

A draft genome of *Yersinia pestis* from victims of the Black Death

Kirsten I. Bos^{1*}, Verena J. Schuenemann^{2*}, G. Brian Golding³, Hernán A. Burbano⁴, Nicholas Waglechner⁵, Brian K. Coombes⁵, Joseph B. McPhee⁵, Sharon N. DeWitte^{6,7}, Matthias Meyer⁴, Sarah Schmedes⁸, James Wood⁹, David J. D. Earn^{5,10}, D. Ann Herring¹¹, Peter Bauer¹², Hendrik N. Poinar^{1,3,5} & Johannes Krause^{2,12}

Technological advances in DNA recovery and sequencing have drastically expanded the scope of genetic analyses of ancient specimens to the extent that full genomic investigations are now feasible and are quickly becoming standard¹. This trend has important implications for infectious disease research because genomic data from ancient microbes may help to elucidate mechanisms of pathogen evolution and adaptation for emerging and re-emerging infections. Here we report a reconstructed ancient genome of *Yersinia pestis* at 30-fold average coverage from Black Death victims securely dated to episodes of pestilence-associated mortality in London, England, 1348–1350. Genetic architecture and phylogenetic analysis indicate that the ancient organism is ancestral to most extant strains and sits very close to the ancestral node of all *Y. pestis* commonly associated with human infection. Temporal estimates suggest that the Black Death of 1347–1351 was the main historical event responsible for the introduction and widespread dissemination of the ancestor to all currently circulating *Y. pestis* strains pathogenic to humans, and further indicates that contemporary *Y. pestis* epidemics have their origins in the medieval era. Comparisons against modern genomes reveal no unique derived positions in the medieval organism, indicating that the perceived increased virulence of the disease during the Black Death may not have been due to bacterial phenotype. These findings support the notion that factors other than microbial genetics, such as environment, vector dynamics and host susceptibility, should be at the forefront of epidemiological discussions regarding emerging *Y. pestis* infections.

The Black Death of 1347–1351, caused by the bacterium *Yersinia pestis*^{2,3}, provides one of the best historical examples of an emerging infection with rapid dissemination and high mortality, claiming an estimated 30–50% of the European population in only a five-year period⁴. Discrepancies in epidemiological trends between the medieval disease and modern *Y. pestis* infections have ignited controversy over the pandemic's aetiological agent^{5,6}. Although ancient DNA investigations have strongly implicated *Y. pestis*^{2,3} in the ancient pandemic, genetic changes in the bacterium may be partially responsible for differences in disease manifestation and severity. To understand the organism's evolution it is necessary to characterize the genetic changes involved in its transformation from a sylvatic pathogen to one capable of pandemic human infection on the scale of the Black Death, and to determine its relationship with currently circulating strains. Here we begin this discussion by presenting the first draft genome sequence of the ancient pathogen.

Y. pestis is a recently evolved descendent of the soil-dwelling bacillus *Yersinia pseudotuberculosis*⁷, which in the course of its evolution

acquired two additional plasmids (pMT1 and pPCP1) that provide it with specialized mechanisms for infiltrating mammalian hosts. To investigate potential evolutionary changes in one of these plasmids, we reported on the screening of 46 teeth and 53 bones from the East Smithfield collection of London, England for presence of the *Y. pestis*-specific pPCP1 (ref. 3). Historical data indicate that the East Smithfield burial ground was established in late 1348 or early 1349 specifically for interment of Black Death victims⁸ (Supplementary Figs 1 and 2), making the collection well-suited for genetic investigations of ancient *Y. pestis*. DNA sequence data for five teeth obtained via molecular capture of the full *Y. pestis*-specific pPCP1 revealed a C to T damage pattern characteristic of authentic endogenous ancient DNA⁹, and assembly of the pooled Illumina reads permitted the reconstruction of 98.68% of the 9.6-kilobase plasmid at a minimum of twofold coverage³.

