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Tuberculosis-like respiratory infection in 245-million-year-old marine reptile suggested by bone pathologies

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An absence of ancient archaeological and palaeontological evidence of pneumonia contrasts with its recognition in the more recent archaeological record. We document an apparent infection-mediated periosteal reaction affecting the dorsal ribs in a Middle Triassic eosauropterygian historically referred to as 'Proneusticosaurus silesiacus'. High-resolution X-ray microtomography and histological studies of the pathologically altered ribs revealed the presence of a continuous solid periosteal reaction with multiple superficial blebs (protrusions) on the visceral surfaces of several ribs. Increased vascularization and uneven lines of arrested growth document that the pathology was the result of a multi-seasonal disease. While visceral surface localization of this periosteal reaction represents the earliest identified evidence for pneumonia, the blebs may have an additional implication: they have only been previously recognized in humans with tuberculosis (TB). Along with this diagnosis is the presence of focal vertebral erosions, parsimoniously compared to vertebral manifestation of TB in humans.

1. Introduction

Pathological conditions observed in fossil skeletal remains provide an exceptional opportunity for epidemiologic study of
disease. Recognition of pathogens permits investigation of how disease affected ancient biota [1,2]. The fossil record, although very selective and incomplete, allows resolution of gaps in our understanding of their evolution and transmission.

Osseous abnormality/pathology is relatively common in reptiles [3,4]. Most of the conditions described thus far have been identified in terrestrial or semi-terrestrial taxa. They include post-traumatic malformation [5,6], congenital defects [7,8] and neoplasms [3,9–11]. Previous reports of Mesozoic marine reptile pathology have been limited to recognition of bone necrosis, of both aseptic [12] and infectious [13] aetiology. Aseptic avascular necrosis associated with decompression syndrome is quite common in ancient marine reptiles, including sea turtles [14,15], mosasaurs [16], ichthyosaurs [12] and sauropterygians [12,17]. The fossil record of infectious disease in marine reptiles is scarce, limited to bacterial infections in Triassic sauropterygians [13] and Cretaceous mosasaurs [18,19]. Fossilization of actual microorganisms is exceptionally rare (e.g. Yersinia strains have been found in Eocene flea) [20].

Pneumonia, the result of infection of lung parenchyma, affects 450 million people yearly [21]. It is one of the oldest diseases known to humankind, described for the first time by Hippocrates [21]. Among the organisms responsible for bacterial pneumonia is the trans-phylogenic zoonosis tuberculosis (TB), caused by Mycobacterium tuberculosis and related organisms. This contagious disease has been previously documented in reptiles [2,3,22–24] and marine tetrapods [22,25–27]. It is commonly recognized in the archaeological record [2,28–32] and also documented in Pleistocene bovids [33–35]. Diagnosis of TB, on the basis of osteological data, M. tuberculosis lipid biomarkers and DNA sequence detection, was established by Hershkovitz et al. [36] in Neolithic human bones and by Rothschild et al. [33] and Lee et al. [37,38] in Pleistocene bison. Vertebral collapse and coalescence, classic complications of TB [2,39], were also documented in a Pliocene macropod (marsupial) [2]. Molecular data suggest that M. tuberculosis diverged from its common ancestor as recently as 17 000 years ago [33,38], but that perspective is difficult to reconcile with actual recognition of the disease in 17 000 and 75 000–100 000 ybp bison [33]. Thus, the antiquity of TB has not been established.

Herein, we report a TB-like respiratory infection in a Triassic (approx. 245 million ybp) marine reptile. The presumptive diagnosis is established on the basis of macroscopic, radiological and histological study of its rib and vertebrae pathology.

2. Methods

2.1. Thin-sectioning

The proximal portion of the rib fragment was thin-sectioned and examined using a Nikon Eclipse 80i transmitted light microscope fitted with a DS-5Mc cooled camera head. The distal part of the same specimen, around 0.5 cm long, was subjected to X-ray microtomography (XMT) scanning.

2.2. X-ray microtomography

Radiologic evaluation was performed with an Xradia MicroXCT-200 imaging system, equipped with a 90 kV/8 W tungsten X-ray source. The scans were performed using the following parameters: voltage: 50 kV; power: 6 W; exposure time: 4 s; voxel size: 4.61 µm. Images were reconstructed with the XMReconstructor software provided with the Xradia system. Three-dimensional images of bone and XMT sections were obtained by processing with the AVIZO 7.0 Fire Edition software.

