Human tuberculosis predates domestication in ancient Syria

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Human tuberculosis predates domestication in ancient Syria


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Title page

Title: Human tuberculosis predates domestication in ancient Syria

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Summary:

The question of pre-neolithic tuberculosis is still open in paleopathological perspective. One of the major interests is to explore what type of infection could have existed around the early stage of animal domestication. Paleopathological lesions evoking skeletal TB were observed on five human skeletons coming from two PPNB sites in Syria, which belongs to the geographical cradle of agriculture. These sites represent respectively pre-domestication phase (Dja’de el Mughara, Northern Syria, 8800-8300 BCE cal.) and early domestication phase (Tell Aswad, Southern Syria, 8200-7600 BCE cal.). MicroCT scan analyses were performed on two specimens (one per site) and revealed microscopic changes in favour of TB infection. Detection of lipid biomarkers is positive for two specimens (one per site). Initial molecular analysis further indicates the presence of TB in one individual from Dja’de. Interestingly, no morphological evidence of TB was observed on animal remains of wild and newly domesticated species, discovered in these sites. These observations strongly suggest the presence of human tuberculosis before domestication and at its early stages.

Keywords: paleopathology of TB, Early Neolithic, PPNB, agriculture cradle, domestication, lipid biomarkers, ancient DNA

Introduction:

Human tuberculosis is a specific disease caused by infection by a member of *Mycobacterium tuberculosis* complex. Although the previous widely accepted hypothesis that the human pathogen (*M. tuberculosis*) derived from cattle pathogen (*M. bovis*) by contamination during the Neolithic, the new evolutionary scenario assess that *M. tuberculosis* actually represents a direct descendant of a most ancient strain that existed before the split of *M. africanum - M. bovis* lineages from the *M. tuberculosis* lineage \(^1\). This suggests that the tubercle bacilli could have been already a human pathogen before animal domestication.

Paleopathological evidence of tuberculosis before the Neolithic is poorly documented. Lesions observed on extinct bison remains discovered in a natural trap
cave in Wyoming and dated from 17000 BP, led to biomolecular identification of *M. tuberculosis* ancient DNA and lipid biomarkers. A diagnosis of leptomeningitis tuberculosa was mentioned about endocranial lesions observed on a fossil hominin attributed to *Homo erectus* from Turkey, but this hypothesis was questioned by other authors. As for human tuberculosis, most ancient cases are dated from 7000 yrs BCE in the Eastern Mediterranean. In the site of Atlit Yam, remains of an adult female and an immature individual presented paleopathological evidence of TB, confirmed by lipid biomarkers and aDNA analyses. In the same geographic area, paleopathological evidences of tuberculosis were previously mentioned for contemporaneous site (Ain Ghazal, ca. 7250 BCE). These cases correspond to the development of the first farming and herding societies in the Fertile Crescent, known to be the cradle of agriculture.

In order to explore the link between human and cattle tuberculosis at the beginning of domestication process, the first purpose of this study is to investigate the paleopathological evidence of tuberculosis from two ancient Neolithic sites representing pre and early domestication phases in ancient Syria. This study will allow shedding further light on the debate concerning the origins and the evolutionary pathway of the *Mycobacterium tuberculosis* complex, for example by dating the emergence of the modern strain of *M. tuberculosis*.

**Materials and methods:**

**Archaeological material**

Studied material is coming from two Pre-Pottery Neolithic B (PPNB) sites from Syria, Dja’de el Mughara and Tell Aswad.

The pre-domestic Neolithic site of Dja’de el Mughara is located on the left bank of the Euphrates River, about 115 Km N/W from Aleppo. It has been excavated from 1991 to 2010 in the framework of the French Archaeological Research Program in Syria, headed by Eric Coqueugniot. The chronology of the human occupation ranged from 9310 to 8200 BCE cal. It has been divided in 3 chrono-cultural periods. The first one (DJ 1) corresponds to the end of PPNA, transition phase to early PPNB (9310-8830 BCE cal.). Human remains associated to this period are scarce: only five individuals
were discovered, represented by four adult skulls and a fragmentary skeleton of an adolescent. The second period (DJ 2) corresponds to early PPNB and is dated from 8800 to 8500 BCE cal. Human settlement is represented by 13 burials that provided skeletal remains of 26 individuals (8 immature individuals and 18 adults). The third period (DJ 3), corresponding to the end of early PPNB, is dated from 8540 to 8290 BCE cal. Human remains associated to this period are more numerous: 99 individuals were identified (37 adult and 62 immature individuals). More than 70 are coming from collective burials (Houses of the Dead). After 8200 BCE, the site was abandoned till the 7th millennium.