To evaluate the suitability of capture-based methods for reconstructing the complete ancient genome, multiple DNA extracts from both roots and crowns stemming from four of the five teeth which yielded the highest pPCP1 coverage³ were used for array-based enrichment (Agilent) and subsequent high-throughput sequencing on the Illumina GAII platform¹⁰. Removal of duplicate molecules and subsequent filtering produced a total of 2,366,647 high quality chromosomal reads (Supplementary Table 1a, b) with an average fragment length of 55.53 base pairs (Supplementary Fig. 4), which is typical for ancient DNA. Coverage estimates yielded an average of 28.2 reads per site for the chromosome, and 35.2 and 31.2 for the pCD1 and pMT1 plasmids, respectively (Fig. 1a, c, d and Supplementary Table 1b, c). Coverage was predictably low for pPCP1 (Fig. 1e) because probes specific to this plasmid were not included on the arrays. Coverage correlated with GC content (Supplementary Fig. 6), a trend previously observed for high-throughput sequence data¹¹. The coverage on each half of the chromosome was uneven due to differences in sequencing depth between the two arrays, with 36.46 and 22.41 average reads per site for array 1 and array 2, respectively. Although greater depth contributed to more average reads per site, it did not increase overall coverage, with both arrays covering 93.48% of the targeted regions at a minimum of onefold coverage (Supplementary Table 1b). This indicates that our capture procedure successfully retrieved template molecules from all genomic regions accessible via this method, and that deeper sequencing would not result in additional data for CO92 template regions not covered in our data set.

Genome architecture is known to vary widely among extant *Y. pestis* strains¹². To extrapolate gene order in our ancient genome, we analysed reads mapping to the CO92 reference for all extracts stemming

¹McMaster Ancient DNA Centre, Department of Anthropology, McMaster University, 1280 Main Street West, Hamilton, Ontario L8S 4L8, Canada. ²Institute for Archaeological Sciences, Rümelinstr. 23, University of Tübingen, 72070 Tübingen, Germany. ³Biology Department, McMaster University, 1280 Main Street West, Hamilton, Ontario L8S 4L8, Canada. ⁴Department of Evolutionary Genetics, Max Planck Institute for Evolutionary Anthropology, 04103 Leipzig, Germany. ⁵Michael G. DeGroot Institute for Infectious Disease Research, McMaster University, 1280 Main Street West, Hamilton, Ontario L8S 4L8, Canada. ⁶Department of Anthropology, University of South Carolina, Columbia, South Carolina 29208, USA. ⁷Department of Biological Sciences, University of South Carolina, Columbia, South Carolina 29208, USA. ⁸Institute of Applied Genetics, University of North Texas Health Science Center, 3500 Camp Bowie Boulevard, Fort Worth, Texas 76107, USA. ⁹Department of Anthropology and Population Research Institute, Pennsylvania State University, University Park, Pennsylvania 16802, USA. ¹⁰Department of Mathematics and Statistics, 1280 Main Street West, Hamilton, Ontario L8S 4K1, Canada. ¹¹Department of Anthropology, McMaster University, 1280 Main Street West, Hamilton, Ontario L8S 4L8, Canada. ¹²Human Genetics Department, Medical Faculty, University of Tübingen, 72070 Tübingen, Germany.

*These authors contributed equally to this work.