3. Results

Pneumonia-related bone involvement was recognized on the visceral surface of dorsal ribs of the holotype MG UWr 4438 (figure 1a,c; electronic supplementary material, figure S1) of the Middle Triassic marine reptile ‘Proneusticosaurus’ silesiacus Volz [40]. We agree with previous authors [41–43], who interpreted the stated genus ‘Proneusticosaurus’ as actually representing the postcranial remains of another taxon, Cymatosaurus. The latter, however, is known thus far only from cranial material. Complete description of the specimen, associated vertebral collapse and fusion of vertebral centra (typical of TB-related Pott’s disease in humans) and additional illustrations of rib pathology are included in the electronic supplementary material.

Bone reaction (figure 1b) is present on the visceral surface of at least four consecutive dorsal ribs on the left side of the animal (figure 1b). The lesions are elliptical in shape, ranging in size from 0.5 to 3 mm, and appear contiguous with, rather than applied to the bone surface. XMT revealed bleb-shaped thickening
The blebs have relatively smooth surfaces and an unbroken outline, although one does have serpentine bifurcating grooves on its surface (figure 1d). The grooves most probably represent the imprint of overlying blood vessels, rather than the result of irregular bone deposition. Very similar, but weaker fingerprint-like texture is also present on non-pathological regions of the ribs and was previously reported on the vertebrae and appendicular skeleton of other sauropterygians (e.g. pachypleurosaurids) [44].

Histological documentation of penetrating branching channels reveals the infectious nature of the pathologic process. The channels appear limited to the internal aspect of the inflated outer cortex. Underlying bone was otherwise unaffected, except for increased local vascular supply. The ribs are composed of a lamellar-zonal bone tissue, with very minimal remodelling. Zonation is of greatest density and best visualized dorsally (figure 3a,b), where growth cycles consist primarily of parallel-fibred bone, grading into lamellar bone and culminating in lines of arrested growth (LAGs). Organization of collagen fibre traces in a wavy manner (figure 3c) marks external migration of blood vessels during bone growth. Some of the LAG waves appear to represent remnants of superficial vascular grooves that were subsequently incorporated into the bone.

This interpretation is supported by XMT revelation of blood vessels crossing LAGs in canal-like waves and continuing along the length of the rib. Vascularization is predominantly longitudinal in distribution (see electronic supplementary material, movie S1). Bone deposition is increased and less organized on the visceral aspect of the rib (figure 3f,g), where vessels are more radially oriented (figure 3d,e,f,g).

The outer zone of bone is sharply demarcated. Its base is dark in transmitted and milky in polarized light. However, this may be an artefact caused by an uneven thickness of the section. Its external section exhibits stratification in transmitted light, which is inconspicuous on polarization. Radial, diverging canals extend to the surface. Older LAGs contain several well-vascularized, bleb-like protrusions of comparable sizes (much larger than typical, vascular canal-scale LAG waves) and location to those present on the surface. These were subsequently covered by younger zones of less-vascularized bone (figure 2; electronic supplementary material, movie S2). This suggests a chronic disease, spanning several seasons.
Figure 2. X-ray microtomographic images and three-dimensional reconstruction of the dorsal rib fragment of *Proneusticosaurus silesiacus* holotype, MG UWr. 4438s. (a–h) XMT virtual sections (distal towards proximal) of the rib fragment showing superficial blebs (arrows). Note the increased vasculature within the blebs, and the bleb-like shape of older zones. Scale bar equals 1 mm. (i–k) Three-dimensional visualizations of several blebs on the rib surface as well as bone vasculature in the cropped fragment of cortical bone. Note the presence of a fingerprint-like system of superficial vascular grooves (i). (j–k) Not to scale. Ventral (visceral) towards the left.

