RESEARCH ARTICLE

Recovery of a Medieval Brucella melitensis Genome Using Shotgun Metagenomics

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ABSTRACT Shotgun metagenomics provides a powerful assumption-free approach to the recovery of pathogen genomes from contemporary and historical material. We sequenced the metagenome of a calcified nodule from the skeleton of a 14th-century middle-aged male excavated from the medieval Sardinian settlement of Geridu. We obtained 6.5-fold coverage of a Brucella melitensis genome. Sequence reads from this genome showed signatures typical of ancient or aged DNA. Despite the relatively low coverage, we were able to use information from single-nucleotide polymorphisms to place the medieval pathogen genome within a clade of B. melitensis strains that included the well-studied Ether strain and two other recent Italian isolates. We confirmed this placement using information from deletions and IS711 insertions. We conclude that metagenomics stands ready to document past and present infections, shedding light on the emergence, evolution, and spread of microbial pathogens.

IMPORTANCE Infectious diseases have shaped human populations and societies throughout history. The recovery of pathogen DNA sequences from human remains provides an opportunity to identify and characterize the causes of individual and epidemic infections. By sequencing DNA extracted from medieval human remains through shotgun metagenomics, without target-specific capture or amplification, we have obtained a draft genome sequence of an ~700-year-old Brucella melitensis strain. Using a variety of bioinformatic approaches, we have shown that this historical strain is most closely related to recent strains isolated from Italy, confirming the continuity of this zoonotic infection, and even a specific lineage, in the Mediterranean region over the centuries.

Brucellosis is a widespread infection of livestock (sheep, goats, cattle, cows, and pigs) and remains one of the most common zoonotic infections, with more than 500,000 new human cases worldwide annually (1). Human brucellosis is most commonly caused by the species Brucella melitensis and is usually acquired through ingestion of unpasteurized dairy products or, less commonly, through ingestion of infected meat or direct occupational contact with animals (1). If left untreated, the infection usually follows a chronic course, spreading systemically to the organs of the reticuloendothelial system and often leading to osteoarticular disease (2).

Brucellosis is an ancient disease. Vertebral lesions consistent with brucellosis have been described in a 2-million-year-old male skeleton of Australopithecus africanus (3). Lesions consistent with brucellosis have been described in Bronze Age skeletons from the Levant and the Basque country, in adult skeletons from Herculaneum, and in medieval human remains (2).

A major ambition of paleopathology is to shed light on the influence of infectious disease on past populations. However, morphological analyses are limited in that few infections produce durable lesions and very different pathogens can produce similar pathologies (4). For example, similar lesions occur in tuberculosis and brucellosis, even though the causative organisms are quite different taxonomically and in cell structure. Amplification of pathogen DNA from human remains via PCR has provided an alternative source of information about a range of past infections (5). However, only a single study has reported success in using PCR amplification to confirm historical brucellosis (6).

In addition, there are problems with amplification-based approaches to the recovery of historical and ancient DNA. First, PCR is a competitive and highly sensitive reaction, prone to contamination even in dedicated facilities (7). Second, these approaches generally provide information on a single gene or gene fragment, affording little or no insight into pathogen biology, evolution, and epidemiology. Third, they require the onerous design and optimization of pathogen-specific primers; this limits the open-
endedness of the approach, so that one generally finds only what one is looking for. This last point also applies to genome capture approaches, which have proven successful in recovering *Yersinia pestis* genomes from samples from the Black Death and the Justinianic plague (8, 9).

Shotgun metagenomics—that is the unbiased sequencing *en masse* of DNA extracted from a sample without target-specific amplification or capture—provides an attractive alternative approach to the detection and characterization of pathogens in contemporary and historical human material. This approach has proven successful in obtaining genome-wide sequence data for *Borrelia burgdorferi*, *Mycobacterium tuberculosis*, and *Mycobacterium leprae* from long-dead human remains (10–12).

When confronted with calcified nodules from a 14th-century skeleton, we initially thought of tuberculosis. However, when we used shotgun metagenomics to identify potential pathogens in the sample, we were instead surprised to recover a medieval *Brucella melitensis* genome sequence.