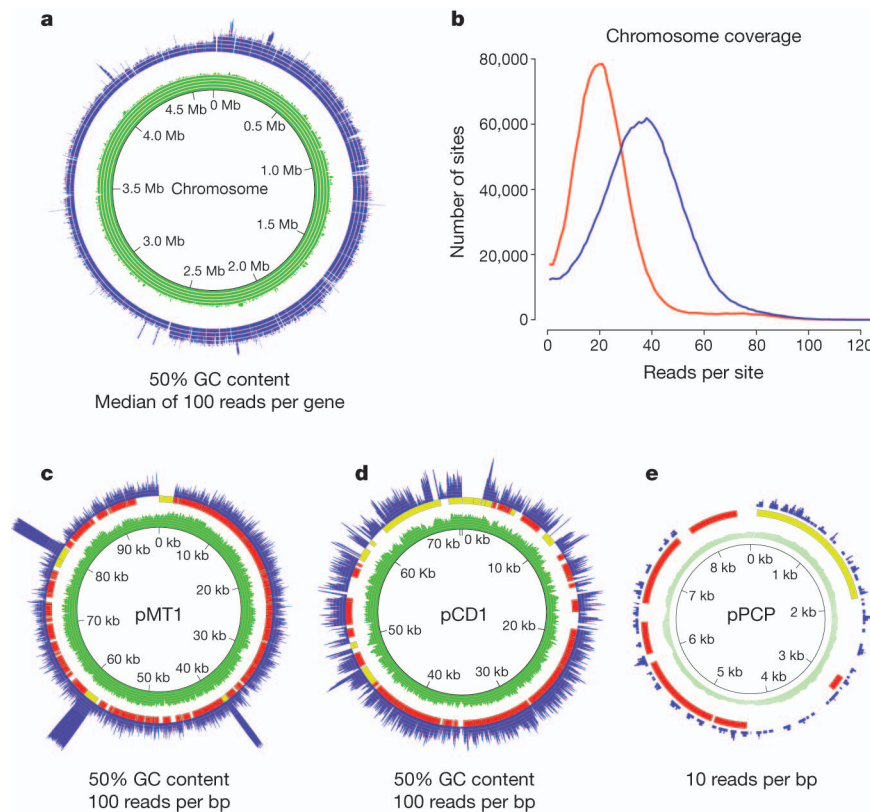


Figure 1 | Coverage plots for genomic regions sequenced. a, c–e, Coverage plots for the chromosome (a) and the plasmids pMT1 (c), pCD1 (d) and pPCP (e). Coverage in blue, GC content in green. Scale lines indicate 10-, 20-, 30-, 40- and 50-fold coverage and 10%, 20%, 30%, 40% and 50% GC content. For plasmids, red corresponds to coding regions, yellow to mobile elements.

Chromosome shows median coverage per gene. Plasmids show each site plotted. Coverage distributions for the plasmids are shown in Supplementary Fig. 5. b, Distributions show chromosomal coverage of array 1 (blue) and array 2 (red), indicating that deeper sequencing increases the number of reads per site, but does not substantially influence overall coverage.

from a single individual who yielded the highest coverage (individual 8291). Despite the short read length of our ancient sequences and the highly repetitive nature of the *Y. pestis* genome, 2,221 contigs matching CO92 were extracted, comprising a total of 4,367,867 bp. To identify potential regions of the ancient genome that are architecturally distinct from CO92, all reads not mapping to the CO92 reference were in turn considered for contig construction. After filtering for a minimum length of 500 bp, 2,134 contigs remained comprising 4,013,009 bp, of which 30,959 stemmed from unmapped reads. Conventional BLAST search queried against the CO92 genome identified matches for 2,105 contigs. Evidence of altered architecture was identified in 10 contigs (Supplementary Table 2). An example of such a structural variant is shown in Fig. 2, where reference-guided assembly incorporating unmapped reads to span the breakpoint validates its reconstruction. This specific genetic orientation is found only in *Y. pseudotuberculosis* and *Y. pestis* strains Mictrotus 91001, Angola, Pestoides F and B42003004, which are ancestral to all *Y. pestis* commonly associated with human infections (branch 1 and branch 2 strains^{13,14}). Furthermore, discrepancies in the arrangement of this region in branch 1 and branch 2 modern *Y. pestis* strains indicate that rearrangements occurred as separate events on different lineages.

