by chromosome) represents the $-\log_{10} P$-value comparing populations in London before and after the plague, points coloured in orange represent the four positions and their associated genes, which are highly differentiated in Denmark as well. d–g. Patterns of allelic change over time for the four strongest candidates for positive selection. Error bars represent the standard deviation based on bootstrapping individuals from that population and each time point 10,000 times. Allele frequencies for London are shown in red and for Denmark are shown in blue. Modern allele frequencies are derived from 1000 Genomes data for Great Britain in London.

The protective allele is associated with higher expression of the gene in the unstimulated condition (Fig. 3b; $P = 2.5 \times 10^{-10}$ but not after $Y. pestis$ stimulation ($P = 0.24$). This effect is intriguing as $TICAM2$ encodes an adaptor protein for $TLR4$. In vivo, $TLR4$ detects $Y. pestis$ via the recognition of lipopolysaccharides (LPS) on the bacterial outer membrane. $Y. pestis$ attempts to circumvent this detection by decaying surface LPS, thereby reducing the binding affinity for $TLR4$ (ref. 30). $TICAM2$ ushers LPS-bound $TLR4$ into endosomes and activates type I interferon responses. It is therefore possible that increased $TICAM2$ expression confers protection against $Y. pestis$ by increasing sensitivity to LPS and promoting an effective immune response.

The strongest association we identified was between rs2549794 and $ERAP2$ expression, in which the protective allele (C) is associated with a fivefold increase in expression of $ERAP2$ relative to the putatively deleterious T allele (Fig. 3c), in both unstimulated ($P = 4.4 \times 10^{-10}$) and $Y. pestis$-challenged ($P = 8.7 \times 10^{-8}$) cells. We observed similarly strong associations in macrophages and monocytes infected with other pathogens ($Salmonella$, $Listeria$, influenza) or stimulated with $TLR$-activating ligands (all $P < 1 \times 10^{-10}$; Extended Data Fig. 6). This pattern also generalizes to $Y. pestis$-infected peripheral blood mononuclear cells (PBMCs) more broadly, suggesting that rs2549794 is associated with $ERAP2$ expression levels regardless of the infectious/inflammatory stimulus or cell type. In detail, we generated single-cell RNA-sequencing data from PBMCs from ten individuals (five homozygous for the selectively favoured rs2549794 C allele and five homozygous for the alternate T allele), both at baseline and after infection with live, fully virulent $Y. pestis$. Across all immune cell types profiled—B cells, CD4$^+$ T cells, CD8$^+$ T cells, natural killer cells and monocytes (Extended Data Fig. 7)—we identified 5,570 genes for which infection with $Y. pestis$ significantly alters gene expression levels (314 to 4,234 genes per cell type, 10% false discovery rate; Supplementary Table 6). Most genes near our candidate loci are differentially expressed in response to $Y. pestis$ infection, but both the magnitude and direction of such effects is cell type-specific (Extended Data Fig. 8). For example, $ERAP2$ is upregulated upon stimulation in all PBMC cell types (Fig. 3d,e), but is downregulated in monocyte-derived macrophages. These differences could be due to differences in the transcription factors and enhancers active in each cell type, differences in our infection models (PBMCs versus monocyte-derived macrophages), or both. Notably, in all cell types and under all conditions, the selectively favoured rs2549794 C allele is associated with increased $ERAP2$ expression compared to the alternative T allele.

