Introduction

Osteomyelitis (OM) is a nonspecific infection of the bone and marrow. It is usually caused by pyogenic bacteria, most notably Staphylococcus, but other pathogens such as fungi and viruses can also be responsible (Aufderheide and Rodríguez-Martín, 1998; Ortner, 2003; Lewis, 2018; Roberts, 2019). Infection occurs either directly, through an open wound or adjacent soft tissue contamination or, more commonly, through indirect hematogenous dissemination. Because OM specifically involves the marrow, non-adult individuals—whose bones contain high quantities of red marrow—are particularly susceptible (Lewis, 2018). The condition is documented most frequently among children aged 3–12 years, typically manifesting as an acute (<4 weeks) infection (Aufderheide and Rodriguez-Martin, 1998; Ortner, 2003; Lewis, 2018; Roberts, 2019). When it spreads hematogenously, as is common among non-adults, then subsequent sepsis may be fatal. In fact, mortality estimates from the pre-antibiotic era were as high as 20–30% (Bancroft, 1921; Bass, 1928; Amberg and Ghormley, 1934; Lacey and Engel, 1939; Aufderheide and Rodríguez-Martin, 1998).

OM is a disease with diverse clinical and skeletal manifestations that reflect the combined effects of pathogen virulence, host resistance, and therapeutic treatment. Its study has been plagued by inconsistencies in terminology and diagnostic criteria, hindering both clinical and paleopathological research (Dormans and Drummond, 1994; Aufderheide and Rodriguez-Martin, 1998; Ortner, 2003; Suei et al., 2005; Baltensperger and Eyrich, 2009a, b; Chang et al., 2015; Santos and Suby, 2015; Lewis, 2018; Roberts, 2019). In paleopathology, in particular, pathognomonic lesions such as sequestra, involucra, and cloacae are not always present on skeletal remains. As a result, few archaeological cases of OM have been published, despite the general understanding of the condition as a relatively common and serious health complication in the past (Baltensperger and Eyrich, 2009b;
Santos and Suby, 2015; Tavares et al., 2017; Lewis, 2018; Roberts, 2019). A thorough documentation of OM prevalence, presentation, and distribution in archaeological and early historical contexts that pre-date the use of antibiotic therapies is paramount to understanding the evolution and long-term impact of the disease (Santos and Suby, 2015). This paper contributes to the paucity of paleopathological literature by presenting a probable case of chronic mandibular OM in a young child from Moyoro, an Okhotsk period site in northern Japan. A differential diagnosis and discussion shed light on the condition’s etiology and pathogenesis. To the best of our knowledge, this is only the second case of pediatric mandibular OM documented thus far from the archaeological record.

Materials and Methods
The Okhotsk were marine-adapted foragers inhabiting Sakhalin, Hokkaido, and the Kuril Islands during the 5th–13th centuries AD (Amano, 2003; Hudson, 2004; Naito et al., 2010; Tsutaya et al., 2015; Junno et al., 2021). Their early occupation (5th–6th century) of Sakhalin and north Hokkaido was followed by rapid eastward expansion into east Hokkaido and the Kuril Islands during middle (7th–9th century) period (Yamaura and Ushiro, 1999; Hudson, 2004; Tsutaya et al., 2015). While the Okhotsk relied largely on hunting and fishing, particularly of marine resources, there is also evidence that they consumed domesticated foods including dogs, pigs, barely, and millet (Ishida, 1988; Amano, 2003; Tsutaya et al., 2014; Liepe et al., 2017). However, their eastward migration necessitated a subsistence shift in response to changing winter sea ice conditions: the east Hokkaido Okhotsk did not keep pigs and relied heavily on marine mammals (Yamaura, 1998; Yamaura and Ushiro, 1999; Hudson, 2004; Tsutaya et al., 2014). After remaining distinct for centuries, the Okhotsk were replaced by/assimilated with Satsumon groups, who had Jomon ancestry, at the end of the 10th century AD in north Hokkaido and during the 12th century AD in east Hokkaido (Amano, 2003; Hudson, 2004; Sato et al., 2007, 2009; Tsutaya et al., 2015). The site of Moyoro (Figure 1) is an east Hokkaido Okhotsk shell midden from which ~250 individuals were excavated between 1926 and 2011 (Kodama, 1948; Komai, 1964; Kiyono, 1969; Abashiri City Board of Education, 2014). Most human remains recovered date between 500 and 900 cal. AD (with application of the Marine13 calibration curve), consistent with the middle Okhotsk period of rapid eastward expansion (Tsutaya et al., 2014, 2015). While several rich Moyoro burials have been noted (Hudson, 2004), there is no information regarding mortuary context for many interments, including the individual discussed here.