### RESULTS

** Metagenomic recovery of *Brucella melitensis* sequences with signatures of medieval origin.

The skeleton of a 50- to 60-year-old male (skeleton 2568) was excavated from the abandoned medieval village of Geridu (Sorso, Sassari, Italy) in northwest Sardinia in December 1997 (Fig. 1) (13). The skeleton showed features of diffuse idiopathic skeletal hyperostosis (DISH), including fusions between the fourth and tenth thoracic vertebrae, fusion of the fifth lumbar vertebra to the sacrum, and extraspinal enthesopathies (14). Thirty-two calcified nodules were found in the pelvic girdle, with diameters ranging from 0.6 by 0.7 cm to 2.2 by 1.6 cm (Fig. 1). A DNA extraction was performed on one of the nodules.

We obtained a DNA yield from the nodule of 30 ng, which was used to construct a TruSeq Nano Illumina library, which was run at low coverage on an Illumina MiSeq sequencer, alongside 10 other bar-coded libraries, 8 from other historical human tissue samples and 2 from blank controls. Just over two million se-
quences were obtained from sample 2568 on this run. An analysis of size distribution was performed, revealing a bimodal distribution with a broad peak running from 50 to 150 bp, with a second much taller peak running up to the maximum read length of 250 bp (Fig. 1). On the assumption that historical DNA fragments were restricted to the smaller peak, sequences over 150 bp in length were excluded from analysis of this run. Homology searches revealed sequences from sample 2568 that could be assigned with confidence to the genus *Brucella*.

We then attempted to map reads from all 11 samples against the genome of the *B. melitensis* reference strain 16 M. We obtained insignificant matches for 10 of the samples (<12 aligned reads per sample), whereas sample 2568 yielded >20,000 paired-end reads (equivalent to 10,000 sequences, as the fragments are shorter than the read length used in paired-end sequences) that mapped against the *B. melitensis* 16 M genome, providing approximately 0.7-fold coverage of a medieval *Brucella* genome from a strain that we have called Geridu-1. A coverage plot (Fig. 1) revealed even coverage across both chromosomes in the 16 M genome, ruling out spurious hits to conserved sequences from environmental bacteria.

To obtain additional sequencing reads, the sample 2568 library was sequenced at ~10-fold-higher coverage on a single dedicated MiSeq run, which yielded just over 20 million paired-end sequences. When these were mapped at high stringency, 23% of these reads mapped against the human genome and 0.48% against the *B. melitensis* 16 M genome (representing 6.5-fold coverage). Interestingly, when the reads that mapped to either the *B. melitensis* or human genomes were reanalyzed, they showed a much tighter size distribution than the library as a whole, with a peak centered on 100 bases (Fig. 1). In addition, the reads mapping to the *B. melitensis* or human genome showed abundant CT and GA base conversions at the 5’ and 3’ ends, which is indicative of the damage typical of ancient or aged DNA (Fig. 1). These findings are supportive of a medieval origin of these sequences (15).

**SNP-based phylogenetic placement shows that the medieval *Brucella* genome is closely related to recent Italian isolates.** Conventional phylogenetic methods based on identification of trusted single-nucleotide polymorphisms (SNPs) cannot be applied to low-coverage genome sequences. However, the technique of “phylogenetic placement” provides an alternative solution (16).

Here, one draws on a fixed reference tree, computed from high-coverage genomes, and places the unknown query sequence on the tree using programs such as pplacer. We used a published set of phylogenetically informative SNPs for *Brucella* spp. (17) and analyzed reads from the initial MiSeq run that aligned to equivalent positions in the 16 M genome. Using this approach, despite the low coverage, we could show confidently (with a posterior probability of 1) that the Geridu-1 strain clustered most closely with the well-characterized Ether strain (ATCC 23458; the reference strain for *B. melitensis* biovar 3) and was nested within a clade of four *B. melitensis* strains (see Fig. S1 in the supplemental material). Indeed, with as few as 250 reads, we were able to accurately assign the Geridu-1 strain to the Ether clade (Fig. S2).

To refine the placement of the Geridu-1 genome, we constructed a broad-based phylogenetic tree from all available modern *B. melitensis* genomes. This showed that Geridu-1’s close relative, the Ether strain, belonged in a distinctive clade along with four other strains, for which only draft genome sequences were available (see Fig. S3 in the supplemental material). We then compared all five strains from the Ether clade and the Geridu-1 genome recovered from the second MiSeq run by calling SNPs against the completed 16 M genome (Table S1) and using them to draw a phylogenetic tree for this clade (Fig. 2). The SNP table and tree showed that the Geridu-1 strain represented the earliest branching lineage within the Ether clade, separated by 450 to 500 SNPs from any other strain in the clade (Table S2).