During this period, people still rely on hunting and gathering. Indeed, the wild resources were still dominating the diet, and there is no morphological evidence that either cereals or animals were domesticated. At the "DJ 3" phase, however, some clues of pre-domestication could be recognized, both for plants (wild cereals) and animals (aurochs) 7, 8.

For this site, skeletal remains of 130 individuals were examined by two researchers of this study (OB and BC).

The Neolithic site of Tell Aswad located around 35 Km from East / Southeast Damascus (Southern Syria) has been discovered in 1967 by H. de Contenson, and excavations were performed between 2001 and 2006 by a French-Syrian archaeological team co-headed by Danielle Stordeur and Bassam Jammous 9.

Numerous skeletal remains were discovered, representing 119 individuals, dating from middle and end of PPNB (8200-7600 BCE cal). The domestic resources have been strongly used since the middle PPNB: cereals are derived from an agricultural economy, and exploitation of domestic animals increases in this period, while hunting decreases.

Methods

**Anthropology and paleopathology**

All the skeletal material was carefully examined in the storage place on site, both for anthropological and paleopathological purposes. Possible changes due to TB were systematically researched on spine, bones and joints. Diagnostic was based upon
the morphology and distribution of the lesions according to criteria commonly accepted in clinical and paleopathological literature \textsuperscript{10-13}, including aspects of spondylodiscitis, osteoarthritis and periostitis. Age estimation and sex determination of the immature and adult individuals were performed according to a set of different methods currently used in biological anthropology\textsuperscript{14-16}.

The skeletal pieces matching with these criteria were selected and analysed by different methods. Morphological methods have used digital imaging acquisitions by laser and X-ray (\(\mu\)CT) with volumetric reconstructions. Lipidic biomarkers were detected by HPLC\textsuperscript{17} and MTBC specific sequence motifs were targeted by ancient DNA based analysis.

In the same time, animal bones found in these sites (wild and domesticated species) were studied by two researchers of this study (DH and LG). No paleopathological lesion was observed. A sample of 9 cattle bones from Dja‘de el Mughara was selected for further analyses (lipid biomarkers and ancient DNA).

\textbf{Digital imaging}

Lesions present on 2 thoracic vertebrae of individual Dja‘de 304 were digitalized by 3D laser scan at low energy (class II), without contact at a mean resolution lower than 0.5mm. Contrarily to X-rays, this surface laser acquisition allows saving morphological information before sampling for molecular analysis while protecting ancient DNA from ionization. Moreover, it allows the 3D printing of the lesions by rapid prototyping (Eden250\textsuperscript{TM} 3D Printing System, Objet, with horizontal layers of 16µm) using VIRCOPAL\textsuperscript{®} technology \textsuperscript{18}.

Two specimens were analysed by microtomodensitometry: distal ulna of individual Aswad 509 on a GE Healthcare eXplore Locus microCT at a resolution of 20µm and lumbar vertebra of young immature individual Dja‘de B108 on a GE V/tome/x at 3µm. 3D reconstructions were performed using TIVMI\textsuperscript{®} software program \textsuperscript{19}.