The $ERAP2$ locus is characterized by two haplotypes (A and B) that are common around the world. Haplotype A encodes the canonical (full-length) ERAP2 protein consisting of 960 amino acids; Haplotype B is characterized by the presence of the G allele of rs2248374, a splice-site-variant leader that leads to the production of a splice isoform with an elongated exon 10 containing two premature

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stop codons. This ERAP2 isoform undergoes nonsense-mediated decay (NMD), resulting in undetectable ERAP2 protein levels\(^{21}\). Even when produced, the shorter protein encoded by Haplotype B has limited aminopeptidase activity\(^{33}\). The selectively favoured rs2549794 C allele is in strong linkage with allele A of rs2248374 (\(R^2 = 0.8, D' = 1.0\)) of Haplotype A, which encodes the full-length ERAP2 protein, whereas the deleterious rs2549794 T allele is in linkage with the rs2248374 G allele of Haplotype B, which encodes the truncated version of ERAP2. Using real-time PCR specific to the mRNA encoded by Haplotype B, we found that in macrophages, the decreased ERAP2 expression in individuals harbouring the deleterious rs2248374 G allele is coupled with a higher expression of the truncated isoform known to undergo NMD (Fig. 3f, \(P = 2.66 \times 10^{-6}\)). PCR amplification of exon 10 further confirmed that individuals homozygous for the protective rs2248374 A allele only express Haplotype A, whereas individuals homozygous for the rs2248374 G allele almost exclusively express the truncated Haplotype B (Extended Data Fig. 9).

ERAP1 and ERAP2 are aminopeptidases that work synergistically to trim peptides for presentation to CD8\(^+\) T cells by major histocompatibility complex (MHC) class I molecules\(^{35}\). Given their central role in antigen presentation, it is not surprising that polymorphisms near these genes, including rs2248374, have been associated with susceptibility to a variety of infectious agents\(^{36,37}\). ERAP2 deficiency leads to a significant remodelling of the repertoire of antigens that are presented by MHC to CD8\(^+\) T cells\(^{36,37}\), including MHC ligands that have high homology to peptide sequences derived from \(Y. pestis\) species\(^{32}\). Having ERAP2-mediated aminopeptidase activity plausibly helps to promote the presentation of a more diverse array of \(Y. pestis\)-derived antigens to CD8\(^+\) T cells, which
in turn has an important role in protection against infection\textsuperscript{38,39}. Indeed, mice depleted of CD8\textsuperscript{+} T cells all die within one-week post-infection with a milder \textit{Yersinia} spp., \textit{Y. pseudotuberculosis}, whereas all wild-type mice survive\textsuperscript{39}. Unfortunately, rodent models only possess a single ERAP aminopeptidase that is homologous to ERAP1, limiting our ability to directly test the function of ERAP2 on antigen presentation and response to \textit{Y. pestis} in vivo.

In addition to its canonical role in antigen presentation and CD8\textsuperscript{+} T cell activation, ERAP2 is also involved in viral clearance and cytokine responses\textsuperscript{40}. We therefore sought to test whether ERAP2 genotype is associated with variation in the cytokine response to \textit{Y. pestis} infection. To do so, we infected a further set of monocyte-derived macrophages from 25 individuals (9 homozygous for the selectively favoured ERAP2 haplotype, 9 heterozygous and 7 homozygous for the deleterious haplotype) with live, virulent \textit{Y. pestis} and measured the protein levels of ten cytokines involved in various aspects of the immune response, at baseline and at 24 h postinfection. No differences in cytokine levels existed at baseline (all \(P > 0.05\); Supplementary Table 7), but four cytokines showed a significant association with ERAP2 genotype upon stimulation. Specifically, the levels of granulocyte colony-stimulating factor (\(P = 0.0155\)), interleukin (IL)-1\( \beta \) (\(P = 0.00248\)) and IL-10 (\(P = 0.0216\)) significantly decreased with the number of protective rs249794 alleles found in each individual: 0, 1 or 2 and the percentage of bacteria killed.

We identified four loci that were strongly differentiated before and after the Black Death in London and replicated in our Danish cohort as the strongest candidates for selection. However, given our small sample sizes and the low sequencing depth from some samples, our replication power is limited. Thus, some of the other 245 highly differentiated loci in London were also probably impacted by natural selection, although they did not survive our conservative filtering criteria. Increased sample sizes coupled with additional functional data will help to further dissect the evolutionary role played by these variants in the immune response to \textit{Y. pestis}.