At least three of the five contemporary strains that belong to the Ether clade originate from the Italian peninsula or from Sicily: F15/06-7 is a 2006 human isolate from Sicily, F5/07-239A is a 2007 small-ruminant isolate from Italy, and the Ether strain itself was isolated from an Italian goat in 1961. There are no other Italian isolates in the set of currently available *B. melitensis* genome sequences, perhaps suggesting that the Ether clade originated in Italy and its associated islands, or at least in the western Mediterranean. We note that in a recent multiple-locus variable-number tandem-repeat analysis (MLVA) study (18), Italian strains clustered separately from most other European strains and we suspect that Geridu-1 belongs to the western Mediterranean cluster defined by MLVA, although additional genome sequencing of isolates from that cluster are required to confirm this.

**Confirmation of phylogenetic placement using insertions and deletions.** To confirm the placement of the Geridu-1 strain within the Ether clade of *B. melitensis*, we drew upon two additional sources of information: the distribution of deletions and the locations of insertion elements. First, we looked for deletions of >100 bases in length that occurred in the Geridu-1 genome in comparison to the 16 M strain. We identified 11 such deletions (see Table S3 in the supplemental material). We determined the distribution of these deletions in all available *B. melitensis* genomes. Nine deletions were found in only Geridu-1 and the five other strains in the Ether clade. Two occurred sporadically in other strains.

Next, we examined the distribution and location of the insertion element IS711 (also called IS6501), which occurs widely in *Brucella* spp. (19). When we mapped reads from the Geridu-1 genome to the insertion element, we obtained 52.3-fold coverage. Dividing that by the average coverage for the Geridu-1 genome (6.5-fold) provided us with an estimate of eight IS711 copies in the medieval strain. By analyzing reads that spanned the ends of IS711 and the adjacent chromosome, we were able to confirm the existence of eight insertion points in Geridu-1, seven of which also occurred in *B. melitensis* 16 M and in all other available *B. melitensis* genomes.

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**Fig 2** Phylogenetic tree showing the position of the medieval Geridu-1 strain within the Ether clade. Only those SNPs which correlated with sufficient coverage in the Geridu-1 alignment were included in the construction of the tree.
sis genomes (Table S3). The IS711 insertion present in Geridu-1 but absent from 16 M was located at position 517784 in chromosome 2, disrupting a gene encoding the hypothetical protein BMEII0494. This insertion point was found in no other B. melitennis strain genome, apart from the other five members of the Ether lineage. These patterns of deletion and IS711 insertions confirm the placement of the Geridu-1 strain within the Ether clade of B. melitensis.

**DISCUSSION**

Here, we have shown that shotgun metagenomics can be used to obtain a Brucella melitensis genome sequence from a medieval sample without target-specific amplification or capture. The recently reported success of this approach with two mycobacterial diseases, leprosy and tuberculosis, has been ascribed to the unique properties of the mycobacterial cell wall in preserving bacterial DNA (11, 12). In contrast, this study confirms that whole-genome sequences from bacterial pathogens without resilient cell envelopes can be recovered from human remains by metagenomics hundreds or even thousands of years postmortem. However, unlike medieval mycobacterial DNA (12), medieval Brucella DNA does show signatures of damage associated with an ancient or historical provenance.

This observation complements several other lines of evidence that support the authenticity of data obtained from this historical sample. First, the sample was processed in facilities dedicated to ancient DNA research, where no work on Brucella cultures or DNA had ever taken place. Second, laboratory contamination was ruled out by the lack of significant hits for *Brucella* in libraries obtained from eight other historical specimens and two blanks sequenced in the same run. Third, we observed conclusive and extensive matches to a dedicated human and animal pathogen, obtaining even genomic coverage of a genome nested within the *B. melitensis* phylogeny. This rules out spurious and patchy hits for conserved genes from related environmental organisms as a source of error. Finally, we took care to sample the interior of the nodule, which eliminated the risk of contamination from soil.

Calcification of soft tissue is a recognized, albeit rare, complication of human brucellosis, so we assume that calcification of abdominal or pelvic tissues accounts for the appearance of pelvic nodules in this individual. The skeletal pathology provides no additional evidence of brucellosis. Instead, we propose that the demineralized nodule was formed after death from a nonhuman source. The calcified nodule was designated as possibly biologic or nonbiologic, with more appropriate terminology pending further study.