\textbf{Lipid extraction}

Analyses were performed on 2 specimens presenting paleopathological lesions (Dja‘de sp483 and Tell Aswad 509). The Dja‘de skeleton sp483 provided 14 vertebrae (Laboratory Numbers DV1 to DV14) and 36 rib (Laboratory Numbers DR1 to DR36) samples for study. Eleven fibula fragments were available from Tell Aswad skeleton 509 (Laboratory Numbers AF1 to AF11). Bone samples AF1 (548 mg), AF3
(573 mg), AF11 (473 mg), DR13 (256 mg), DR36 (316 mg), DV7 (214 mg) and DV13 (70 mg) were chosen for lipid biomarker analysis. Specimens were hydrolysed by heating with 30% potassium hydroxide in methanol (2 ml) and toluene (1 ml) at 100°C overnight. In parallel, standard biomass from Mycobacterium tuberculosis was processed. Long-chain compounds were extracted as described previously and the extract was treated with pentafluorobenzyl bromide, under phase-transfer conditions, to convert acidic components into pentafluorobenzyl (PFB) esters. Subsequent separation on an Alltech 209250 (500 mg) normal phase silica gel cartridge gave fractions containing non-hydroxylated fatty acid PFB esters, mycolic acid (MA) PFB esters. The MA PFB esters were reacted with pyrenebutyric acid (PBA) to produce PBA-PFB MA derivatives, which were purified on an Alltech 205250 (500 mg) C₁₈ reverse phase cartridge. The PBA-PFB mycolates were analysed by reverse phase HPLC, as described previously. The non-hydroxylated PFB ester fractions were refined on an Alltech 205250 (500 mg) reverse phase silica gel cartridge, using a water-methanol/methanol/methanol-toluene elution sequence. A fraction enriched in mycocerosic acid and other longer chain (> C₂₀) PFB esters was eluted by 100% methanol with the more usual C₁₂ to C₂₀ esters eluting in the earlier water/methanol fractions. The fractions containing possible mycolipenate and mycocerosates, were analysed by negative ion chemical ionization gas chromatography mass spectrometry (NICI-GCMS), essentially as previously described. Technical details of the GC-MS analyses are provided in Supplementary Information.

**Ancient DNA analysis**

Molecular analyses were conducted at the ancient DNA laboratory of the EURAC Institute for Mummies and the Iceman, Bolzano, Italy. Sample preparation and DNA extraction was performed in a dedicated pre-PCR area following the strict procedures required for studies of ancient DNA: use of protective clothing, UV-light exposure of the equipment and bleach sterilization of surfaces, use of PCR workstations and filtered pipette tips. Within a designated sample preparation room the outer surface of the bone samples was mechanically removed by using a Dremel speed rotary tool. The surfaces of hard tissue material were subsequently subjected to an 15min UV treatment. Finally, the cleaned samples were pulverized using a Retsch mixer mill.
Hz, 15 sec). A complete list of the bone samples used for molecular analysis is provided in the Table 1.

DNA extraction was performed with approximately 250 mg of tissue powder using a silica-based DNA extraction described by Rohland et al. with minor modifications. Different primer combinations were used to initially check for the presence of human mitochondrial DNA and to further diagnose TB. Details of the PCR conditions and to the oligonucleotide primers used in this study are provided in Supplementary Information. The obtained PCR products were separated by electrophoresis on a 2.85 % agarose gel and visualized on a UV-screen after staining with ethidium bromide. The nucleotide sequences of the PCR products were determined by direct sequencing. The PCR products of amplification were checked and purified (EXO-SAP treatments) and four microliters of the PCR amplification cleaned-product were sequenced on an ABI Prism 310 DNA automated sequencer, using the BigDyeTM Terminator Cycle Sequencing Ready Reaction Kit version 3.1 (Applied Biosystems, Foster City, CA, USA). Primers used for the sequencing were the same as those used for the PCRs. Results were subjected to sequence analysis via NCBI blastN.

Results

Paleopathology

Dja’dé el Mughara site

Among the skeletal remains of the site representing a total of 135 individuals, 4 individuals showed paleopathological lesions that are consistent with a diagnosis of skeletal tuberculosis.

Two cases are attributed to the early stage of PPNB, "DJ 2" (8800-8500 BCE cal.)

The first specimen (sp483) is an immature individual of about 1 year old at death (determination based on dental maturation stage). The individual was buried in a rectangular structure against a wall, in a primary flexed position. The structure is dated from the "DJ 2b" phase, an initial stage of the early PPNB, which is only known at Dja’dé el Mughara. The skeleton is fragmented, mainly represented by fragmented skull (frontal, occipital, parietal bones); 19 vertebrae (unfused); fragmented claviculae, humerii, ulnae, radii, right femora, tibia and fibula, incomplete metatarsal
and foot phalanges. The lesions are represented by symmetrical periosteal reaction on long bones of upper limbs, characterized by thin unilamellar layer of woven bone ‘appliqué’ on the diaphyseal shafts.

