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Ancient DNA analysis – an established technique in charting the evolution of tuberculosis and leprosy

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34 Summary

35 Many tuberculosis and leprosy infections are latent or paucibacillary, suggesting a long time-scale for 36 host and pathogen co-existence. Palaeopathology enables recognition of archaeological cases and PCR 37 detects pathogen ancient DNA (aDNA). Mycobacterium tuberculosis and Mycobacterium leprae cell 38 wall lipids are more stable than aDNA and restrict permeability, thereby possibly aiding long-term 39 persistence of pathogen aDNA. Amplification of aDNA, using specific PCR primers designed for short fragments and linked to fluorescent probes, gives good results, especially when designed to target 40 41 multi-copy loci. Such studies have confirmed tuberculosis and leprosy, including co-infections. Many 42 tuberculosis cases have non-specific or no visible skeletal pathology, consistent with the natural history 43 of this disease. M. tuberculosis and M. leprae are obligate parasites, closely associated with their 44 human host following recent clonal distribution. Therefore genotyping based on single nucleotide 45 polymorphisms (SNPs) can indicate their origins, spread and phylogeny. Knowledge of extant genetic 46 lineages at particular times in past human populations can be obtained from well-preserved specimens 47 where molecular typing is possible, using deletion analysis, microsatellite analysis and whole genome sequencing. Such studies have identified non-bovine tuberculosis from a Pleistocene bison from 17,500 48 49 years BP, human tuberculosis from 9000 years ago and leprosy from over 2000 years ago. 50 51 Key words: Ancient DNA; evolution; Mycobacterium leprae; Mycobacterium tuberculosis; molecular

52 typing

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55 **1. Introduction**

According to the World Health Organisation,¹ one third of the global human population is infected 56 57 with tuberculosis (TB) but most are latent infections. In people with no underlying risk factors, approximately 10% will develop an active infection during their lifetime.¹ However, underlying 58 deficiencies in immunity caused by co-infections such as HIV, or co-morbidities such as cancer, greatly 59 increase the chance of active infection. This high level of latent infection suggests a period of close co-60 evolution of *Mycobacterium tuberculosis* and its human host.² Phylogenetics indicate that the 61 Mycobacterium tuberculosis complex (MTBC) emerged via an evolutionary bottleneck and that 62 existing lineages have emerged after a succession of unidirectional deletion events.³ M. tuberculosis is 63 an obligate pathogen and has no environmental reservoir. There appears to be an association between 64 65 *M. tuberculosis* lineages with different human populations around the globe and this persists within modern cities with a population of diverse origins.⁴ An association has been found between population 66 density and the emergence of human infectious diseases.⁵ This association is apparent in the early 67 Neolithic period of human development and in the case of TB appears to be continuing with the 68 emergence of highly transmissible and virulent strains of *M. tuberculosis* in major cities that have a 69 70 long record of continuous habitation.⁶

71 Leprosy is a chronic human infection caused by Mycobacterium leprae. This has declined in recent years but caused approximately 219,000 new cases in 2011,⁷ mainly in South East Asia, Africa and 72 73 South America. It is a major cause of preventable disability and of social exclusion due to stigma. M. 74 *leprae* is extremely slow growing and requires to be in an intracellular environment within a host, primarily human. *M. leprae* also appears to have experienced an evolutionary bottleneck and 75 subsequent clonal expansion between pathogen and host.⁸ Different strains of *M. leprae* can be 76 77 distinguished by variable number tandem repeat (VNTR) and short tandem repeat typing. These can 78 indicate short-term transmission via microsatellite analysis but are unstable due to poor DNA repair by *M. leprae.*⁹ Stable long-term changes can be monitored by synonymous single nucleotide 79 polymorphisms (SNPs) and these have identified lineages that are also associated with different human 80 populations.¹⁰ 81

82 Much can be inferred by the study of modern isolates of both *M. tuberculosis* and *M. leprae*.

83 However, the direct study of ancient specific biomarkers for these pathogens, such as ancient DNA

84 (aDNA) and cell wall lipid biomarkers, has distinct advantages.¹¹ These biomarkers enable