4. Discussion

4.1. Differential diagnosis

The observed blebs are typical of those occasionally seen in humans with *Mycobacterium tuberculosis*-derived tuberculosis [45,46]. XMT examination documents the blebs as solid (branching channels excepted), ruling out taphonomic blister formation and that related to vitamin C deficiency (scurvy). The periosteum is separated from the underlying cortical bone in scurvy [2,47]. Potential taphonomic
Figure 3. Histology of the dorsal rib of 'Proneusticosaurus’ silesiacus holotype, MG UWr. 4438s. (a,b) The dorsal region of the rib composed of avascular lamellar-zonal bone with well-pronounced zonation, and the medullary area in transmitted light (a) and polarized light with \( \lambda \) compensation (b). Arrows show LAGs. (c) Anterior region of the rib in polarized light exhibiting the vascularization, and rate of bone deposition gradually increasing towards the ventral (visceral) region. Note the wavy organization of the tissue. (d,e) Ventro-posterior region of the rib in polarized light without (d) and with (e) \( \lambda \) compensation showing the vasculature increasing even more and attaining radial organization towards the ventrum. (f,g) Ventral region of the rib in transmitted (f) and polarized light with \( \lambda \) compensation (g), presenting the radial vasculature and the bleb. Indicated is the LAG separating the pathological outer zone of the bone (dotted line) and the area shown in panel (h). (h) Close-up of the bleb in transmitted light. Scale bars for panels (a–g) equal 0.5 mm, for panel (h) equals 0.1 mm. In all panels ventral (visceral) towards the right-hand side.

Explanation of branching channels is essentially limited to fungal disease, excluded because of size discrepancy. The diameter of fungal hyphae is five times larger [48]. Branching channels of a size similar to that of the vascular rib channels are, however, rarely noted with the branching bacterium Actinomyces [2].

There is no bone deformation, indicative of a healed fracture. Localization of the pathology to the internal (visceral) rib surface rules out a traumatic explanation. While developmental, dysplastic (e.g. fibrous dysplasia) and metabolic disorders (e.g. renal osteodystrophy) can involve ribs, their effect is limited to the internal bone structure. The periostium is not altered, in the absence of subjacent bone involvement [2,39]. One exception is melorheostosis, a phenomenon in which ‘drip-patterns’ of periosteal reaction may occur without apparent underlying bone involvement [39]. However, that is macroscopically recognized by continuous dispersal down long bones, quite different from the focal spots (blebs) observed in MG UWr. 4438s. Pathology in the latter is clearly distinguishable from that of hypertrophic osteoarthropathy, in which periosteal reaction is present on the distal diaphyses of the appendicular skeleton [49]. These were apparently unaffected in MG UWr. 4438s. Rib involvement by cancer is typically related to metastatic disease directly invading the bone [2,39,50]. No such evidence was found in MG UWr. 4438s.
Primary bone tumours (e.g. osteoid osteoma) only secondarily affect the periostea, and are recognized because of internal bone structure modification [39]. Gardner’s syndrome, caused by a dominantly inherited mutation in the adenomatous polyposis coli tumour suppressor gene, may give a superficially similar osteological morphology [51]. Additionally, bone involvement in Gardner’s syndrome is in the form of osteomas. They are spherical protuberances and do not have the flattened teardrop shape observed in our specimen. This is also true for button osteomas, which are also rounded, not tear-shaped. Further, such osteomas consist of Haversian bone [52], not the lamellar bone noted in MG UWr. 4438s. Other conditions that can produce periosteal proliferation are associated with osteolytic or osteoblastic internal bone reaction, absent in MG UWr. 4438s.

### 4.2. Rationale for suggesting tuberculosis as an inferred diagnosis for MG UWr. 4438s

The multidisciplinary approach used in the current analysis revealed a chronic, multi-seasonal disease affecting bone tissue. The presence of bone reaction and protrusions, localized to the visceral surface of the paravertebral area of the ribs, suggest a TB-type pulmonary infection, the oldest known such observation to date.

TB is well represented as a trans-phylogenetic disorder. All previously identified causative organisms have been members of the genus *Mycobacterium*, with species-dependent (phylogenetic) host specificity [23]. Most infections in humans are caused by *Mycobacterium tuberculosis*. Other mycobacterial infections (e.g. that caused by *M. kansasii*, *M. avium-intracellularis* and *M. thamnopheos*) have been recognized in extant animals, reptiles included [22–24]. Representatives of the genus *Mycobacterium* exhibit high tolerance to increased water salinity [53]. The genus is therefore believed to have emerged in the remote past as a primarily saprophytic, non-pathogenic [54] bacterium. It apparently acquired its virulence (pathogenicity) during long-term interactions with poikilothermic and subsequently with homoeothermic animals, given its wide taxonomic distribution [22,25].

