Death metal: Evidence for the impact of lead poisoning on childhood health within the Roman Empire

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Funding information
NERC IAPETUS, Grant/Award Number: NE/L002590/1

Abstract
The use of lead was ubiquitous throughout the Roman Empire, including material for water pipes, eating vessels, medicine, and even as a sweetener for wine. The toxicity of lead is well established today, resulting in long-term psychological and neurological deficits as well as metabolic diseases. Children are particularly susceptible to the effects of lead, and it is likely that the widespread use of this deadly metal among Roman populations led to a range of adverse health effects. Indeed, lead poisoning has even been implicated in the downfall of the Roman Empire. This research examines, for the first time, the direct effect of lead poisoning on the inhabitants of the Empire. It explores whether the dramatic increase in lead during this period contributed to the failure to thrive evident within the skeletal remains of Roman children. Lead concentration and paleopathological analyses were used to explore the association between lead burdens and health during the Roman period. This study includes 173 individuals (66 adults and 107 non-adults) from five sites, AD 1st–4th centuries, located throughout the Roman Empire. Results show a negative correlation between age-at-death and core tooth enamel lead concentrations. Furthermore, higher lead concentrations were observed in children with skeletal evidence of metabolic disease than those without. This study provides the first bioarchaeological evidence that lead poisoning was a contributing factor to the high infant mortality and childhood morbidity rates seen within the Roman world.

KEYWORDS
bioarcheology, ICP-MS, infant mortality, lead concentrations, tooth enamel

1 | INTRODUCTION

Few historical subjects evoke more fervent debate than what brought about the fall of the Roman Empire. For centuries, scholars have put forth arguments for a plethora of singular causes for its decline, positing everything from the conversion to Christianity, to environmental catastrophe in the wake of a volcanic eruption (Gilfillian, 1990; Harper, 2017). It is, however, the notion that lead poisoning was a key contributing factor behind its decline that has captured the interest of scholars and general enthusiasts alike. The urban myth-like quality of this theory has ensured its endurance. Historical texts describe a range of maladies associated with lead poisoning, affirming that Roman populations did indeed suffer the deleterious effects of lead toxicity (Lessler, 1988; Needleman, 2009; Retief & Cilliers, 2006;
Diachronic trends in British human tooth enamel lead concentrations (ppm) from the Neolithic to the 19th century. Adapted from Montgomery et al. (2010), with additional Roman data from Shaw et al. (2016) and 18th–19th century data from Millard et al. (2014). The red dashed line indicates the threshold at which symptoms of lead induced metabolic disease begin to manifest (1.5 ppm) [Colour figure can be viewed at wileyonlinelibrary.com]
2.2 Osteological analyses

Adult sex was determined using the sexually dimorphic traits of the pelvis and skull as described by Phenice (1969), Ferembach et al. (1979), and Walker (2005). Following standard practice (e.g., Buikstra & Ubelaker, 1994), sex assessment was not attempted for the non-adult individuals, all of which were categorized as indeterminate.

### TABLE 1

| Age category | Site                  | Total |
|--------------|-----------------------|-------|
|              | Alba Iulia (Romania)  | 7     |
|              | Beirut (Lebanon)      | 0     |
|              | Barcelona (Spain)     | 1     |
|              | Tarragona (Spain)     | 1     |
|              | Lisieux (France)      | 1     |
| Fetal        |                       | 10    |
| 0–1 year     |                       | 22    |
| 2–6 years    |                       | 33    |
| 7–12 years   |                       | 28    |
| 13–18 years  |                       | 14    |
| Adult (18 + years) |                   | 66    |
| Total        |                       | 173   |
Non-adults were categorized as those aged less than 18 years, assessed using dental development (AlQahtani et al., 2010), long bone length (Scheuer et al., 1980; Scheuer & Black, 2000) and epiphyseal fusion (Scheuer & Black, 2000). Adult age was estimated using morphological changes in the pubic symphysis (Brooks & Suchey, 1990), the auricular surface (Buckberry & Chamberlain, 2002; Lovejoy et al., 1985), cranial suture closure (Meindl & Lovejoy, 1985), and dental wear (Brothwell, 1981).

2.3 Paleopathological analyses

Lead is an insidious poison and the gradual accumulation of the metal in bodily tissues becomes increasingly toxic. Due to the systemic nature of lead poisoning, the clinical manifestations of toxicity are varied and complex. With the exception of lead lines visible at the growing ends of bones (metaphyses) on radiographs, no specific skeletal lesions have been associated with lead poisoning (Rabinowitz et al., 1993). This is most likely due to the toxicodynamics of absorbed lead culminating in clinical manifestations that are common to many other disease processes. However, with its propensity to disrupt metabolic pathways, it is unsurprising that both modern and historical clinical literature associate lead poisoning with a number of metabolic diseases, such as rickets, scurvy and anemia (Rabinowitz et al., 1993; Caffey, 1938; Waldron, 1966; Smith et al., 1938). Therefore, it is probable that individuals who died suffering the ill effects of chronic lead poisoning would exhibit pathological skeletal alterations consistent with these metabolic diseases.