85 confirmation of infection in skeletal or soft tissue remains with non-specific or no palaeopathology, as

86 only about 5% of TB cases are believed to result in bony changes. Ancient biomarkers may also answer

historical questions, such as the nature of pre-Columbian TB or the role played by the slave trade
across the Atlantic in the dispersal of TB and leprosy to the Americas. Mixed infections can also be
detected. Analysis of aDNA may permit the determination of genetic lineages, genotypes or subgenotypes in specimens of known age, thus providing real time calibration of the date of their
emergence. The association of ancient pathogen genotypes with different host populations may also
pinpoint and date human migrations.^{10,12}

93 A useful approach in palaeomicrobiology is to obtain independent verification of findings by seeking 94 different specific biomarkers in individual specimens. Our group has concentrated on initial examination for aDNA and subsequent independent analysis of mycobacterial specific cell wall 95 lipids.¹³ Until recently, our aDNA data have been obtained by DNA amplification using PCR, first 96 97 conventional single-stage or nested PCR, and more recently using real-time PCR with specific 98 fluorescent probes and primers for selected target regions. As aDNA is often highly fragmented, the 99 use of specific probes has been very productive as they enable specific detection of selected target 100 regions of as little as 60–80 base pairs (bp). The development of Next Generation Sequencing (NGS) 101 and sophisticated bioinformatic analysis has enabled sequencing and analysis of non-amplified DNA using targeted enrichment approaches.^{14,15} Rarely, in exceptionally well-preserved material, it is 102 possible to perform shot-gun sequencing without target enrichment and to obtain an analysis of the 103 entire DNA within a sample. This has been achieved for *M. leprae* found in mediaeval dental pulp¹⁵ 104 and in extremely well preserved lung tissue from a naturally mummified individual from 18th century 105 Vác, Hungary.¹⁶ 106

107

108 **2. Ancient DNA methodology**

109 2.1 Extraction of M. tuberculosis and M. leprae aDNA

110 The following protocol gives sufficient time for samples to be disaggregated, but includes vigorous 111 bead beading and snap freezing in dry ice to release aDNA from association with any residual lipids 112 from the lipid-rich mycobacterial cell wall. Small samples (bone scrapings 20-80 mg; mummified 113 tissue10-40 mg) are taken from human remains, according to recommended protocols for aDNA with separate rooms and equipment for different stages of the process.¹⁷ Skeletal material is crushed in a 114 sterile pestle and mortar and samples are added to 400 µl of Proteinase K/EDTA. The slurry is 115 116 incubated at 56°C and mixed on a bead beater daily until solubilised. An aliquot is treated with 40 µl of 0.1 mol⁻¹ of N-phenacylthiozolium bromide (PTB), to cleave any covalent cross-links thus enabling 117

DNA strand separation and amplification.¹⁷ As PTB is inhibitory in the PCR reaction, an aliquot 118 119 without PTB is processed in parallel, so that short DNA fragments can be precipitated from PTB-free 120 silica supernatants (see below). Sample tube contents are transferred into lysis buffer containing 5 mol⁻¹ 121 guanidium thiocyanate and incubated for 1–3 days at 56°C. To complete the disruption of bone and any 122 mycobacterial remnants, samples are boiled, then snap-frozen in liquid nitrogen and thawed in a 65°C 123 water bath. This procedure is repeated twice. Sample tubes are centrifuged at 5000g for 15 mins at 5° C 124 and the supernates carefully removed into clean, sterile tubes. DNA is captured with silica suspension (NucliSens[®]) and mixed on a rotator wheel for 1 hour. Tube contents are centrifuged and silica pellets 125 washed once with wash buffer (NucliSens[®]), twice with 70% (v/v) ethanol (-20 °C) and once with 126 127 acetone (-20 °C). After drying in a heating block, DNA is eluted using 60µl elution buffer 128 (NucliSens[®]), aliquoted and used immediately or stored at -20°C. Silica supernates (500 µl) from PTB-129 negative samples are also taken from the lysis buffer, and 2.0 ml screw-capped Eppendorf tubes used to 130 wash the silica. After chilling at 5°C, supernates are mixed vigorously for 20 sec with 200 µl of Protein 131 Precipitation Solution (SLS Ltd, UK) and centrifuged for 3 min at 10,000 g. Any pellet is discarded and 132 600 μl isopropanol (-20°C) added to 550 μl of each supernate. Tubes are mixed by inversion 50 times and spun 3 min. Supernates are discarded and tubes washed once with 500 µl 70% ethanol (-20°C). 133 After draining, tubes are dried in a heating block. Any precipitated DNA is re-hydrated with 60µl 134 elution buffer (NucliSens[®]), aliquoted and used immediately or stored at -20°C. Negative extraction 135 136 controls are processed in parallel with the test samples.