Single-nucleotide differences between our ancient genome and the CO92 reference surprisingly consisted of only 97 chromosomal positions, and 2 and 4 positions in the pCD1 and pMT1 plasmids, respectively (Supplementary Table 3), indicating tight genetic conservation in this organism over the last 660 years. Twenty-seven of these positions were unreported in a previous analysis of extant *Y. pestis* diversity¹⁴ (Supplementary Tables 3 and 4). Comparison of our ancient genome to its ancestor *Y. pseudotuberculosis* revealed that the medieval sequence contained the ancestral nucleotide for all 97 positions, indicating that

it does not possess any derived positions absent in other *Y. pestis* strains. Two previously reported chromosomal differences³ were not present in our genomic sequence data, suggesting that they probably

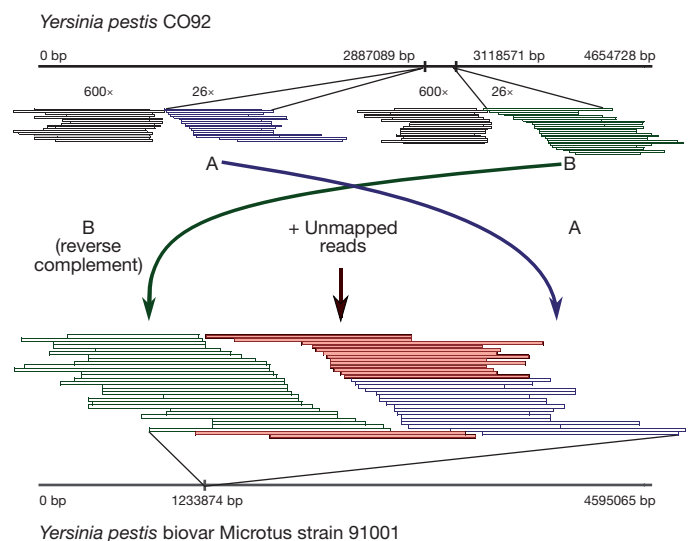


Figure 2 | Alignment of mapped reconstructed contigs against CO92 and Mictrotus genomes. Reads mapped at positions A (blue) and B (green) are 231 kb apart in the linearized CO92 genome. Adjacent sequence is high coverage although only 18 \times and 20 \times is shown due to space constraints (black) for A and B, respectively. The structural variant was assembled using reads that did not map to CO92 (red). Its position is shown on the linearized Mictrotus 91001 chromosome where the 9,096 bp contig maps with 100% identity.

derived from deaminated cytosines that would have been removed in the current investigation via uracil-DNA-glycosylase treatment before array capture.

To place our ancient genome in a phylogenetic context, we characterized all 1,694 previously identified phylogenetically informative positions¹⁴ (Supplementary Table 4), and compared those from our ancient organism against aggregate base call data for 17 publicly available *Y. pestis* genomes and the ancestral *Y. pseudotuberculosis*. When considered separately, sequences from three of the four victims fall only two substitutions from the root of all extant human pathogenic *Y. pestis* strains (Fig. 3a), and they show a closer relationship to branch 1 *Y. pestis* than to branch 2; however, one of the four victims (individual 6330) was infected with a strain that contained three additional derived positions seen in all other branch 1 genomes¹⁴. This suggests either the presence of

multiple strains in the London 1348–1350 pandemic or microevolutionary changes accruing in one strain, which is known to occur in disease outbreaks¹⁵. Additional support for *Y. pestis* microevolution is indicated by the presence of several variant positions for which sequence data from one individual shows two different nucleotides at comparable frequencies (Supplementary Table 5). Position 2896636, for example, is a known polymorphic position in extant *Y. pestis* populations¹⁴, and this position shows the fixed derived state in one individual (6330) and the polymorphic state in another (individual 8291) at minimum fivefold coverage (Supplementary Fig. 7). This provides a remarkable example of microevolution captured during an historical pandemic. The remaining variance positions are unchanged in the 18 extant *Yersinia* genomes, thus they may be unique to the ancient organism and are, therefore, of further interest. Additional sampling