The bleb-like periosteal bone apposition recognized in MG UWr. 4438s has been previously reported in humans with TB [45,46]. However, bone pathology is uncommon in humans (found in 1–2% of individuals with pulmonary disease) and blebs are even rarer. No other individual among the 10,000 animals examined to date by one of us (B.M.R.) has been found with such blebs (see also [2–4]). It is unclear why they have been so infrequently recognized. The mechanism of this particular morphology remains unknown. They may represent an unusual bone response in an immunosuppressed individual or even an aspect of disease pathophysiology (developmental course) that is so evanescent as to escape notice. It is apparent from the XMT scans of MG UWr. 4438s that the blebs developed during cycles of bone apposition. They were initially bordered by consecutive LAGs and eventually overlaid by newly deposited bone tissue during the next cycle, during which new blebs formed in neighbouring areas (electronic supplementary material, movie S2). This apparently was the response to local inflammation at the interface between the bone and surrounding tissues, possibly related to irritation by bacterial cells or their products. There is no clear evidence of gradual migration between spots. However, each bleb ends with a LAG. The bleb-inducing factor may have migrated, but no continuous record of such movement is preserved in the bone tissue, due to the cyclical arrest of bone apposition. Alternatively, the inflammation foci might have been stationary, appearing and disappearing in various locations during each cycle of bone growth. Dependence on some inducing factor, and the ability to resume normal bone growth in the area previously occupied by a bleb, allows consideration of the possibility that the blebs may be present only temporarily through the disease course, leaving no external traces. In animals (including many mammals) lacking or rarely developing LAGs or those with intensive bone remodelling, bleb ‘history’ may not be assessable. Nonetheless, determination of the occurrence and prevalence of blebs in the general zoological or anthropological record is beyond the scope of this study and will be monitored in a prospective manner.

Typical TB-related vertebral modification, referred to as Pott’s disease [55], seems present in MG UWr. 4438s (see the electronic supplementary material). Although the specimen described herein is much too old to permit molecular-based (e.g. DNA) analysis, previous studies document that morphological clues may be successfully used to diagnose TB in a trans-phylogenetic manner and through geologic time [35]. Mycobacteriosis is common in modern reptiles [22], fish [56,57] and aquatic tetrapods [25–27], related to facile transmission of *Mycobacteria* spp. in aquatic medium [22]. Virulence of an ancient member of the *Mycobacterium tuberculosis* complex or of some other closely related fossil epizootic bacterial taxon developed much earlier than previously considered.

Many pathogens transfer horizontally across species, acquiring the ability during their evolution to colonize new groups of hosts or vectors. New hosts are sometimes acquired, independent of their
evolutionary history and phylogenetic relationships. They originate as opportunists, infecting any taxa possible, only later specializing to infection or infestation of specific hosts. The susceptibility of those animals, whether pathologically affected or simply serving as vectors for transmission to disease-susceptible animals, is determined by initial (from an evolutionary perspective): (i) deficits of immunity against the given pathogen; (ii) physiological nuances (e.g. basal resting temperature); (iii) ecologically driven exposition; or (iv) newly acquired pathogen virulence [58,59]. This is illustrated by the distribution of modern pathogens, which is often spotty when it comes to taxa within families, orders or classes, but at the same time may involve more than one class. Therefore, the presence of infectious diseases in extant taxa does not necessarily indicate that their ancestors were similarly afflicted.

4.3. The impact of infection on the life of ‘Proneusticosaurus silesiacus’

Given uncertainty as to whether the early sauropterygians lived in herds or exhibited any other gregarious behaviour, it is difficult to postulate the full epidemiologic potential of TB. Modern-day epizootic studies of the spread of mycobacteriosis in free-ranging animals have been reported for fish [60,61], birds [62] and mammals [63], but data regarding reptiles are scarce. Lack of good sauropterygian analogues in modern ecosystems compromises further phylogenetic analysis at this time. However, another analytic pathway derives from consideration of ecotrophism. As the semi-aquatic lifestyle of nothosaurs is often compared to that of modern-day seals, it may be hypothesized that the character of disease transmission might have been comparable. Ecotrophism, an often overlooked aspect of organismal behaviour and spread of infectious disease, should be considered. Pinnipeds appear to be the marine mammal group most susceptible to TB [64]. Analogously, the first case of a TB-like disease is similarly found in a resident of a marine habitat.