On the anterior part of the vertebral bodies, besides an enlargement of the vascular foramina (anterior venous plexuses) a periosteal reaction is also visible. Similar pattern has been described at the end of the 19th century as a special expression of spine TB infection, which differed from Pott’s disease. The skeleton sp675 belonged to 8-10 years old individual (determination based on dental maturation and epiphyseal fusion stages) who was primarily buried half-lied down, half-seated in a narrow pit. The burial had no direct link with architecture, but according to the stratigraphic level the skeleton can be dated to the "DJ 2" phase. The skeleton is represented by fragmented skull (mainly facial bones), quite well preserved claviculae, humerii, radii, ulnae, femorae, tibiae, fragmented fibulae, incomplete numbers of hand an foot short and tubular bones. Lesions are represented by a periosteal reaction on both ulnae and tibiae, having the same global pattern (symetrical, woven bone, ‘appliqué’) but thinner than for the previous case. A periosteal reaction is clearly visible on the internal side of 3 preserved ribs, that is has been presented as a paleopathological marker of TB. Several vertebral bodies present the same pathological pattern on their anterior part (enlargement of the venous foramina, periosteal reaction) than those described for the young immature sp483.

Two other cases are attributed to "DJ 3" phase (8540-8290 BCE cal.). The first case came from the 304 collective grave, which was not fully excavated, as a large part was situated in the baulk. The minimal number of individuals represented in this burial is of 5 individuals, 3 adults and 2 immature individuals (4-6 years and 10-14 years old at time of death). The individual which exhibits typical lesions of vertebral TB infection is one of the 3 adults. Sex determination was not possible due to the poor preservation of coxal bones. Due to the collective nature of the burial and mixing of skeletal elements, it was not possible to associate precisely the spine of this individual with the rest of the skeleton, as the skeletons of the 3 adults were mingled. The axial skeleton is represented by vertebrae from T2 to L4, sacrum and fragments of the two coxal bones. Therefore, sex determination is not possible, but small marginal osteophytosis present from 4th to 8th vertebrae can help in age estimation, indicating that this spine belonged to an adult individual. Typical lesions
are present on the 9th and 10th thoracic vertebrae (figure 1a). They are mainly lytic. The inferior part of the 9th thoracic vertebrae is completely destroyed, and the upper part of the 10th thoracic vertebrae shows cystic rounded cavitations expanded to the vertebral body, having a space occupying mass aspect (figure 1b). Due to the lack of any changes in the facet joints, it does not seem that vertebral collapse did occur, in spite of the important lytic destruction of the two vertebral bodies. The cavity was probably occupied by a renitent TB granulomatous mass that therefore offered a biomechanical protection against collapse. This aspect matches with criteria of cystic aspect of spinal TB\textsuperscript{27, 28}

Using 3D reconstruction from laser surface acquisition, it has been possible to reassemble precisely the zygapophyseal joints and thus to get a more precise view of the spinal static. Indeed, it seems that the vertebrae did not collapse in spite of the importance of the destruction of the vertebral bodies, which is confirming the existence of a space occupying mass (such as a granulomatous cyst) that has probably protected the thoracic spine from crush. This pattern is not commonly described in modern clinical practice, but can be seen by medical imaging on living patients\textsuperscript{29} and mummified bodies\textsuperscript{28} and has been clearly identified on skeletal material\textsuperscript{27}, related to tuberculosis of spine.

The second case is labelled B108, coming from a collective burial containing 16 individuals. The skeleton, in good preservation state, is almost complete, excepting the lower limbs that are missing. Dental maturation stage indicated the individual died at an age of about 4-5 years. Lesions were tiny, located on 7th to 12th thoracic vertebrae and on the first 3 preserved lumbar vertebrae. The global pattern is characterized by a slight periosteal reaction on the anterior part of the vertebral body that is developed around the anterior venous plexus foraminae. This aspect can matches with an early stage of superficial vertebral tuberculosis\textsuperscript{25}.

Tell Aswad site

Among the hundred individuals excavated in this site, we examined 25 skeletons older than 15 years. The rest of the individuals are represented by young immature individuals and adult plastered skulls.
The skeleton 509 coming from an individual burial dated from the Middle PPNB period (8200-7600 BCE cal.) is well preserved, excepting the axial skeleton (spine, pelvic bones and ribs) that is vanished. Age estimation from the physeal fusion indicated a result of 20-22 years old with a Bayesian probability of 97% \(^1\). Long and tubular bones (humerii, radii, ulnae, metacarpal, femorae, tibiae, fibulae, metatarsals) exhibited a striking pattern of symmetrical diffuse periosteal reaction, made of thin layers of woven bone (figure 2b). The distal part of the right ulna is swollen. The symmetrical periosteal reaction involving long and short tubular bones is characteristic of Hypertrophic Pulmonary Osteoarthropathy (HPOA), syndrome that is due to chronic pulmonary disease, including TB \(^3\). TB is here the much probable aetiology, due to the young age of this individual, rather than a malignant process that mainly concern elderly people.