**Discussion**

Our results provide strong empirical evidence that the Black Death was an important selective force that shaped genetic diversity around some immune loci. Although our candidate selected loci are generally involved in the response to pathogens, and not just \textit{Y. pestis}, our unique sampling design minimized the degree to which other historical events (such as \textit{tuberculosis} or famine\textsuperscript{41-43}) could have affected the inference of selection. To support our ancient genomic data, we confirmed that the strongest candidates for positive selection are directly involved in the immune response to \textit{Y. pestis} using functional data from in vitro infection experiments.

![Fig. 4 ERAP2 genotype is associated with cytokine response to Y. pestis stimulation. a-d. Effect of genotype upon cytokine levels for granulocyte colony-stimulating factor (G-CSF) (a), interleukin-1\( \beta \) (IL-1\( \beta \)) (b), interleukin 10 (IL-10) (c) and C-C motif chemokine ligand 3 (CCL3) (d). Remaining cytokines showed no significant effects and are included in Supplementary Table 7. e. Boxplots showing the percentage of bacteria killed (y axis) by macrophages infected for 24 h as a function of ERAP2 genotype (x axis). The percentage of bacteria killed was calculated as the CFU\textsubscript{24h} – CFU\textsubscript{2h}/CFU\textsubscript{2h}, where CFUs is colony-forming unit. The \(P\) value results from a linear model examining the association between ERAP2 single nucleotide polymorphism genotypes (SNP) (coded as the number of protective rs249794 alleles found in each individual: 0, 1 or 2) and the percentage of bacteria killed.](https://example.com/image.png)
of the canonical full-length ERAP2 protein. We suggest that this protein increases the presentation of Yersinia-derived antigens to CD8+ T cells, stimulating a protective immune response against Y. pestis. Furthermore, we show that macrophages from individuals possessing the selected ERAP2 allele engage in a unique cytokine response to Y. pestis infection and are better able to limit Y. pestis replication in vitro.

In general, individuals with more copies of the selectively advantageous haplotype displayed a weaker cytokine response to infection but a better ability to limit bacterial growth. For example, levels of IL-1β, a key proinflammatory cytokine often associated with pyroptotic cell death, were threefold lower in individuals homozygous for the advantageous ERAP2 genotype when compared to individuals homozygous for the putatively deleterious one. Therefore, subjects with the advantageous haplotype are both more efficient at controlling internalized bacteria and at resisting Y. pestis-induced cell death than subjects with the deleterious haplotype—abilities that may help to reduce bystander tissue damage during infection. However, as our experiments were done in vitro we were unable to directly evaluate the impact of ERAP2 genotype on tissue damage, immune cell recruitment and survival.

More broadly, our results highlight the contribution of natural selection to present-day susceptibility towards chronic inflammatory and autoimmune disease. We show that ERAP2 is transcriptionally responsive to stimulation with a large array of pathogens, supporting its key role in the regulation of immune responses. Therefore, selection imposed by Y. pestis on ERAP2 probably affects the immune response to other pathogens or disease traits. Consistent with this hypothesis, the selectively advantageous ERAP2 variant is also a known risk factor for Crohn’s disease, and ERAP2 variation has also been associated with other infectious diseases. Thus, selection for pathogen defence in the presence of pathogens such as Y. pestis may be counterbalanced against the costs of immune disorders, resulting in a long-term signature of balancing selection. Likewise, another of our top candidate loci (rs11571319 near CTLA4) is associated with an increased risk of rheumatoid arthritis and systemic lupus erythematosus, such that retaining the putatively advantageous allele during the Black Death confers increased risk for autoimmune disease in present-day populations. To date, most of the evidence for an association between autoimmunity risk alleles and adaptation to past infectious diseases remains indirect, primarily because the aetiological agents driving selection remain uncertain. Our ancient genomic and functional analyses suggest that Y. pestis has been one such agent, representing empirical evidence connecting the selective force of past pandemics to present-day susceptibility to disease.