137 2.2 DNA amplification and detection

138 In the current protocol, two specific regions of each organism are targeted, using repetitive elements 139 to increase the likelihood of detection of pathogen aDNA. For the M. tuberculosis complex, IS6110 (1-25 copies/cell) and IS1081 (6 copies/cell) are used.¹⁸ For M. leprae, RLEP (37 copies/cell) and 140 REPLEP (15 copies/cell) are used.¹⁹ Initially, conventional PCR was used, with primers targeting DNA 141 142 regions of around 90-bp to 123-bp. PCR was performed in two stages, with 45 rounds of amplification 143 followed, if necessary, by a nested reaction using internal primers, with a further 25 cycles of 144 amplification. PCR products were detected by agarose gel amplification, gel slices were removed, the 145 PCR products purified and sequenced. As aDNA is highly fragmented specific primers and fluorescent 146 probes have since been designed to enable shorter DNA fragments to be specifically detected (Table 1). The Qiagen QuantiTect[®] Probe reaction mix is used with additional 2 mM BSA to reduce PCR 147 148 inhibition and additional 2.0 mM MgCl₂ to facilitate primer binding. A hot-start *Taq* polymerase is 149 used to minimise non-specific primer and template binding. Negative DNA extraction and PCR

150 controls are processed alongside the test sample. Amplification is performed in a final volume of $25 \ \mu l$

using the Qiagen RotorGene[®] real-time platform. After enzyme activation for 15 min at 95°C,

amplification consists of 50-55 cycles of strand separation at 94° C for 10 sec, primer binding at 60° C

153 for 20 sec and strand extension at 72°C for 10 sec. The probes enable direct observation of specific

amplicons and the determination of cycle threshold (Ct) indicates relative concentration of template.

155 Findings may be confirmed by sequencing. Analysis of cell wall lipid biomarkers, based on the direct

detection of cell wall components without any amplification of the signal, enables independent

157 verification of the presence of the target pathogen in the sample.^{13,20}

158

159 **3. Significant questions answered by aDNA studies**

160 3.1 The M. tuberculosis lineages prevalent in early human history

TB is spread by infectious aerosols from an infected person, which results in lung infections, 161 162 although bacteria may spread via the bloodstream to all other parts of the body. Infection of the 163 vertebrae results in pathology typical of TB, such as Pott's disease, that was recognized in ancient 164 Egypt and early Neolithic communities. However, in the majority of cases there is non-specific 165 palaeopathology or none at all, so the extent of past TB infections was greatly underestimated.¹¹ However, it was noted that there was an increase in TB in the Neolithic compared with hunter-166 167 gatherers and this appeared to be associated with animal domestication. This led initially to an 168 assumption that human TB was derived from animals and that Mycobacterium bovis, the principal 169 cause of TB in domesticated animals, was the ancestor of *M. tuberculosis*. This was disproved once the phylogenomics of the MTBC was determined,² that demonstrated *M. tuberculosis* was more ancestral 170 171 than *M. bovis*. The earliest demonstration of the MTBC, in Pleistocene bison, suggested that the pathogen resembled *M. tuberculosis* or *M. africanum*²¹ but at that time the significance of the smooth 172 173 colony "Mycobacterium canettii" group was not realized. This very diverse group is believed to be 174 most similar to the original ancestor of the MTBC and demonstrates greater variability than any other member of the MTBC.^{2,22} 175