Contrasted with extensive study of nothosaurid long bones [65–68], rib histology has received less attention [65]. Histological section of the MG UWr. 4438s rib reveals dense, very compact bone with a small medullary area (figure 3a,b). Such conditions seem to be consistent with that observed in the earliest (Early Anisian) representatives of Nothosaurus spp., characterized by increased bone mass. The latter is an adaptation, facilitating diving behaviour [65]. The ‘Proneusticosaurus silesiacus’ holotype is also characterized by relatively broad gastralia and unusually broad ribs (e.g. electronic supplementary material, figure S2), classified as pachyostotic [69]. Such adaptations are known in the secondarily aquatic tetrapods inhabiting shallow waters. They provide extra body mass, to compensate the positive buoyancy caused by atmospheric air accumulated in the lungs [70]. Contrasted with the physiology of open marine animals that are able to accumulate oxygen in the body tissues [71], the lungs in these primitive secondarily aquatic tetrapods are the main oxygen reservoir [71]. As pulmonary TB reduces respiratory function [72], breathing capacity was probably compromised in MG UWr. 4438s. This potentially constrained diving behaviour related to feeding or escape from predators. Decompression syndrome-related avascular necrosis, however, is relatively rare in nothosaurs [12,73]. Limited lung capacity might not have been detrimental for nothosaurs, if they were not routinely deep divers. While the part of the slab with the complete hand of MG UWr. 4438s is now missing, documentation provided by Volz [40] revealed neither hyperphalangy nor polydactyly. It had a typical nothosaurian autopodium structure, indicative of semi-aquatic adaptation.

It is clear that MG UWr. 4438s survived several seasons of disease, before it perished. It cannot be determined whether the death of MG UWr. 4438s was an independent phenomenon or the result of compromise of hunting efforts due to limited mobility related to diminished lung capacity or the result of TB-related failure of internal organs.

5. Conclusion

The case presented here is the oldest record of pneumonia and possibly pushes the earliest record of mycobacteriosis back to the Early Mesozoic, the dawn of the age of reptiles. Although the palaeopathological studies described herein are restricted in terms of available material and no typical microbiological or biochemical clues are accessible at this time, our diagnosis is supported by several lines of evidence—external and internal morphology of the blebs on the ribs, their distribution in the rib cage and the presence of vertebral damage comparable to that observed in Pott’s disease. This is the first evidence of these in fossil reptiles and in such a distant past. The presence of probable TB-related changes on a skeleton of an extinct reptile living before the ascent of mammals supports the view of an ancient origin of mycobacterioses and their long-lasting influence on various vertebrate taxa.
References