### 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/s41586-022-05349-x.

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

Data availability
Hybridization capture data from the ancient individuals have been deposited in the NCBI Sequence Read Archive (SRA) under BioProject PRJNA798381. Expression data have been deposited into the NCBI Gene Expression Omnibus (GEO) under project GSE194118 (for macrophages) and the NCBI SRA under project accession PRJNA871128 (for PBMCs). Cytokine data is available in Supplementary Table 8 and CFU data in Supplementary Table 9.

Code availability
Scripts for all data analyses are available at github.com/TaurVil/Vil-galysklunk_yersinia_pestis/.

Acknowledgements
We thank all members of the Barreiro laboratory and the Poinar laboratory for their constructive comments and feedback. We thank J. Tung for her comments and edits to the manuscript. Computational resources were provided by the University of Chicago Research Computing Center. Sequencing was performed at the Farncombe Sequencing Facility McMaster University. We thank the Cytometry and Biomarkers platform at the Institut Pasteur for support in conducting this study, with a special thanks to C. Petitdemange for help running the Luminex assay. We thank X. Zhang for assistance in simulating allele frequency changes under neutral evolution. This work was supported by grant R01-GM134376 to L.B.B., H.P. and J.P.-C., a grant from the Wenner-Gren Foundation to J.F.B. (8702), and the UChicago DDRCC, Center for Interdisciplinary Study of Inflammatory Intestinal Disorders (C-IID) (NIDDK P30 DK042086). The SSHRC Insight Development Grant supported the collection of the Danish samples (430-2017-01193). H.N.P. was supported by an Insight Grant no. 20008499 from the Social Sciences and Humanities Research Council of Canada (SSHRC) and The Canadian Institute for Advanced Research under the Humans and the Microbiome programme. T.P.V. was supported by NIH F32GM140568. X.C. and M. Steinrücken were supported by grant R01GM146051. We also thank the University of Chicago Genomics Facility (RRID:SCR_019196), especially P. Faber, for their assistance with RNA sequencing. H.P. thanks D. Poinar for continued support and manuscript suggestions and editing.

Author contributions
L.B.B. and H.N.P. directed the study. J.K. designed the enrichment assays and generated ancient genomic data. T.P.V. led all data and computational analyses, with contributions from J.-C.G. X.C. performed all analyses to estimate selection coefficients under the supervision of M. Steinrücken. J.F.B., J.P. and T.P.V. performed challenge experiments with macrophages and heat-killed Y. pestis. M. Shiratori and A. Dumaine performed the infection experiments on PBMCs and generated the single-cell RNA-sequencing data, with assistance from D.E. and D.M. C.E.D. and J.P.-C. performed and designed the infection experiments with live Y. pestis on macrophages and generated both cytokine and CFU data, with assistance from J.M. and R.B. C.I.Y. and M.B. designed the probes to quantify the isoform encoding the short version of ERAP2 and M.I.P. performed the experiments. R.R., S.N.D., J.A.G. and J.L.B. provided access to samples, archaeological information, including dating, and other relevant information. A.C. and N.V. provided insights into historical context. K.E. and G.B.G. provided additional sampling and bioinformatic processing and cluster maintenance. A. Devault and J.-M.R. provided insight on targeted enrichment and modified versions of baits used for immune enrichment. G.A.R. provided genomic input on loci and contributed financially to the sequencing of targets. T.P.V., J.K., H.N.P. and L.B.B. wrote the manuscript, with input from all authors.

Competing interests
J.K., A. Devault and J.-M.R. declare financial interest in Daicel Arbor Biosciences, which provided the myBaits hybridization capture kits for this work. All other authors declare no competing interests.