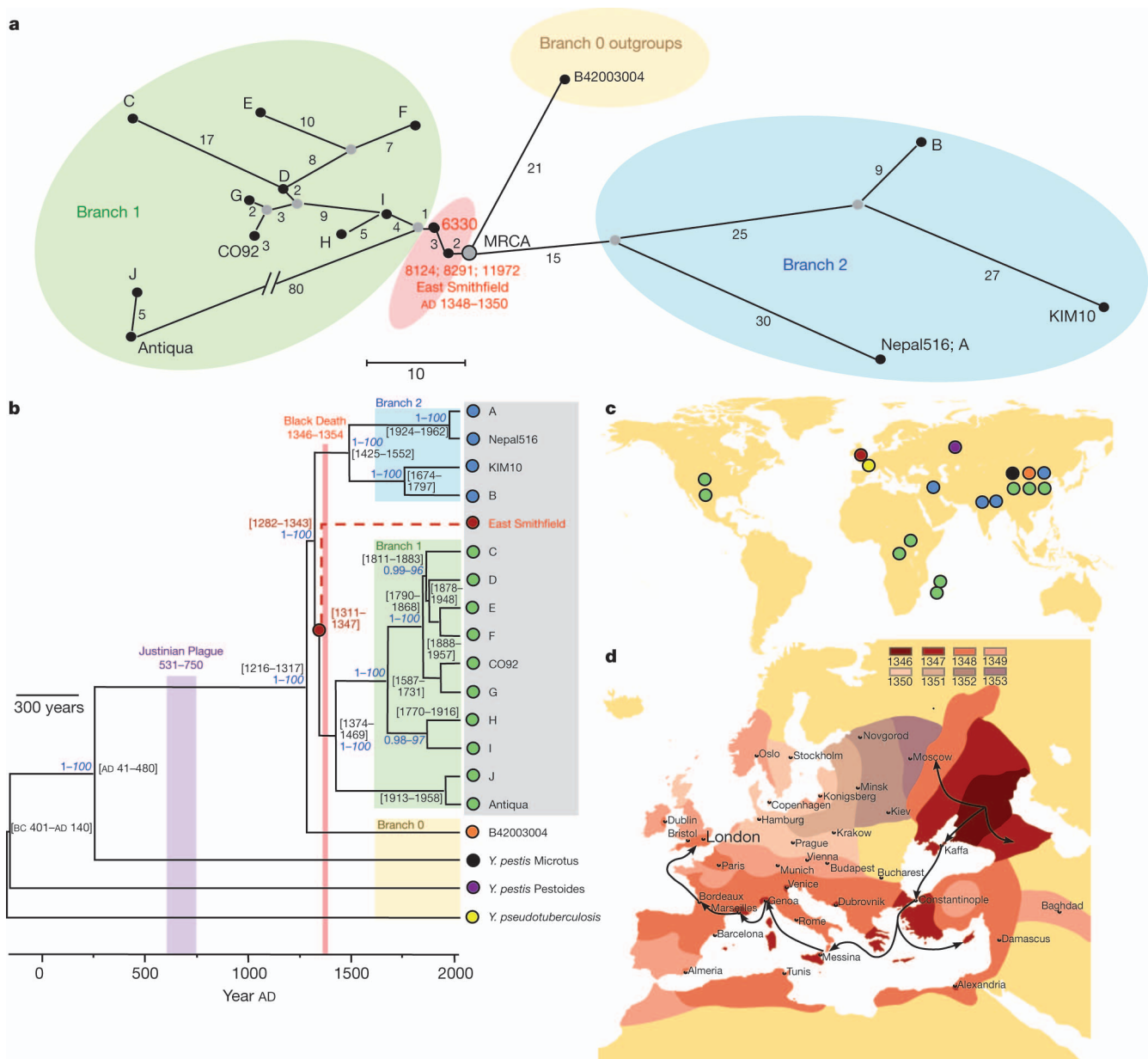


Figure 3 | Phylogenetic placement and historical context for the East Smithfield strain. **a**, Median network of ancient and modern *Y. pestis* based on 1,694 variant positions in modern genomes¹⁴. Coloured circles represent different clades as defined in ref. 13. Gray circles represent hypothetical nodes. **b**, Phylogenetic tree using 1,694 variable positions. Divergence time intervals are shown in calendar years, with neighbour-joining bootstrap support (blue italic) and Bayesian posterior probability (blue). Grey box indicates known