M. tuberculosis aDNA from ancient Egypt that had not experienced the TbD1 deletion has been
reported from Ancient Egypt.²³ Such strains are still extant in the Far East around the Pacific Rim and
are believed to be the oldest *M. tuberculosis* lineage. However, the majority of modern *M. tuberculosis*strains are TbD1-deleted and this lineage was demonstrated 9000 years ago in the pre-pottery Neolithic
site of Atlit-Yam, in the Eastern Mediterranean.²⁴ *M. africanum* was found in Middle Kingdom ancient

181 Egypt²³ but *M. bovis* is very rare in the archaeological record. It was found in a group of Iron Age 182 Siberian pastoralists (4th century BC – 4th century AD) who over-wintered in huts with their animals.²⁵

183 The interest in *M. tuberculosis* lineages has been driven partly because of the realization that the 184 clinical presentation of TB results from a combination of factors related to host susceptibility and *M.* 185 *tuberculosis* virulence.² This can be studied directly in past populations, allowing the effects of 186 industrialization, population density and large population movements to be examined.^{3,11}

187 3.2 The nature of past M. leprae infections

188 Leprosy was recognized in antiquity by the characteristic clinical symptoms. M. leprae targets nerves and the bacteria invade the Schwann cells that are essential for the transmission of nerve 189 190 impulses. The clinical presentation depends upon the host immune reactions. A predominant humeral 191 response leads to multibacillary, or lepromatous leprosy, where there is a strong antibody response that 192 is useless. This is the form of leprosy that results in gross changes to the nasopharyngeal region, the 193 hands and feet, with destruction of tissue and gross deformity. However, in the presence of an effective 194 cell-mediate host response, symptoms are minor although late autoimmune reactions can lead to 195 destruction of nerve function and disability. The disease has been described in ancient China, Egypt 196 and India⁷ although it is sometimes difficult to distinguish between leprosy and other diseases with 197 similar external symptoms. In archaeological cases, it is lepromatous leprosy that is recognized. Leprosy occurred during the Roman empire¹⁰ and was spread by traders and invading armies. 198

199 A phylogenetic study of global *M. leprae*, that included both modern and aDNA, demonstrated a 200 clear link between global populations and the *M. leprae* genotype and subgenotypes, as determined by SNPs.^{10,19} It appears that SNP type 2 strains are associated with early strains that migrated from the 201 202 Middle East to South East Asia but recent work shows they also spread westwards to northern and 203 western Europe.¹² SNP 3 strains are found in North Africa and the Eastern Mediterranean and were very common in Mediaeval Europe. Different migratory routes were suggested for the spread of M. 204 205 *leprae* from the proposed source near the Horn of Africa – a land route from the Mediterranean east to 206 central China, and sea routes via India and South East Asia. The genotypes found today in these regions support this theory of dispersal.²⁶ 207

208 3.3 Past co-infections

The ability of aDNA studies to detect and characterize mixed infections had not been an original aim and it was by chance that co-infections of *M. tuberculosis* and *M. leprae* were detected.²⁷ Using

conventional PCR, a decision has to be made on which organisms to target in a sample and choice of

primers is made accordingly. Mixed infections were discovered when mediaeval leprosy samples were examined for evidence of *M. tuberculosis* after contemporary co-infections were reported. This has led to an on-going debate on the possible sequence of events in the decline of leprosy in late Mediaeval Europe and whether TB had any role in bringing this about.

216 Thereafter, other examples of co-infections were sought. Another disease known in antiquity was 217 malaria, in particular that caused by *Plasmodium falciparum*. Co-infection of *M. tuberculosis* and 218 parasites is an important public health problem in co-endemic areas of the world today, and is therefore 219 likely to have been so in the past. This has been demonstrated in ancient Lower Egypt dating to c. 800 BC, where four mummies were found with aDNA from both *M. tuberculosis* and *P. falciparum*.²⁸ 220 221 Intestinal and systemic parasites were widespread in the past, also was the carriage of ectoparasites. In 222 addition, it is highly likely that future whole genome studies will identify multiple bacterial and viral 223 infections within individual human archaeological remains, in addition to associations of infection with 224 co-morbidities such as host immune or genetic disorders and cancer. Such complex scenarios form the 225 backdrop to the emergence of modern pathogens and we should endeavour to increase our 226 understanding of the factors involved.