Additional information
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41586-022-05349-x.
Correspondence and requests for materials should be addressed to Hendrik N. Poinar or Luis B. Barreiro.
Peer review information Nature thanks M. Thomas P. Gilbert and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.
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Extended Data Fig. 1 | Differences in recombination rate and background selection are insufficient to explain the marked enrichment of high $F_{st}$ values among immune loci. Comparison of recombination rate (A) and background selection levels (B) between neutral loci and our candidate regions. Candidate regions were stratified into those which were tested and those which were candidates for positive selection based on high differentiation in London pre- vs post-BD. (C) Forward simulations matched for the rates of recombination and background selection of the regions targeted in our study show a slight enrichment of highly differentiated sites in candidate regions, but far from the level of enrichment observed in our collected data (D), replicated from Fig. 2a for comparison. For example, whereas our data differentiation at immune loci exceeded the 99th percentile of neutral variants at 2.4x the rate expected by chance (among variants with a MAF > 10%), the same enrichment is less than 1.2x in the simulated data.
Extended Data Fig. 2 | Estimates of the selection coefficients for the four SNPs of interest and power of the inference procedure. (A) Distributions of \( \hat{s}_{\text{MLE}} \) for the four SNPs when replicates are simulated with the corresponding bootstrapped allele frequency distributions as initial conditions and bootstrap-corrected estimates \( \hat{s}_{\text{MLE}} \). Whiskers on the violin plots label the 2.5-, 50-, and 97.5-percentiles of their respective distributions. (B) ROC and (C) Precision-Recall curves for the estimation procedure to distinguish replicates under selection from those under neutrality.
Extended Data Fig. 3 | Principal components of gene expression for macrophages stimulated with heat-killed *Y. pestis*. The first principal component clearly separates stimulated samples from matched controls.
Extended Data Fig. 4 | Response to *Y. pestis* is similar between macrophages stimulated with heat-killed and live *Y. pestis*. Effect size of *Y. pestis* stimulation compared between heat-killed *Y. pestis* (x-axis, n = 33 individuals) and live *Y. pestis* (y-axis, n = 8 individuals). (A) shows all genes, with a blue line representing the best fit line ($r = 0.88$). (B) compares effect sizes at genes near candidates for positive selection profiled in both expression datasets (red: heat-killed; purple: live bacteria). Error bars represent the standard error in estimating the effect size. The direction of effect is consistent except for *LNPEP* (which is not significant in either analysis) and *ERAP1, CTLA4* and *TICAM2* are not shown because they were not expressed at sufficiently high levels in the macrophages from the 8 individuals infected with live *Y. pestis*. Asterisks placed near the point estimate of each value represent the significance: *** $p < 0.001$; ** $p < 0.01$; * $p < 0.05$. 
Extended Data Fig. 5 | Transcriptional changes of genes nearby candidate loci in response to bacterial and viral stimuli. Data are derived from Nedelec et al.27, and Quach et al.28 Nedelec et al. measured the gene expression response of monocyte-derived macrophages to infection with two live intracellular bacteria: *Listeria monocytogenes* (a Gram-positive bacterium) and *Salmonella typhimurium* (a Gram-negative bacterium). Quach et al. characterized the transcriptional response at 6 h of primary monocytes to bacterial and viral stimuli ligands activating Toll-like receptor pathways (TLR1/2, TLR4, and TLR7/8) and live influenza virus. The data for *Y. pestis* are the fold change responses observed in response to heat-killed bacteria. A negative estimate in plot (purple) indicates that the gene is downregulated and a positive value (red) indicates that the gene is up-regulated. The statistical support for the reported changes is given by the associated *p* values. Larger circle sizes represent smaller *p* values and empty circles refer to cases where the gene was not expressed in that dataset.
Extended Data Fig. 6 | Genotype effects on transcription at candidate loci.