human pathogenic strains. A, NZ ACNQ01000; Nepal516, NC 008149; KIM10, NC 004088; B, NZ AAYT01000; C, NZ ABAT01000; D, NZ ACNS01000; E, NZ AAYS01000; F, NZ AAOS02000; CO92, NC 003143; G, NZ ABCD01000; H, NZ AAYV01000; I, NC 014029; J, NZ AAYR01000; Antiqua, NC 008150. **c**, Geographical origin of genome sequences used in **a** and **b**. **d**, Geographical spread of the Black Death from infection routes reported in ref. 4.

of ancient genomes will assist in determining the frequency of these mutations in co-circulating *Y. pestis* strains, and will clarify the emergence of branch 2 strains that are as yet unreported in ancient samples.

Consistent tree topologies were produced using several construction methods and all major nodes were supported by posterior probability (pp) values of >0.96 and bootstrap values >90 (Fig. 3b and Supplementary Figs 8 and 9). The trees place the East Smithfield sequence close to the ancestral node of all extant human pathogenic *Y. pestis* strains (only two differences in 1,694 positions) and at the base of branch 1 (Fig. 3b). A secure date for the East Smithfield site of 1348–1350 allowed us to assign a tip calibration to the ancient sequence and thus date the divergence time of the modern genomes and the East Smithfield genome using a Bayesian approach. Temporal estimates indicate that all *Y. pestis* commonly associated with human infection shared a common ancestor sometime between 668 and 729 years ago (AD 1282–1343, 95% highest probability density, HPD), encompassing a much smaller time interval than recently published estimates¹⁴ and further indicating that all currently circulating branch 1 and branch 2 isolates emerged during the thirteenth century at the earliest (Fig. 3b), potentially stemming from an Eastern Asian source as has been previously suggested¹⁴. This implies that the medieval plague was the main historical event that introduced human populations to the ancestor of all known pathogenic strains of *Y. pestis*. This further questions the aetiology of the sixth to eighth century Plague of Justinian, popularly assumed to have resulted from the same pathogen: our temporal estimates imply that the pandemic was either caused by a *Y. pestis* variant that is distinct from all currently circulating strains commonly associated with human infections, or it was another disease altogether.

Although our approach of using an extant *Y. pestis* reference template for bait design precluded our ability to identify genomic regions that may have been present in the ancient organism and were subsequently lost in CO92, genomic comparisons of our ancient sequence against its closest outgroups may yield valuable insights into *Y. pestis* evolution. The *Microtus* 91001 strain is the closest branch 1 and branch 2 relative confirmed to be non-pathogenic to humans¹⁶, hence genetic changes may represent contributions to the pathogen's adaptation to a human host. Comparisons against this outgroup revealed 113 changes (Supplementary Table 6a, b), many of which are found in genes affecting virulence-associated functions like biofilm formation (*hmsT*), iron-acquisition (*iucD*) or adaptation to the intracellular environment (*phoP*). Similarly, although its virulence potential in humans has yet to be confirmed to our knowledge, *Y. pestis* B42003004 isolated from a Chinese marmot population¹⁷ has been identified as the strain closest to the ancestral node of all *Y. pestis* commonly associated with human plague, and thus may provide key information regarding the organism's evolution. Full genome comparison against the East Smithfield sequence revealed only eight single-nucleotide differences (Supplementary Table 6c), six of which result in non-synonymous changes (Supplementary Table 6d). Although these differences probably do not affect virulence, the influence of gene loss, gene gain or genetic rearrangements, all of which are well documented in *Y. pestis*^{12,18}, is as yet undetermined. In more recent evolutionary terms, single-nucleotide differences in several known pathogenicity-associated genes were found between our ancient genome and the CO92 reference sequence (Supplementary Table 3), which may represent further adaptations to human hosts.