Paleopathological analysis of the non-adult individuals focused on the identification of metabolic diseases associated in the clinical literature with lead poisoning (Landrigan, 1989; Landrigan & Todd, 1994). The following section outlines the paleopathological features of the metabolic stress often associated with lead poisoning and the parameters used to diagnose these diseases within the skeletal assemblages. No radiographs were available for any of the study sites; therefore, diagnoses were made solely from macroscopic examination of...
individuals with observable elements. Examples of the skeletal manifestation of these metabolic diseases are presented in Figure 3.

2.3.1 | Rickets

Rickets is generally identified by bowing of the long bones and/or the presence of widened, cupped and porous/frayed (“brush end”) epiphyses, sternal rib-end flaring (“rachitic rosary”) and cranial vault thinning (Waldron, 2009, p. 129). Additional manifestations of the disease in non-adults can also include orbital roof porosity, deformation of the mandibular ramus, porosity of the sternal rib-ends and deformation of the ribs (Brickley et al., 2005; Brickley & Ives, 2010; Mays et al., 2006; Ortner, 2003; Ortner & Mays, 1998). Using the published diagnostic criteria outlined in Table 2, macroscopic lesions were recorded as either present or absent. Because many of the lesions associated with rickets can have multiple etiologies (Mays et al., 2006; Ortner & Mays, 1998), a diagnosis of rickets was only recorded if three or more probable rachitic features were present, or if there were bending deformities of the long bones plus one other feature. This was done to avoid over diagnoses of rickets in the sample population. Individuals exhibiting no probable features but three or more possible features alongside any non-diagnostic features were considered as possibly rachitic. Using Ortner and Mays (1998) definition, a distinction was also made between healed and active rickets.

2.3.2 | Scurvy

Pathological alterations indicative of scurvy primarily consist of abnormal cortical porosity (often with subperiosteal new bone formation) on the ectocranial surface, scapulae, long bone metaphyses, and mandible (Schattmann et al., 2016; Snoddy et al., 2018; Stark, 2014). These lesions tend to manifest bilaterally and are thought to be caused by chronic, low-grade hemorrhage of weakened blood vessels, predominantly at muscle attachment sites, which stimulates an inflammatory response (Ortner et al., 1999, 2001; Ortner & Ericksen, 1997). Although abnormal cortical porosity is the primary lesion associated with scurvy, it is also common to many other disease processes such as specific and non-specific infection, hemoglobinopathies, anemias, and other metabolic disorders (Brown & Ortner, 2011; Lagia et al., 2007). It is therefore important to analyze the porosity in relation to its distribution across the entire skeleton. Using the published diagnostic criteria outlined in Table 3, macroscopic lesions were recorded as either present or absent. In line with recommendations by Snoddy et al. (2018), individuals were recorded as scorbutic if two or more diagnostic scorbutic features were present. If the individual revealed multiple suggestive features, they were considered as possibly scorbutic.

2.3.3 | Cribra orbitalia and porotic hyperostosis

Cribra orbitalia was identified as porotic changes of the orbital roofs and recorded for each orbit using the Stuart-Macadam (1991) grading system. Any individual with cribrotic lesions over the age of 10 was designated as having the healed form of cribra orbitalia due to the red-to-yellow marrow conversion that occurs around this age, thereby precluding the formation of these lesions (Simonson & Kao, 1992). Porotic hyperostosis was identified as abnormal cortical porosity of the cranial vault, and recorded as either present or absent (Mann & Hunt, 2013, p. 28; Waldron, 2009, p. 137). Cross-sections of parietal bones that showed abnormal widening/thickening of the diploic space was also noted as potential narrow hyperplasia (thickening) of the cranial vault.

| Diagnostic category | Probable | Possible | Non-diagnostic |
|---------------------|----------|----------|---------------|
| Cranial             |          |          |               |
| Deformed mandibular ramus | Cranial vault porosity | Delayed closure of fontanelles |
|                      | Orbital roof porosity | Craniotabes (softening of bone behind ears over occipital region and adjacent to lambdoid suture) |
|                      | Layers of speculated, irregular porous bone can occur during healing when osteoid is mineralising | Frontal and parietal bossing |
| Post-cranial        |          |          |               |
| Deformed arm bones  | Flaring of sternal rib-ends | Superior flattening of the femora |
| Deformed leg bones  | Porosity of sternal rib-ends | |
| Ilium concavity     | Long bone metaphyseal flaring | |
| Altered rib angle   | Long bone thickening | |
|                     | Porous roughening of long bone metaphyses | |
|                     | Long bone concave curvature porosity | |