1. Rothschild BM. 2016 Odyssey in the evolution of a paleopathologist. Foss. Rec. 20, 37–45. (doi:10.5949/fr-20-37-2016)
2. Rothschild BM, Martin LD. 2006 Skeletal impact of disease. Albuquerque, NM: New Mexico Museum of Natural History.
3. Rothschild BM, Schultz H-P, Pellegrini R. 2012 Herpetological osteopathology. Annotated bibliography of amphibia and reptiles: Berlin, Germany: Springer Science + Business Media.
4. Rothschild BM, Schultz H-P, Pellegrini R. 2013 Osseous and other hard tissue pathologies in turtles and abnormalities of mineral deposition. In Morphology and evolution of turtles (eds DB Brinkman, PA Holroyd, JD Gardner), pp. 501–534. Dordrecht, The Netherlands: Springer Science + Business Media.
5. Hedrick BP et al. 2016 An injured Pirotosaurus (Dinosauria: Ceratopsia) from the Yixian formation (Liouning, China): Implications for Pirotosaurus biology. Anat. Rec. 299, 897–906. (doi:10.1002/ar.23363)
6. Anie J, Garwood RJ, Lowe T, Withers PL, Manning PL. 2015 Interpreting pathologies in extinct and extinct archosaurs using micro-CT. PeerJ 3, e1150. (doi:10.7717/peerj.1150)
7. Szczysielie T, Saurik D, Kapsycińska A, Rothschild BM, 2017 The oldest record of aquatic amniote congenital scoliosis. PLoS ONE 12, e0165310. (doi:10.1371/journal.pone.0165310)
8. Witzmann F, Rothschild BM, Hampe O, Sobral G, Gubin YM, Ashab P. 2014 Congenital malformations of the vertebral column in ancient amphibians. Anat. Histol. Embryol. 43, 90–102. (doi:10.1111/ahex.12050)
9. Rothschild BM, Witske BJ, Herszkovitz I. 1999 Metastatic cancer in the Jurassic. Anat Rec. 354, 980–984. (doi:10.1002/(SICI)1097-0185(199910)354:6<980::AID-AR2>3.0.CO;2-9)
10. de Souza Barbosa FH, Pereira PVG, da C, Bergqvist LP, Rothschild BM. 2016 Multiple neoplasms in a single sauropod dinosaur from the Upper Cretaceous of Brazil. Cretae. Rev. 62, 13–17. (doi:10.1615/cretes.2016.01.010)
11. Dumbraivá MD, Rothschild BM, Weishampel DB, Ciski-sava Z, Andrei RA, Achenach KA, Codrea VA. 2016 A dinozaurul facial deformity and the first occurrence of ameloblastoma in the fossil record. Sci. Rep. 6, 29271. (doi:10.1038/srep29271)
12. Rothschild BM, Xiaoting Z, Martin LD. 2012 Adaptations for marine habitat and the effect of Tetrican and Jurassic predator pressure on development of decompression syndrome in ichthyosaurs. Naturwissenschaften 99, 443–448. (doi:10.1007/s11010-012-0918-0)
13. Szczygielski T, Surik D, Rothschild BM, Dulski M, Janiszewska K. 2017 Two types of bone nodule in the Middle Tertiary Pirotosaurus longaevus bones: the results of integrated studies. R. Soc. open sci. 4, 170204. (doi:10.1098/rsos.170204)
14. Rothschild BM. 1987 Decompression syndrome in fossil marine turtles. Ann. Carnegie Mus. 56, 253–358.
15. Rothschild BM, Naples V. 2013 Decompression syndrome and diving behavior in Odontochelys, the first turtle. Acta Palaeontol. Pol. 69, 161–167. (doi:10.4202/app.2012.0113)
16. Rothschild BM, Martin LD. 1987 Avascular necrosis: occurrence in diving Cretaceous mosasaurs. Science 236, 75–77. (doi:10.1126/science.236.48177.75)
17. Rothschild BM, Stoens GW. 2003 Decompression syndrome in plesiosaurs (Sauropterygia: Reptilia). J. Verteb. Paleontol. 23, 324–328. (doi:10.1002/jvp.2003.0230230402.0.02)
18. Linham-Sollar T. 2004 Palaeopathology and injury in the extinct mosasaur (Lepidosauromorpha, Squamata) and implications for modern reptiles. Lethaia 37, 255–262. (doi:10.1080/0023560X.2004.106659)
19. Schulz AS, Walsenberg GH, Hoffman PA, Stuyp Y, Rothschild BM. 2006 Chronic bone infection in the jaw of Mosasauros hoffmanni (Squamata). Oryctos 6, 41–52.
20. Poinar Jr GO. 2015 The geological record of parasitic nematode evolution. In Advances in parasitology, vol. 80 (eds K De Baets, TJ Littlewood), pp. 53–92. New York, NY: Academic Press.
21. Thomas PK. 2002 Investigations for pneumonia. J. Assoc. Physicians India 60, 13–16.
22. Mitchell MA. 2012 Mycobacterial infections in reptiles. Veterinary Clin. North Am. Exot. Anim. Pract. 15, 101–111. (doi:10.1016/j.cvjp.2011.10.002)
23. Mitchell RI. Friends of the National Zoo. 1970 Proc. Symp. on Mycobacterial Infections of Zoo Animals, Washington, DC, USA, 6–8 October 1976. Washington, DC: Smithsonian Institution Press.
24. Cowan DF. 1968 Diseases of captive reptiles. J. Am. Vet. Med. Assoc. 153, 848–859.
25. Waltzek TB, Cortés-Hinjosa G, Wellehan Jr JFX, Gray GC. 2012 Marine mammal zoosones: a review of disease manifestations. Zoonoses Public Health 59, 521–535. (doi:10.1111/zph.12049.0.01202)
26. Brock JA, Nakamura MY, Miyahara HY, Chen EML. 2011 Tuberculosis in Pacific green sea turtles, Chelonia mydas. Trans. Am. Fish. Soc. 105, 546–566. (doi:10.1577/1548-8659(2010)03546:TITMPS>2.0.CO;2)
27. de Amorim DB et al. 2014 Mycobacterium pinnipedi in a stranded South American sea lion (Otaria byronia) in Brazil. J. Wildl. Dis. 50, 419–422. (doi:10.7589/2013-05-124)
28. Arriaza BT, Salo W, Aufderheide AC, Holcomb TA. 1995 Pre-Columbian tuberculosis in northern Chile: molecular and skeletal evidence. Am. J. Phys. Anthropol. 98, 37–45. (doi:10.2307/238014)
29. Herszkovitz et al. 2015 Tuberculosis origin: the Neolithic scenario. Tuberculosis 95, 512–516. (doi:10.1016/j.tube.2015.02.021)
30. Sager F, Scholzner M, Möller-Christensen V. 1972 A case of spondylitis tuberculosis in the Danish Neolithic Age. Dan. Med. Bull. 19, 176–180.
31. Zink AR, Grabner W, Reischl U, Wolf H, Nierich AG. 2003 Molecular study on human tuberculosis in three geographically distinct and time delineated populations from ancient Egypt. Epidemiol. Infect. 130, 221–239. (doi:10.1017/S0950268802009257)
32. Zimmerman MM. 1979 Pulmonary and osseous tuberculosis in an Egyptian mummy. Bull. N. Y. Acad. Med. 55, 604.
33. Rothschild BM, Martin LD, Lev G, Bercover H, Bar-Gal GK, Greenblatt C, Donoghue H, Spigelman M. 2001 The archaeology of western New York State. In Archaeology of western New York State (ed. RS Laub),
pp. 171–175. Buffalo, NY: Bulletin of the Buffalo Society of Natural Sciences.