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**MATERIALS AND METHODS**

**Source material.** The source material consisted of 1 of 32 calcified nodules found in the pelvic girdle of an adult male skeleton (sample 2568) that was excavated from 1997 to 1999 from a cemetery in the medieval rural settlement of Geridu (Sorso, Sassari, Italy), in northwest Sardinia. Twenty-five single pit graves were identified in a well-organized part of the cemetery (sector 2500). Stratigraphic analysis identified two distinct burial phases. Phase I (9 single burials) dates to the first half of the 14th century CE; phase II (16 single burials) dates to the final period in which the cemetery was used (1350 to 1400 CE). Following the historical reports, the settlement of Geridu was definitely abandoned in 1426 CE. The sample 2568 remains were retrieved from 1 of the 16 burials dating to the second half of the 14th century CE (13). Sex determination was performed on the basis of the morphological features of the skull. Age at death was estimated on the basis of dental wear and sternal rib end modification (22). Lesions indicative of pathologies were recorded in accordance with the methods and standards set out in the Global History of Health Project (23).

**DNA extraction.** DNA extraction and library preparation were carried out in a dedicated ancient DNA laboratory in which no strains of *Brucella* had ever been cultured, no pathogen-specific PCR amplification had ever been performed, and in which the handler wore gloves, together with a mask, a gown, and a hood. The surface of the calcified nodule was removed using a drill bit that had been cleaned with bleach to eliminate potential surface contaminants. The undersurface was then removed by drilling at low speed to produce approximately 20 mg of powder. The sample was incubated at 37°C with shaking to demineralize it in 400 μl CTAB (cetyltrimethylammonium bromide) solution and 40 μl proteinase K for 1 week.

DNA was isolated with chloroform using the DNeasy plant minikit (Qiagen United Kingdom), with the following modifications to the manufacturer’s protocol. Three volumes of buffer AW1 was added to each sample without target-specific amplification or capture. The reaction was incubated at 37°C with shaking to demineralize it in 400 μl CTAB (cetyltrimethylammonium bromide) solution and 40 μl proteinase K for 1 week. DNA was eluted in 50 μl of 0.2 M chloroform and the tube was inverted for 3 min, and the supernatant was transferred to 2-ml tubes. The manufacturer’s DNeasy plant minikit protocol was followed from step 6 with the following additional modifications. Three volumes of buffer AW1 was added to each sample and incubated at room temperature for 2 to 3 h. After the addition of buffer AW2, the samples were centrifuged for 3 min. DNA was eluted in two 50-μl aliquots (100-μl total). Extracted DNA was quantified in 5 μl of sample using the Qubit high-sensitivity double-stranded DNA (HS dsDNA) assay according to the manufacturer’s instructions (Invitrogen Ltd., Paisley, United Kingdom) and then stored at −20°C until library preparation.

**Sequencing.** DNA extracted from sample 2568 was converted into a TruSeq Nano library for sequencing on an Illumina MiSeq sequencer according to the manufacturer’s low-sample protocol (Illumina UK, Little Chicago, IL). In brief, the demineralized sample was centrifuged at 20,000 × g for 10 min and the supernatant was collected. To homogenize the sample, 400 μl of 0.2 M chloroform was added and the tube was inverted for 10 min. The sample was centrifuged at 6,000 × g for 3 min, and the supernatant was transferred to 2-ml tubes. The manufacturer’s DNeasy plant minikit protocol was followed from step 6 with the following additional modifications. Three volumes of buffer AW1 was added to each sample and incubated at room temperature for 2 to 3 h. After the addition of buffer AW2, the samples were centrifuged for 3 min. DNA was eluted in two 50-μl aliquots (100-μl total). Extracted DNA was quantified in 5 μl of sample using the Qubit high-sensitivity double-stranded DNA (HS dsDNA) assay according to the manufacturer’s instructions (Invitrogen Ltd., Paisley, United Kingdom) and then stored at −20°C until library preparation.

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Chesterford, United Kingdom), with the following minor modifications. No fragmentation step was included, given the expectation that ancient DNA would already be heavily fragmented. DNA was end-repaired within the ancient-DNA laboratory, with the modification of incubation for 90 min at 30°C and size selection for <350-bp inserts. Analysis on an Agilent Bioanalyzer 2100 system provided an estimated size distribution of fragments with a peak length of 235 bp.