Note: After Brickley and Ives (2010), Hess (1930), Mays et al. (2006), Ortner and Mays (1998), and Pettifor (2011).
2.4 Lead concentration analysis

Initial sample preparation was carried out at the Archaeological Isotope and Peptide Laboratory (AIPRL), Durham University, following procedures outlined by Montgomery (2002), briefly described here: The enamel surface was abraded using a tungsten carbide dental bur to remove surface contamination. Following this, a chip of enamel was removed using a flexible diamond edged rotary saw, all exposed surfaces of the chip were abraded to remove any adhering dentine and potential sources of contamination. Enamel chips were stored in clean micro-centrifuge tubes for transfer to the National Environmental Isotope Facility (NEIF), British Geological Survey, Keyworth. All dental tools were cleaned between samples via ultra-sonication in Decon for 5 min and rinsed three times with ultra-pure de-ionized water.

Trace element analysis was carried out at NEIF using an Agilent 7500cx ICP-MS fitted with a CETAC ASX-520 autosampler. Sample introduction from the autosampler to the Inductively Coupled Plasma Mass Spectrometry (ICP-MS) was controlled by a CETAC ASXpress + vacuum pump. Multi-element quality control (QC) check standards, containing the trace elements of interest at 25 μg/L, and a separate major element QC were analyzed at the start and end of each run and after no more than every 20 samples. To overcome polyatomic interferences the ICP-MS collision cell was operated in He mode at a flow rate of 5.5 mL min⁻¹ for all analytes except Se, for which H₂ gas was used at 4.5 mL min⁻¹ due to the more intense interferences experienced with Se because of argon (Ar) dimers formed in the plasma. Samples were diluted with 1% v/v HNO₃. 0.5% v/v HCl before analysis. Quantitative data analysis was carried out using MassHunter Workstation software (Agilent),

TABLE 3 Scorbutic lesions used in the identification of scurvy and their diagnostic category

| Diagnostic category | Probable | Possible | Non-diagnostic |
|---------------------|----------|----------|----------------|
| Cranial             | Porosity and/or new bone formation on the greater wing of the sphenoid Porosity on the posterior aspect of the mandible Porosity on the temporal bone | Porosity in the mandibular coronoid fossae Porosity and/or new bone formation on the lesser wing of the sphenoid Porosity at the infraorbital foramen on the maxilla Porosity and/or new bone formation on the orbital roof Porosity and/or new bone formation on the pars basilaris | Porosity on the palate of the maxilla Porosity in the maxilla and/or mandibular alveola processes Porosity and/or new bone formation on the endocranium |
| Post-cranial        | Porosity and/or new bone formation in the supraspinous and/or infraspinous fossae | Metaphyseal flaring of long bones Flaring of sternal rib-ends | Porosity and/or new bone formation on the long bones Metaphyseal porosity |

Note: After Brickley and Ives (2006, 2010), Geber and Murphy (2012), Moore and Koon (2017), Ortner (2003), Ortner et al. (1999, 2001), and Ortner and Ericksen (1997).
3 | RESULTS AND DISCUSSION

Lead is incorporated into the mineral matrix of tooth enamel during childhood and does not alter in vivo or undergo diagenetic changes from the burial environment after the tissue mineralises; therefore, lead concentrations acquired in this way represent an individual’s childhood exposure to the toxic metal (Montgomery et al., 2010). Early studies show that deciduous tooth enamel often has higher lead concentrations than permanent tooth enamel, and this is thought to be because younger children absorb higher quantities of lead relative to older children with similar exposure (Shapiro et al., 1972). If age-related absorption rates led to higher lead concentrations in deciduous teeth, regardless of exposure level, then co-forming permanent teeth would be expected to yield similarly high lead concentrations. However, a recent study shows no significant difference in median tooth enamel lead concentration between earlier and later permanent tooth types (e.g., second and third molars) (Moore, 2019). This suggests that age-related lead absorption rates are not the dominant

FIGURE 6 Prevalence of metabolic diseases in the observable (a) non-adult (n = 65) and (b) adult (n = 51) sample populations [Colour figure can be viewed at wileyonlinelibrary.com]
factor influencing lead concentrations in tooth enamel, and that comparing lead concentrations in deciduous and permanent tooth enamel provides a useful means of investigating the effects of lead exposure on childhood health (ibid). In this study a comparison of the lead concentration data from the adult and non-adult individuals show that those who survived into adulthood had lower childhood lead burdens (median = 2.6 ppm) than those who died during childhood (median = 7.2 ppm) (see Figure 4). The results of a Kruskal-Wallis test showed that the median lead concentrations in these two groups were statistically significantly different ($X^2 = 12.181, p = 0.0005$). Children have more than double the lead concentrations observed in adults, suggesting that higher lead burdens are accompanied by lower life expectancies. These results offer the first bioarchaeological evidence that lead poisoning resulted in increased frailty for citizens of the Roman Empire.