Moyoro 165, the subject of this paper, represents the incomplete skeleton (Figure 2) of a young child. The remains are directly dated to 1871 ± 32 BP, or 736–977 cal. AD (68.3%, ±1 SD), with the application of the IntCal20 and Marine20 calibration curves and assuming 90% contribution from marine proteins and a local reservoir correction of +248 ± 99 years (based on eight dates from Kunashiri, Shikotan, and Sakhalin; Kuzmin et al., 2001; Yoneda et al., 2007), as calculated with the 14CHRONE Marine20 Reser-
voir Database. The lesion of interest affects the mandible, which is represented by two fragments: (1) most of the left mandibular body and a small part of the right, and (2) the posterior aspect of the right mandibular body and ramus (Figure 3). An inventory and developmental summary of the mixed mandibular dentition are presented in Table 1, indicating a dental age of 3.5–5.5 years (Ubelaker, 1979; Schaefer et al., 2009; AlQahtani et al., 2010). Of particular note—and directly relevant to the lesion (see below)—is the antemortem loss of both left deciduous molars and the developing replacement permanent premolars. The cranial and postcranial remains are fragmented and incomplete, with occipital and vertebral fusion indicating an age consistent with dental development (Schaefer et al., 2009). Slight porosity was observed on the right orbital roof, and two linear enamel hypoplastic defects were noted on the cervical crowns of each developing permanent mandibular canine (Figure S1), suggesting stress events during the 12–18 months before death (Schaefer et al., 2009; AlQahtani et al., 2010). No other pathological changes were documented, although skeletal preservation is admittedly poor.

**Results**

The lesion itself encompasses most of the left mandibular body and exhibits two main features (Figure 3). The first is cortical thickening along the length of the body, most pronounced between the canine and first permanent molar, reflecting periosteal new bone formation on the lateral (buccal), inferior, and medial (lingual) surfaces. The cortex in this area, especially on the lateral surface, is distinctly swol-
len and slightly roughened, with increasing porosity toward the superolateral margin (Figure 3b, c). The left mandibular body measures 21.8 mm mediolaterally at its greatest extent (mesial to the first permanent molar), while the right measures 12.5 mm at the same location. The second lesion feature is a large (33.2 mm × 21.4 mm) lytic defect of the left alveolar bone between the canine and first permanent molar (Figure 3a). This lesion completely obliterated the sockets of both deciduous molars, the crypts of both developing permanent premolars, and most of the mandibular canal anterior (mesial) to the first permanent molar. The lateral and medial margins of the defect are elevated (superiorly) relative to its interior and are distinctly porous, particularly on the lateral side (Figure 3c). The entire base of the lytic lesion consists of undulating trabecular bone, with the course of the mandibular canal being vaguely distinguishable (Figure 3a). The base of the defect was further examined via a handheld digital microscope (Figure 4).

Clinical computed tomography (CT) scanning was conducted with a NAOMi-CT 3D-M (RF Co.) at the Sapporo Medical University by H. Matsumura. The images were reconstructed with isotropic 0.071-mm voxels. CT imaging confirms that the expansion of the left mandibular body reflects periosteal new bone formation on the lateral, inferior,
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and medial surfaces (Figure 5, Figure 6). Much of this bone is at least partially remodeled and integrated with the original cortex, but areas of lamellated (‘onion-skin’) deposits are visible on the lateral mandibular body (Figure 5a–f, Figure 6a, c–d). The lytic alveolar lesion appears to interrupt the periosteal new bone in the area of the deciduous molars and permanent premolars (Figure 5c–e, Figure 6c). There is no evidence of these teeth or their dental crypts, but the crowns and partial roots of the developing left permanent incisors are visible in the anterior portion of the mandible (Figure 5g–h). The mandibular canal appears to be intact posterior (distal) to the alveolar defect (Figure 5b, Figure 6a, c).

The CT scans also reveal (1) dense trabecular bone in the interior mandibular body, especially inferior to the alveolar lesion (Figure 5a–c, Figure 6a, b); (2) undulating trabeculae comprising the base and margins of the lytic lesion (Figure 5e–f, Figure 6b–d); and (3) numerous small canals extending outward from the internal mandibular body, particularly from the area of the lytic lesion (Figure 5b–f, Figure 6).

Most of these canals, especially on the more inferior portion of the bone, do not perforate the external cortex. Those that do appear to correlate with the porosity observable on the lateral surface (Figure 3b, c). No evidence of trauma (e.g., fractures) or abscess formation (e.g., cloacae) was observed on the mandible macroscopically, microscopically, or via CT imaging. No destructive analyses were permitted for these remains.