Once adapters (including bar codes) had been ligated, the sequencing library was moved to a PCR laboratory. In accordance with the manufacturer’s instructions, the library fragments were PCR amplified; however, we used 10 PCR cycles instead of the usual 8. The library was quantified (2 μl per sample) using the Qubit HS dsDNA assay according to the manufacturer’s instructions (Invitrogen Ltd., Paisley, United Kingdom) and then stored at −20°C. The sample 2568 library was diluted to 4 nM, as determined by analysis on an Agilent Bioanalyzer 2100 and using the Qubit HS dsDNA assay, and then pooled in equimolar amounts with 10 other bar-coded libraries (8 from other historical human tissue samples and 2 from blank controls). The entire library pool was then diluted to 12 pM and sequenced on the first MiSeq run using the Illumina MiSeq v2 2 × 250-bp paired-end protocol. In a second MiSeq run, diluted entirely to the 2568 library, the 4 nM library was diluted to 12 pM without pooling.

Identification of Brucella sequences. Metagenomic sequence reads from both MiSeq runs using sample 2568 have been deposited in the European Nucleotide Archive (project accession number PRJEB6045). Sequences derived from sample 2568 in the first MiSeq run that had lengths of ≥150 bp were subjected to a BLASTN search against the NCBI NR database, and the results were analyzed with MEGAN (24). Reads from all samples on the initial MiSeq run were analyzed with Bowtie2 version 2.1.0 (25), allowing only 1 mismatch per 33 bases of the read (under the following settings: --mp 1,1; --ignore-quals; --score-min L,0,−0.033). The reads were mapped against the genomes of the following pathogens: Plasmodium falciparum 3D7, Leishmania infantum JPCM5, Yersinia pestis CO92, Mycobacterium tuberculosis H37Rv, and Brucella melitensis 16 M (GenBank accession numbers AL844501 to AL844509, AE011485 to AE011488, AE001362, FR796433 to FR796468, NC_003143, AL123456, NC_003317, and NC_003318).

Phylogenetic placement of Brucella sequences from Geridu at low coverage. Previously described lineage-defining SNPs (17) were used to construct a tree using FastTree 2.7.1. Using this extended matrix, concatenated SNPs for each strain were then called from the mapped bam file using Samtools (29), with requirements that there was at least 6-fold coverage and that the mutant allele accounted for at least 80% of aligned sequences. The SNPs from Geridu-1 were combined with those from the Ether clade strains, excluding any SNP that occurred at a location where there was less than 6-fold coverage in the Geridu-1 sequence alignment. The remaining 2,332 SNPs (Table S1) were then used to construct a tree using FastTree 2.7.1 and to produce a pairwise dissimilarity matrix.

Analysis of breakpoints in Geridu-1 and other strains. From manual scrutiny of the coverage plot, we identified regions in the 16 M reference genome where for a span of >100 bp, no reads mapped from the Geridu-1 genome. To confirm the existence and refine the boundaries of these deletions, the Geridu-1 reads were remapped against 16 M using the --local option of Bowtie2, which allows the beginning and end of reads to be soft-clipped to obtain an improved alignment. The clipped regions of the reads at the edges of candidate deletions were used to identify the breakpoint created by the deletion. The distribution of the breakpoints in other strains was determined by retrieving the sequences flanking the deletion breakpoints in the Geridu-1 strain and performing a BLASTN search of available B. melitensis genomes. Similar approaches were used to determine the identity and distribution of insertion breakpoints associated with IS11.

Nucleotide sequence accession numbers. Metagenomic sequence reads from this study have been deposited in the European Nucleotide Archive (project accession number PRJEB6045). The following URL will pull all of the sequences that were used in our searches: http://www.ncbi.nlm.nih.gov/nuccore/AL844501,AL844502,AL844503,AL844504,AL84505,AL844506,AL844507,AL844508,AL844509,AE011485,AE011486,AE011486,AE011487,AE011488,AE001362,FR796433,FR796434,FR796435,FR796436,FR796437,FR796438,NC_003143,AL123456,NC_003317,NC_003318.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl?doc-id=10.1128/mBio.01337-14/-/DCSupplemental.

Figure S1, PDF file, 0.2 MB.
Figure S2, PDF file, 0.1 MB.
Figure S3, PDF file, 0.5 MB.
Table S1, PDF file, 0.1 MB.
Table S2, PDF file, 0.4 MB.
Table S3, PDF file, 0.1 MB.

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R.B., B.B., and M.J.P. designed the research; M.M., R.B., V.G., P.B., and G.L.K. performed the research; G.L.K., M.J.S., and M.J.P. analyzed the data; and M.J.P. and R.B. wrote the paper.

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