It is evident from the archeological record that there is a real failure to thrive in children throughout the Roman period (Carroll, 2014; Rohnbogner, 2017; Rohnbogner & Lewis, 2017). It is estimated that up to 50% of children died before the age of 10 years old, with 20–40% of these not reaching the age of 1 year (Carroll, 2014, 2018). Children are more susceptible to lead poisoning than adults as their developing bodies are prone to absorbing higher quantities of ingested lead. To explore whether the high lead burdens characteristic of Roman individuals contributed to the high childhood mortality rates in the Roman Empire, lead concentrations were further compared to age-at-death (see Figure 5). A negative correlation between lead concentration and age-at-death is evident, again indicating that individuals with lower lead burdens lived longer than those with higher lead burdens. This is particularly interesting with regards to children under the age of 1 year. Explanations for high infant mortality rates during the Roman period have ranged from malnutrition and disease to infanticide and exposure (Gowland et al., 2014; Mays, 1993; Pilkington, 2013; Rohnbogner, 2017). The results of this study offer new insights into the previously overlooked role that lead may have played in these high infant mortality rates. Unfortunately, little research has been done to understand how lead concentrations in tooth enamel reflect in vivo lead burdens, or how they correlate to manifestations of lead poisoning during life (Grobler et al., 2000; Rabinowitz et al., 1993). As such, identifying high lead concentrations in archeological remains alone is unlikely to be sufficient to determine those who may have died from lead poisoning. However, using modern clinical literature and the known biochemical pathogenesis of lead toxicity it may be possible to further elucidate the effect of lead poisoning on the health of archeological populations.

Of the sample population, 65/107 non-adults and 51/66 adults demonstrated suitable preservation for paleopathological evaluation. Results indicate that 46/65 (71%) non-adults with observable elements exhibit pathological lesions diagnostic or consistent with rickets, scurvy, and/or cribra orbitalia and porotic hyperostosis (see Figure 6, S1). This is considerably lower when compared to the adult “survivor” population, of which 15/51 (29%) demonstrate lesions consistent with these metabolic diseases.

The non-adult individuals exhibiting pathological lesions diagnostic of metabolic disease had significantly higher lead concentrations (median = 8.1 ppm) than those without (median = 4.9 ppm) (see Figure 7). A Kruskal-Wallis test showed a statistically significant difference in lead concentrations between these two groups ($X^2 = 4.007, p = 0.0453$). While these metabolic diseases have multiple etiologies, the association with lead supports the presupposition that high lead concentrations are also implicated in Roman-period skeletons. Thus, elevated levels of environmental lead pollution characteristic of the Roman period did have a negative impact upon childhood health. In future, it is worth exploring if differences in lead concentrations exist chronologically (i.e., earlier vs. later Roman contexts), examining the extent to which geographical differences influence childhood lead burdens, and exploring how lead isotope ratios vary between different regions to better understand mobility and trade networks within the Empire.

4 | CONCLUSION

Through the combination of paleopathological and trace element analyses, lead poisoning can tentatively be identified in archeological human remains. The results of this study demonstrate that increased exposure to anthropogenically produced lead was a contributing factor to the ill health and failure to thrive seen in many Roman
infants and children, thereby providing the first bioarchaeological evidence that lead poisoning may have been a contributing factor to the high infant mortality rates seen in Roman skeletal populations. The introduction of a bioarchaeological perspective to the decades old debate regarding the impact of lead on Roman health offers new insights into the effects of environmental lead pollution on child health during this period.

ACKNOWLEDGMENTS
The authors are grateful to the Museu Nacional Arqueologia de Tarragona (MNAT), Museu d’Història de Barcelona (MUHBA), Université de Caen, Université “1 Decembre 1918” Alba Iulia, the Lebanese Ministry of Culture, and Lebanese Directorate General of Antiquities (with special thanks to Drs. Assaad Seif and Georges Abou Diwan) for access to skeletal material and site information. Simon Chenery (NIGL/BGS) is thanked for his support and assistance in processing the trace element analysis, and the authors thank Sal Kellett (Durham University) for producing Figure 2. Funding to J. Moore under NERC IAPETUS studentship NE/L002590/1 supported this research.

CONFLICT OF INTEREST
The authors declare that they have no conflict of interest.

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