35. Rothschild BM. 2005 Series, experience, limited experience and semantics of proboscidean forefoot and hindfoot erosions: distinguishing mycobacterial infection and spondyloarthropathy. J. Vertebrae Paleontol. 25, 1064–1074.

36. Hershkovitz I et al. 2008 Detection and molecular characterization of 9000-year-old Mycobacterium tuberculosis from a Neolithic settlement in the eastern Mediterranean. PLoS ONE 3, e3426. (doi:10.1371/journal.pone.0003426)

37. Lee OY-C et al. 2012 Mycobacterium tuberculosis complex lipid virulence factors preserved in the 17 000-year-old skeleton of an extinct bison, Bison antiquus. PLoS ONE 7, e41923. (doi:10.1371/journal.pone.0041923)

38. Lee OY-C, Wu HHT, Besa GS, Rothschild BM, Spigielman MA, Hershkovitz I, Bar-Gal GK, Donoghue HD, Minnin DE. 2015 Lipid biomarkers provide evolutionary synapomorphs for the oldest known cases of tuberculosis. Tuberculosis 95, 5127–5132. (doi:10.1016/j.tube.2015.02.013)

39. Resnick DL. 2002 Diagnosis of bone and joint disorders, 4th edn. Philadelphia, PA: Saunders.

40. Volz W. 1902 Proneusticosaurus, ein neues Sauropotyrgier-Gattung aus dem Unteren Muschelkalk Oberschlesiens. Palaeontographica 49, 121–162.

41. Rieppel O, Hagdorn H. 1997 Paleobiogeography of Middle Triassic Sauropterygia in central and western Europe. In Ancient marine reptiles (eds. JM Callaway, EL Nicholls), pp. 121–144. San Diego, CA: Academic Press.