227 Whole genome sequencing makes it feasible to detect different strains of the same pathogen within a host. This scenario was demonstrated in 18^{th} century naturally mummified lung, where two M. 228 229 *tuberculosis* strains were detected, apparently one more ancestral than the other and both resembling a modern outbreak strain, which is closely related to modern Haarlem and Erdman strains.¹⁶ This may be 230 relevant as a recent study of current mixed strain infections²⁹ found that the Haarlem and Beijing 231 genotypes are more likely to occur in a mixed infection than any of the other genotypes tested 232 233 suggesting pathogen-pathogen compatibility. There is evidence for intra-strain gene flux in M. tuberculosis²² and this is likely to be significant in the emergence of modern M. tuberculosis strains 234 that are rapidly diversifying, due to mutation, recombination and natural selection.³⁰ 235

236

4. Conclusions

During the past twenty years, since study of the palaeomicrobiology of human infectious diseases became feasible, the nature of the research questions addressed has broadened in scope and become far more sophisticated. Early palaeopathologists wished for validation of their diagnoses that were based on skeletal morphological changes. However, it soon became clear that the scale of past TB infections was far greater than previously envisaged, as many infections do not have skeletal involvement.

Palaeomicrobiology has been used to answer historical questions, such as whether the European colonialists brought TB to South America. Indeed they did, but pre-Columbian TB existed and its nature is still the subject of study. Population studies of past TB and leprosy enable epidemiological studies from the pre-antibiotic era. Information on living and burial conditions can highlight whether there was social stigma or whether infected individuals were integrated into their society.

As *M. tuberculosis* and *M. leprae* are obligate pathogens, their geographical distribution illustrates past human migrations or dispersal around the globe. Recent developments in genomics have increased our understanding of modern strains of *M. tuberculosis* and *M. leprae*, thereby providing comparators for pathogen aDNA. It is now appreciated that palaeomicrobiology enables direct comparison of ancient and modern lineages. One of the greatest benefits is that palaeomicrobiology enables direct calibration of the timescale over which changes have occurred, in the absence of modern evolutionary pressures caused by antimicrobial therapy and mass transport around the globe.

255

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266 Author contributions

HD and MS conceived the original aDNA studies and DM and GB the lipid work. MS, IP and IS

268 collected or supplied specimens. HD performed experiments and analyzed aDNA data. JO'G designed

- the PCR probes and primers. OL and HW performed lipid experiments. DM, GB, OL and HW
- analyzed data. HD wrote the manuscript and all authors approved the final version.
- 271 Competing interests
- None declared.

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367 **Table 1**

368 PCR primers and probes to detect *M. tuberculosis* (IS6110, IS1081) and *M. leprae* (RLEP, RepLep)

PCR locus	Primer name	Primer sequence (5'–3')	Amplicon
			size (bp)
IS6110	6110 Probe	5'-FAM-ACCTCACCTATGTGTCGACCTG-BHQ1-3'	,
IS6110	6110F	CACCTAACCGGCTGTGG	
IS6110	6110R	TGACAAAGGCCACGTAGG	75
IS1081	1081 Probe	5'-FAM-GGGCTACCGCGAACGCA-BHQ1-3'	
IS1081	NF	TGATTGGACCGCTCATCG	
IS1081	NR	CTTGATGGGGGGCTGAAGC	72
RLEP	RLEP Probe	FAM-5'- CTCAGCCAGCAAGCAGGCAT-3'-BHQ2	
RLEP	RLEPF	CGCTGGTATCGGTGTCG	
RLEP	RLEPR	ACACGATACTGCTGCACC	80
REPLEP	REPLEP	5'-FAM-CATGTCTATCTCCGTACGCAGCTG-	
	Probe	BHQ1-3'	
REPLEP	REPLEPF	GACTGTACTTCTTGGCCAGC	
REPLEP	REPLEPR	GCAAGGTGAGCGTTGTGG	66
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