Effect of genotype at nearby loci on the expression of ERAP2 (top) and TICAM2 (bottom), across experimental conditions in this study and previously published\(^{26,27}\). For ERAP2, the protective “C” haplotype increases expression in all conditions ($p < 0.001$). For TICAM2, the protective reference haplotype decreases expression only in the unstimulated condition ($p = 2.5 \times 10^{-6}$; $p > 0.05$ in all other conditions).
Extended Data Fig. 7 | UMAP projection of single cell data. UMAP projection of single-cell RNA sequencing data of non-infected cells and cells infected with live Y. pestis for five hours, after integrating samples. Major immune cell types cluster separately and cells are colored by the cell type to which they were assigned.
Extended Data Fig. 8 | Transcriptional changes of genes nearby candidate loci in response to Y. pestis infection across cell types. For each cell type profiled using single-cell RNA sequencing, we show the effect of Y. pestis infection upon gene expression. A negative estimate (purple) indicates that the gene is downregulated and a positive value (red) indicates that the gene is up-regulated. The statistical support for the reported changes is given by the associated p values. Larger circle sizes represent smaller p values and empty circles refer to cases where the gene was not expressed in that cell type.
Extended Data Fig. 9 | Expression of ERAP2 isoforms. Bioanalyzer traces showing the results of the PCR amplification of cDNA across the exon 10 splice junction from macrophages of 10 individuals with different genotypes for the splice variant rs2248374. The genotype of each individual is shown on top. A negative PCR control was also performed using water. The G allele at rs2248374 is predicted to produce an elongated exon 10 containing two premature stop codons (red rectangles), leading to nonsense mediated decay.
Extended Data Fig. 10 | Schematics for estimating selection coefficients.
The time axis serves as an approximate reference of the relative sampling times for the empirical samples. Dashed vertical lines indicate the relative sampling time for each group of samples considered in the analysis, and the floating boxes with orange and blue dots represent pools of samples from a bi-allelic locus. Above and below the time axis are sketches that respectively correspond to the simulation scheme and the likelihood computations. The shaded red horizontal tree represents the population-continuity along approximate time (x-axis), with the Black Death pandemic occurring in the dark shaded period. The shortened branch with a skull at the end represents people who died of the disease. In each simulated replicate, $\Delta p$ and $\Delta p$ mark the respective changes of allele frequency during the pandemic in the mid-pandemic and post-pandemic sample pools. In the inference schematics, each horizontal straight line represents a sampling scheme from which a likelihood was computed. Lightning bolts labeled with $s$ or $-s$ represent the selection coefficients.
Reporting Summary

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For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
- A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
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*Only common tests should be described solely by name; describe more complex techniques in the Methods section.*
- A description of all covariates tested
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- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted

*Give P values as exact values whenever suitable.*
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

*Our web collection on statistics for biologists contains articles on many of the points above.*

Software and code

Policy information about availability of computer code

Data collection: no software was used in data collection

Data analysis: leeHom (v1.2.5); network-aware-bwa (v0.5.10-evan.10); biohazard (v1.0.2); samtools (v1.9); mapDamage (v2.0); GATK (v4.1.4.1); LCLAE (2016); TrimGalore (v0.2.7); STAR (v2.4.1d); SLUM (v3.7.1); RSEM (v1.2.21); cellranger (v3.0.2); soupconcell (v2.0); singularity (v3.4.0); R (v4.1) with limma (v3.48.3), textTinyR (v1.1.3), ggplot2 (3.3.6), and seurat (v4.0.4) packages. Novel code for the analyses in this project can be found at https://github.com/steinru/diplo_locus/.

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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 description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

Hybridization capture data from the ancient individuals have been deposited in the NCBI Sequence Read Archive (SRA) under BioProject PRJNA798381. Expression data have been deposited to GEO under accession number GSE194118 (for macrophages) and the NCBI SRA under accession number PRJNA871128 (for PBMCs). Cytokine and CFU data are available in Table S8 and S9, respectively.
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
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For a reference copy of the document with all sections, see 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

Sample sizes for ancient genomics were based on the number of individuals who could be sampled and for whom remains were enriched for sufficient endogenous DNA to sequence. Power to detect selection was confirmed via simulation. Sample sizes for expression data were selected based upon previous analyses in Dr. Barreiro’s lab demonstrating sufficient power to detect infection effects and some genetic effects in similarly designed studies.