Through enrichment by DNA capture coupled with targeted high throughput DNA sequencing, we have reconstructed a draft genome for what is arguably the most devastating human pathogen in history, and revealed that the medieval plague of the fourteenth century was probably responsible for its introduction and widespread distribution in human populations. This indicates that the pathogen implicated in the Black Death has close relatives in the twenty-first century that are both endemic and emerging¹⁹. Introductions of new pathogens to populations are often associated with increased incidence and severity

of disease²⁰ and although the mechanisms governing this phenomenon are complex²¹, genetic data from ancient infectious diseases will provide invaluable contributions towards our understanding of host–pathogen coevolution. The Black Death is a seminal example of an emerging infection, travelling across Europe and claiming the lives of an estimated 30 million people in only 5 years, which is much faster than contemporary rates of bubonic or pneumonic plague infection²² and dissemination^{7,8}. Regardless, although no extant *Y. pestis* strain possesses the same genetic profile as our ancient organism, our data suggest that few changes in known virulence-associated genes have accrued in the organism's 660 years of evolution as a human pathogen, further suggesting that its perceived increased virulence in history²³ may not be due to novel fixed point mutations detectable via the analytical approach described here. At our current resolution, we posit that molecular changes in pathogens are but one component of a constellation of factors contributing to changing infectious disease prevalence and severity, where genetics of the host population²⁴, climate²⁵, vector dynamics²⁶, social conditions²⁷ and synergistic interactions with concurrent diseases²⁸ should be foremost in discussions of population susceptibility to infectious disease and host–pathogen relationships with reference to *Y. pestis* infections.

METHODS SUMMARY

DNA from dental pulp was extracted and converted into sequencing libraries as previously described³. Potential sequencing artefacts resulting from deaminated nucleotides were eliminated by treatment of the DNA extracts with uracil-DNA-glycosylase and endonuclease VIII. DNA extracts were subsequently converted into sequencing libraries and amplified to incorporate unique sequence tags on both ends of the molecule. Two Agilent DNA capture arrays were designed for capture of the full *Y. pestis* chromosome (4.6 megabases), and the pCD1 (70 kb) and pMT1 (100 kb) plasmids using the modern *Y. pestis* strain CO92 (accession numbers NC_003143, NC_003131, NC_003134) for bait design with 3 bp tiling density. Serial array capture was performed over two copies of each array using the enriched fraction from the first round of capture as a template for a second round. The resulting products were amplified and pooled in equimolar amounts. All templates were sequenced for 76 cycles from both ends on the Illumina GAII platform, and reads merged into single fragments were included in subsequent analyses only if forward and reverse sequences overlapped by a minimum of 11 bp. Reads were mapped against the CO92 genome using the software BWA, and molecules with the same start and end coordinates were removed with the *rmDup* program in the *samtools* suite. Reference-guided sequence assembly was performed using Velvet version 1.1.03, with mapped and unmapped reads supplied in separate channels. Single-nucleotide differences were determined at a minimum of fivefold coverage and base frequency of at least 95% for both a pooled data set for all individuals and one in which all individuals were treated separately. A median network was constructed on these base calls using SplitsTree4. Phylogenetic trees were constructed using parsimony, neighbour-joining (MEGA 4.1) and Bayesian methods, and coalescence dates were determined in BEAST using both a strict and a relaxed molecular clock (Supplementary Fig. 9).

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

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CORRIGENDUM

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& Johannes Krause

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In this Letter, the GenBank accession number was wrongly printed as SRA045745.1. It should be SRA045745. Supplementary Tables 3a and 4 have also been corrected.