MicroCT
The first specimen is the vertebra of the young immature Dja'de B108. In contrast with minor morphological changes, the 3D reconstruction evidenced clear focal microarchitectural changes on the antero-lateral inner part of the body, when it is sectioned along a horizontal axis. These lesions can be interpreted as the first step of the process of tubercular infection of spine corresponding to a hematogenous spread via lateral equatorial branches of the vertebral artery (for detailed description and illustration, see Coqueugniot et al, this volume) \(^2\).

The second specimen is the ulna of the individual 509 from Tell Aswad. Slices revealed a paucilamellar periosteal reaction, with an "appliqué" aspect. The cortical bone showed an increasing of the porosities predominant on the endosteal side. The cortical bone showed an increasing of the porosities predominant on the endosteal side. The 3D reconstruction (figure 2b) revealed that the reorganization on the endosteal side (looking like a kind of medullar expansion into the cortical bone) corresponds to a probable myelitis, with spread of the infection to the cortical bone through the vascular canals. Moreover, 3D viewing showed some communications between these dilated channels and the space localized under the periosteal reaction. This aspect is typical of the one of osteomyelitis \(^1\). Given the fact that the periosteal reaction is diffuse on all long bones and tubular bones of this individual, only tuberculosis can be responsible of this diffuse skeletal infection.
Lipid biomarkers

Profiles of total mycolic acids are shown in Figure 3 and mycocerosate/mycolipenate profiles are in Figure 4. The reverse phase HPLC profiles of total mycolates (Figure 3) were relatively weak for the Dja’de specimens, particularly DV13; correspondence with the total mycolates from standard *M. tuberculosis* was imprecise. The Tell Aswad extracts were essentially negative, with some unknown earlier eluting material (10 – 15 minutes). Owing to the low abundance of these mycolates, it was not productive to examine these extracts by sequential normal and reverse phase HPLC, a procedure that can provide additional high quality diagnostic information.

The mycocerosate/mycolipenate profiles in Figure 4, were more informative for both the Dja’de and Aswad specimens. In the case of the Dja’de 483 vertebra (DV13), clear peaks were recorded for C\textsubscript{30} and C\textsubscript{32} mycocerosates and C\textsubscript{27} mycolipenate, but the same signals were weaker for DR13. Another Dja’de 483 rib sample (DR36) also provided clear C\textsubscript{29}, C\textsubscript{30} and C\textsubscript{32} mycocerosates but C\textsubscript{27} mycolipenate was not detected. Mycolipenate and mycocerosates were not found in extracts of DV7. The three Aswad 509 fibula samples (AF1, AF3 and AF11) had strong peaks for C\textsubscript{27} mycolipenate. Aswad 509 AF1 and AF3 had weak C\textsubscript{30} and C\textsubscript{32} mycocerosate signals, but only C\textsubscript{32} mycocerosate was recorded for AF11.

Ancient DNA

Molecular paleomicrobiological analysis was used to diagnose MTBC bacteria in selected specimens. Four of the five individuals displaying paleopathological lesions indicative for TB were subjected to further molecular analysis. If possible, both one long bone section and one pathological bone section (rib and vertebra) was analysed. Initially, the analysis of a short fragment of the human mtDNA hypervariable region 1 indicated a poor overall preservation of DNA in the human skeletal remains. Only in three of the nine analysed bone samples, human mtDNA was detected via PCR (Table 1). The presence of TB DNA was assessed by applying a nested PCR-based assay targeting the MTBC multicopy IS6110 region. Within the nine bone samples analysed in this study two specimens of individual 483 of the Dja’de site showed positive molecular results for MTBC pathogens. An IS6110 123 bp product from the rib (DR36) and a 92 bp nested IS6110 product from the vertebra (DV8) and rib (DR36) sample were obtained and confirmed by sequencing.
Discussion:
Paleopathological results are strongly suggestive of tubercular infection for two individuals: Dja’dé 304 which showed typical Pott's disease and Aswad 509 which presented a diffuse periosteal reaction characteristic of HPOA. MicroCT analyses with 3D reconstruction revealed a typical pattern of osteomyelitis. Moreover, microarchitectural changes of trabecular vertebral bone observed on the young child Dja’dé 108B keenly supports the hypothesis of early stage of tubercular spondylitis. Mycolic acid profiles did not give clear diagnoses for tuberculosis (Figure 3). Some consistent, but poorly shaped, peaks were seen for DR13, DR36 and DV7 specimens from Dja’dé 483, but the DV13 trace was uninformative. It is likely that the mycolates are substantially degraded. The Aswad 509 profiles had negligible evidence for mycobacterial mycolic acids, but AF1, AF3 and AF11 consistently had substantial amounts of early eluting peaks whose rather uniform profiles suggested a non-microbial origin (Figure 3).