42. Sues H-D. 1987 Postcranial skeleton of Pistosaurus and interrelationships of the Sauropterygia (Diapsida). Zool. J. Linn. Soc. 90, 105–111. (doi:10.1111/1096-3642.1987.tb01351.x)

43. Voeten DFA, Sander PM, Klein N. 2015 Skeletal material from larger Eusauropotheria (Reptilia: Eosauropterygia) with nothosaurian and cymatosaurian affinities from the Lower Muschelkalk of Winterswijk, The Netherlands. Palaeontol. Z. 89, 943–960. (doi:10.1515/palz.2015.024)

44. Sander PM. 1989 The pachypleurosaurids (Reptilia: Nothosauria) from the middle Triassic of Monte San Giorgio (Switzerland) with the description of a new species. Phil. Trans. R. Soc. Lond. B 325, 561–666. (doi:10.1098/rstb.1989.0103)

45. Anson C, Rothschild BM, Naples V. 2012 Soft tissue contributions to pseudopathology of ribs. Adv. Anat. Morphol. 2, 57–63. (doi:10.4236/aam.2012.2.20007)

46. Naples VL, Rothschild BM. 2011 Do ribs actually have a bare area? A new analysis. J. Comp. Hum. Biol. 62, 368–373. (doi:10.1016/j.jchb.2011.08.001)

47. Lyman RL. 1994 Vertebrate taphonomy. Cambridge, UK: Cambridge University Press.

48. Bennett JE. 1995 Mycobacs. In Mandell, Douglas and Bennett’s principles and practice of infectious diseases (eds JE Bennett, RJ Dolin, ML Blaser), pp. 2288–2289. New York, NY: Churchill Livingstone.

49. Rothschild BM, Rothschild C. 1998 Recognition of hypertrophic osteoarthropathy in skeletal remains. J. Rheumatol. 25, 2221–2227.

50. Rothschild BM, Rothschild C. 1995 Comparison of radiologic and gross examination for detection of cancer in deformed skeletons. Am. J. Phys. Anthropol. 96, 357–363. (doi:10.1002/aja.1330960404)

51. Meuten DJ. 2016 Tumors in domestic animals. New York, NY: John Wiley & Sons.

52. Eshed V, Latimer M, Greenwald CM, Jelloma LM, Rothschild BM, Weirh Baratuz S, Hershkovitz I. 2002buton osteoma: its etiology and pathophysiology. Am. J. Phys. Anthropol. 118, 217–230. (doi:10.1002/aja.10087)

53. Asmar S, Sassi M, Phelippeau M, Drancourt M. 2016 Mycobacteriosis-associated mortality in wild bison, Bison antiquus, from a Neolithic settlement in the Giorgio (Switzerland) with the description of a new species. PLoS ONE 11, e0154948. (doi:10.1371/journal.pone.0154948)

54. Klein N. 2010 Long bone histology of sauropterygia from the lower muschelkalk of the Germanic basin provides unexpected implications for phylogeny. PLoS ONE 5, e1613. (doi:10.1371/journal.pone.001613)

55. Krah A, Klein N, Sander PM. 2013 Evolutionary implications of the divergent long bone histologies of Nothosaurus and Pistosaurus (Sauropterygia, Triassic), BMC Evol. Biol. 13, 1. (doi:10.1186/1471-2148-13-32)

56. Klein N, Griebeler EM. 2016 Bone histology, microanatomy, and growth of the nothosaur Simosaurus galliardoi (Sauropterygia) from the Upper Muschelkalk of southern Germany/Baden-Württemberg, C.R. Palaevol 15, 142–162. (doi:10.1002/cvr.2015.02.009)

57. Houssaye A. 2009 ‘Pachyostosis’ in aquatic amniotes: a review. Integr. Zool. 4, 325–340. (doi:10.1111/j.1749-4877.2009.00146.x)

58. de Ricqlès A, de Buffrénil V. 2001 Bone histology, heterochronies, and control of the return of tetrapods to life in water: where are we? In Secondary adaptation of tetrapods to life in water (eds J-M Mazin, V de Buffrénil), pp. 289–310. Munich, Germany: Friedrich Pfeil.

59. Snyder GK. 1983 Respiratory adaptations in diving mammals. Respir. Physiol. 54, 269–294. (doi:10.1016/0034-5687(83)90072-5)

60. Simson DG, Kuschnier M, Clementson J. 1963 Respiratory function in pulmonary tuberculosis. Am. Rev. Respir. Dis. 87, 1–16.

61. Surmik D, Rothschild BM, Pawlicki R. 2017 Unusual intraspecies fossilized soft tissues from the Middle Triassic Nothosaurus bone. Sci. Nat. 104, 25. (doi:10.1007/s12047-017-1453-y)