Data exclusions

Ancient genomic data were excluded when they failed to meet DNA quality or coverage standards. No expression data were excluded. One sample was excluded from the CFU data analyses because the cell cultures were noticeably contaminated.

Replication

All attempts at replication were successful, including strong concordance in expression results between heat-killed and live yersinia pestis. Analysis of ancient DNA cannot be replicated due to sample availability; however sampling at two separate locations serves as pseudo-replication and increases confidence in the strongest signatures of selection.

Randomization

Ancient genomes represent a natural experiment and were therefore not capable of being randomized. For expression analyses, cells from each individual were split into two batches, one of which was exposed to yersinia pestis and the other mock-infection control.

Blinding

not applicable as all outcome measures were objectively measured.

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 | Methods |
|---------------------------------|---------|
| n/a                             | n/a     |
| □  |   | □  |   | □ |  | □ |   | □ |  |
| Involved in the study           | Involved in the study |
| Antibodies                      | □ ChiP-seq |
| Eukaryotic cell lines           | □ Flow cytometry |
| Palaeontology and archaeology   | □ MRI-based neuroimaging |
| Animals and other organisms     |         |
| Human research participants     |         |
| Clinical data                   |         |
| Dual use research of concern    |         |

Antibodies

Antibodies used: Mouse Anti-Human CD14 (BD Biosciences)

Validation: The purity of the isolated monocytes was verified using an antibody against CD14 (BD Biosciences) and only samples showing > 90% purity were used to differentiate into macrophages.

Palaeontology and Archaeology

Specimen provenance: The specifics of each sample location and date are provided in Table S1 of the current manuscript. Briefly, samples come from the UK, London specifically from three burial sites, East Smithfield and St. Mary Graces, St. Nicholas Shambles, St. Mary Spital. For those stemming from Denmark and a wider geographic spread, they come from Nordby, Ribe, Viborg, Haagerup and Horsens.

Specimen deposition: Samples from London have been provided to us by Museum of London Center for Human Bioarchaeology. Samples from Denmark...
stem from the Horsens Museum, the Viborg Museum, the Moesgaard Museum, Museum Lolland-Falster, Øhavmuseet, Faaborg, and ADBOU.

**Dating methods**
The London cemeteries are dated based on a combination of primary sources, Bayesian radiocarbon dating, and archaeological remains (primarily coins). We did the same for our Danish samples, as best we could from associated dates, which are less precise due to the lack of records associated with many of the cemeteries and the imprecision of the dating method based on arm position at burial used to date medieval burials in this region. We therefore group Danish samples into either an Early (~850-1350) or Late (1350-1800) period corresponding to before or after the Black Death period.

☐ Tick this box to confirm that the raw and calibrated dates are available in the paper or in Supplementary Information.

**Ethics oversight**
No ethical approval was required given that none of the samples could be linked to living descendants. Usage of the samples was approved by the Museum of London’s Collection Committee.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

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**Human research participants**

**Policy information about** [studies involving human research participants](#)

**Population characteristics**
Human derived cell for infection experiments were obtained from healthy male donors aged 18 to 55 years old of European and African American descent. Each donor’s blood was tested for Hepatitis B, Hepatitis C, Human Immunodeficiency Virus (HIV), and West Nile Virus, and only samples negative for all of the tested pathogens were used and we controlled for genetic ancestry (previously inferred from whole genome sequencing) in all analyses.

**Recruitment**
Depending on the experiments, samples were collected by the Indiana Blood Center (Indianapolis, IN, USA) or the “Établissement Français du Sang” in Paris. Individuals consented their blood to be used for research purposes.

**Ethics oversight**
This study has been approved by the Institutional Review Board at the University of Chicago [protocol #: IRB19-0432].

Note that full information on the approval of the study protocol must also be provided in the manuscript.