In the Aswad 509 individual, the tuberculosis-specific C$_{27}$ mycolipenic acid biomarker was prominent in the three fibula cases AF1, AF3 and AF11 (Figure 2; C$_{27}$ m/z 407). This positive diagnosis was backed up by the presence of small but clearly positive C$_{32}$ (m/z 479) mycocerosates in all cases and additional C$_{30}$ (m/z 451) mycocerosates in extracts of AF1 and AF3 (Figure 4). Taken together, these data confirm tuberculosis in Aswad 509. In contrast, Dja’dé 483 specimens displayed a different balance of mycolipenate and mycocerosic biomarkers with nice clear signals for C$_{30}$ (m/z 451) and C$_{32}$ (m/z 479) mycocerosates, backed up with smaller peaks representing C$_{27}$ mycolipenate (m/z 407) (Figure 2). Again, tuberculosis is confirmed in individual Dja’dé 483 for both rib (DR13, DR36) and vertebra (DV13) samples. The lipid biomarker profiles, recorded here, resemble those recorded for the 17,000 year old bison, where mycolic acids were very degraded but diagnostic mycolipenates and mycocerosates were present $^{20}$.

Ancient DNA first results evidenced the presence of typical IS6110 product for one individual of Dja’dé 483 (young child), confirming the morphological diagnosis of early stage of vertebral tubercular infection, in accordance with the lipid biomarkers. These pluridisciplinary observations strongly suggest the presence of human tuberculosis
before domestication and at its early stages. Further molecular studies on this material will improve our knowledge about the evolution of *Mycobacterium tuberculosis* complex, particularly as they can determine when emerged the modern strain of bacilli responsible for human tuberculosis.

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**References:**


List of figures

Figure 1a: Spinal involvement of individual Dja'de 304 showing antemortem destruction of the body of 9th & 10th thoracic vertebrae.
Figure 1b: Reconstruction of laser surface acquisition of the typical aspect of vertebral tuberculosis observed on the 9th and 10th thoracic vertebrae of individual Dja'de 304.
The cavity was probably occupied by a renitent TB granulomatous mass (pseudo-eosinophilic granuloma) that therefore offered a biomechanical protection against collapse. This aspect matches with criteria of cystic aspect of spinal TB.

Figure 2a: Symmetrical diffuse periosteal reaction observed on long bones of the individual 509 from Tell Aswad (a-humerus; b- right radius & ulna ; c- left radius & ulna ; d- right tibia ; g- left tibia ; e- fragment of left coxal bone ; f- left third metatarsal bone). This paleopathological aspect is in favour of a diagnosis of hypertrophic pulmonary osteoarthropathy.
Figure 2b: 3D reconstruction of microCT (resolution 20 µ) of the ulna of the individual 509 from Tell Aswad.
Various 3D reconstruction A: semi-transparent, longitudinal sections; B: transparent longitudinal section; C: transparent, horizontal sections, are clearly revealing: 1- the reorganization on the endosteal side (aspect of medullar expansion into the cortical bone) 2- dilated channels communicating with the space localized under the periosteal reaction 3- periosteal reaction.
This aspect corresponds to an initial myelitis, with secondary spread of the infection to the cortical bone and subperiosteal space through the cortical vascular canals.

Figure 3: Mycolates profiles

Figure 4: Mycocerosates profiles

Table 1: Summary of the morphological, chemical and molecular analyses of the sampled bone material of Dja'de and Tell Aswad (5 individuals).
<table>
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<th>Lipid biomarkers</th>
<th>Sample type</th>
<th>mtDNA</th>
<th>IS6110</th>
<th>IS6110 intern</th>
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<tr>
<td>Dja’d</td>
<td>DJ III</td>
<td>Early PPNB</td>
<td>B108</td>
<td>4-5 yrs</td>
<td>Superficial anterior vertebral changes</td>
<td>Early stage of spondylitis</td>
<td></td>
<td>Mycolate Mycocerosates Mycolipenate</td>
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<td></td>
<td>DJ III</td>
<td>Early PPNB</td>
<td>304</td>
<td>Adult</td>
<td>Spinal tuberculosis</td>
<td>Cystic vertebral tuberculosis</td>
<td></td>
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<td>Sacrum fragments</td>
<td>+</td>
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<tr>
<td></td>
<td>DJ II</td>
<td>Early PPNB</td>
<td>675</td>
<td>8-10 yrs</td>
<td>Diffuse periosteal reaction</td>
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<td></td>
<td></td>
<td>Tibia, right, fragments</td>
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<td>-</td>
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<tr>
<td></td>
<td>DJ II</td>
<td>Early PPNB</td>
<td>483</td>
<td>1 yr</td>
<td>Superficial anterior vertebral changes</td>
<td></td>
<td></td>
<td></td>
<td>Vertebra (DV13) weak C30(+++), C32(++++), C27(+++)</td>
<td>Vertebra (DV4)</td>
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<td>-</td>
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<td></td>
<td></td>
<td>Vertebra (DV8)</td>
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<td>-</td>
<td>+</td>
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<td>Rib (DR36) weak C29, C30(+), C32(++)</td>
<td>Rib (DR36)</td>
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<td>+</td>
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<tr>
<td>Aswad</td>
<td>Middle</td>
<td>Middle PPNB</td>
<td>509</td>
<td>20-22 yrs</td>
<td>Hypertrophic osteoarthropathy (ulna)</td>
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<td></td>
<td></td>
<td>Fibula (AF1) weak C30, C32, C27(++++)</td>
<td>Fibula (AF1)</td>
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<td>Fibula (AF11) weak C32</td>
<td>Fibula (AF11)</td>
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</table>
Reverse phase HPLC of total mycolates

- DR13
- DR36
- DV7
- DV13
- AF1
- AF3
- AF11
- M. tuberculosis
- Blank

Minutes
**M. tuberculosis**

- C<sub>27</sub> m/z 407: 10.04 (70)
- C<sub>27</sub> m/z 409: 9.66 (3)
- C<sub>29</sub> m/z 437: 11.92 (33)
- C<sub>30</sub> m/z 451: 12.00 (20)
- C<sub>32</sub> m/z 479: 15.39 (100)

**M. tuberculosis (AF3 & DR13)**

- C<sub>27</sub> m/z 407: 13.76 (99)
- C<sub>27</sub> m/z 409: 13.32 (5)
- C<sub>29</sub> m/z 437: 15.72 (40)
- C<sub>30</sub> m/z 451: 15.79 (24)
- C<sub>32</sub> m/z 479: 19.28 (100)
SUPPORTING INFORMATION

Mycocerosic and mycolipenic acid analysis

A Thermo Scientific DSQII Mass Spectrometer coupled to a Thermo Scientific TRACE GC Ultra gas chromatograph, was used at Swansea University. The column was a Phenomenex Zebron ZB-5 (5% phenyl, 95% dimethylpolysiloxane; 30 m × 0.25 mm i.d. × 0.25 µm film thickness), using He as carrier gas (constant flow mode 1.2 ml min\(^{-1}\)) and ammonia as the CI reagent gas. A GC oven temperature gradient from 200 to 300 °C at 17.5 °C min\(^{-1}\) was used, the final temperature being held for 17.5 min. The ion source temperature was 170 °C, the injector used was a programmable temperature vapourising injector, which started at 50 °C for 0.2 min and increased to 300 °C at a rate of 10 °C s\(^{-1}\) where it stayed for 0.5 min.

PFB esters, on NICI-GCMS, fragment to produce negative carboxylate [M – H]\(^{-}\) ions, which can be detected at high sensitivity. Selected ion monitoring (SIM) was used to search for mycocerosate carboxylate ions at \(m/z\) 367.6311 (C\(_{24}\)), 395.6844 (C\(_{26}\)), 409.7111 (C\(_{29}\)), 437.7645 (C\(_{29}\)), 451.7911 (C\(_{30}\)), 479.8445 (C\(_{32}\)), 493.8712 (C\(_{33}\)) and 507.8978 (C\(_{34}\)) \[1\]. Additionally, \(m/z\) 407.6952 was monitored for the presence of the C\(_{27}\) mycolipenate carboxylate ion \[1\]. Partial racemisation of mycocerosates during the alkaline hydrolysis leads to the formation of diasteroisomers, which resolve on gas chromatography to give characteristic doublets; in contrast, mycolipenates are singlets as they cannot racemise \[1\]. Standard GC-MS traces, for \(M.\) tuberculosis, are shown below in Figure S1.

![Figure S1](image-url)

Figure S1. Selected ion monitoring (SIM) negative ion-chemical ionization gas chromatography-mass spectrometry (NI-CI GC-MS) profiles of mycolipenate and mycocerosate pentafluorobenzyl (PFB) esters from extracts of \(M.\) tuberculosis, recorded on different occasions. A. Profiles recorded to correlate with extracts of AF1, AF11 and DV13. B. Profiles recorded to correlate with extracts of AF2 and DR13. Circumstances did not allow recording of a standard to accompany DR36.
Ancient DNA analysis

To assess DNA extraction efficiency and to test for possible PCR inhibitors each DNA extract was subject to a PCR-based assay using a primer pair targeting the mtDNA hypervariable region 1 [2]. The presence of TB DNA was assessed by applying a PCR-based assay targeting the MTBC multicopy IS6110 region [3]. To increase the sensitivity of the assay, a nested PCR strategy was applied [4]. Initially, conventional PCR was performed using primer IS6110F and the IS6110R primer to generate a 123 base pair product. Two microliters of the initial PCR reaction was used as the template for the nested PCR.

Table S1: List of DNA Primers used in this study and the corresponding PCR conditions

<table>
<thead>
<tr>
<th>Molecular target</th>
<th>Primer</th>
<th>Sequence (5’-3’)</th>
<th>Product (bp)</th>
<th>PCR conditions</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>human mtDNA HVS-I</td>
<td>L16117</td>
<td>TACATTACTGCCAGCCACCACCAT</td>
<td>162</td>
<td>95°C, 5 min; 95°C, 55°C, and 72°C, 45 sec for 38 cycles</td>
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<tr>
<td></td>
<td>H16233</td>
<td>GCTTTGGAGTTGCAGTGTGTTGT</td>
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<tr>
<td>MTBC, IS6110</td>
<td>IS6110F</td>
<td>CTCGTCACCGCCGCTCGG</td>
<td>123</td>
<td>95°C, 5 min; 95°C, 68°C, and 72°C, 45 sec for 45 cycles; 72°C, 4 min</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>IS6110R</td>
<td>CCTGCGAGCGTAGGGCGTGG</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>MTBC, IS6110 internal</td>
<td>IS6110intF</td>
<td>TTCGGACCACCAGCACCTAA</td>
<td>92</td>
<td>95°C, 5 min; 95°C, 58°C, and 72°C, 45 sec for 40 cycles; 72°C, 4 min</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>IS6110intR</td>
<td>TCGGTGACAAAAGGCCACGTA</td>
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<td></td>
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</tbody>
</table>

For the IS6110 PCR assay resulting in a 123 bp large fragment [3] the AmpliTaq Gold Mastermix 360 (Applied Biosystems, Foster City, CA, USA) was used according to the manufactures instructions and 2µl of GC Enhancer and 5µl of DNA were added to a final volume of 25µl. The PCR reaction mix for the mitochondrial PCR and the IS6110 internal nested PCR contained 10 mM tris-HCl (pH 8.3), 50 mM KCl, 1.875 mM MgCl2, 200 µM of each deoxynucleotide trisphosphate, 0.5 µM of each primer, 0.1 mg/ml Bovine serum albumin, 0.05 U/µl AmpliTaq Gold (Applied Biosystems, Foster City, CA, USA) and 5 µl of extracted DNA for the mitochondrial PCR or 2µl of PCR product of the IS6110 external PCR for the internal nested PCR to a final volume of 50 µl. Polymerase chain reaction was carried out according to the parameters in Table S1.
References