Differential Diagnosis

The two distinct lesion features on the Moyoro 165 mandible, one osteoblastic and the other osteolytic, complicate differential diagnosis somewhat. Pronounced periosteal new bone formation on the mandibular body may reflect a number of different pathological conditions in a child, including hypervitaminosis A, infantile cortical hyperostosis, malignant neoplasia, treponemal disease, fibro-osseous lesions, Langerhans cell histiocytosis, and osteomyelitis (Belli et al., 2002; Fukuda et al., 2017; Lewis, 2018). Lytic alveolar destruction, on the other hand, can result from some of the same conditions—malignant neoplasia, Langerhans cell histiocytosis, and osteomyelitis—as well as tuberculosis (Can et al., 2005; Baltensperger and Eyrich, 2009a; Lewis, 2018; Hwang et al., 2019).

Hypervitaminosis A, or vitamin A toxicity, reduces bone formation and enhances its resorption (Binkley and Kreuger, 2000). The condition has been associated with a proliferation of coarse periosteal bone on the skull and long bones and, in non-adults, delayed ossification (Naz and Edwards, 1952; Woodard et al., 1961; Walker et al., 1982; Außerheide and Rodriguez-Martin, 1998). In the case of the Moyoro 165, bone formation appears to be limited to the mandible and is distinct in its lamellated appearance. This, and the singular lytic alveolar defect, is inconsistent with hypervitaminosis A. Infantile cortical hyperostosis (ICH) has a poorly understood etiology that may include viral infection, trauma, arterial morphology, and even hypervitaminosis A (Lewis, 2018). It is characterized by widespread formation of peri-
osteoal new bone and cortical thickening on multiple elements, most often the skull, long bones, and ribs (Kamoun-Goldrat and le Merrer, 2008; Lewis, 2018). On the mandible, the condition is more common on the ramus and angle, rather than the body (Karjodkar, 2009). The diffuse lesion distribution of ICH and its occurrence on young infants, typically manifesting before five months of age (Caffey, 1945; Kamoun-Goldrat and le Merrer, 2008), make it an unlikely diagnosis in this case.

Malignant neoplastic conditions affecting young children, such as leukemia, Ewing’s sarcoma, and metastatic neuroblastoma, may also account for the periosteal bone forming lesion and/or lytic alveolar defect on the Moyoro 165 mandible. Leukemia, acute leukemia in particular, is the most common malignant condition affecting children (Kobayashi et al., 2005), with bony involvement present in at least 50% of cases (Sugihara et al., 1989). Skeletal changes are typically multifocal and characterized by widespread periosteal bone deposition and osteolysis. In the jaws, lytic lesions can result in trabecular loss, thinning dental crypts, and tooth displacement (Sugihara et al., 1989; Kobayashi et al., 2005; Lewis, 2018). In this case, the localized and partially lamellated periosteal bone formation and single alveolar defect are inconsistent with leukemia. Furthermore, some of the most characteristic features of childhood leukemia, enlarged vascular foramina and exaggerated porosity and sulci on the metaphyseal cortices (Grauer, 2019), are absent on this individual. Ewing’s sarcoma and neuroblastoma are two other childhood malignancies that also cause osteoblastic lesions, including ‘onion skin’, sunburst, or hair-on-end reactions. However, these are typically aggressive bony expansions accompanied by osteolysis (e.g., mottling) visible on radiographs (Campanacci et al., 1990; Belli et al., 2002; White and Pharoah, 2009; Akgül et al., 2018; Botía González et al., 2018; Lewis, 2018), which are absent here. The localized and partially remodeled bone formation on Moyoro 165 does not support a malignant diagnosis.

Fibro-osseous lesions that may account for mandibular expansion in children include fibrous dysplasia and cherubism. Both are benign conditions reflecting an overproduction of fibrous tissue within bone (Papadaki et al., 2012; Lewis, 2018; Marques, 2019). Fibrous dysplasia can affect one or several bones, the latter typically asymmetrically, while cherubism is usually symmetrical and limited to the mandible and maxillae. Radiographically, both conditions are distinguishable by their ‘ground glass’ appearance (Kambadakone et al., 2008; Craig and Craig, 2013; Willmon et al., 2013; Lewis, 2018; Marques, 2019), which is absent in this case (Figure 5, Figure 6). Fibro-osseous lesions are therefore inconsistent with the osteoblastic lesion on Moyoro 165.

Infectious diseases—specifically treponematosis (one of four conditions caused by Treponema pallidum) and tuberculosis (caused by Mycobacterium, usually M. tuberculosis)—are also possible diagnoses. Treponemal infections potentially affecting the jaws of young children are largely limited to congenital syphilis (Roberts and Buikstra, 2019). Associated skeletal changes include profuse periosteal bone formation, gummatous lesions on the skull, and ‘saddle nose’ facial deformities. Pathognomonic dental changes, such as hypoplastic teeth, mulberry molars, and Hutchinson’s incisors, are most common, occurring in up to 45% of infected individuals (Lewis, 2018; Roberts and Buikstra, 2019).
In the case of Moyoro 165, the solitary and localized nature of the bone deposition and the typical morphology of all ten (nine mandibular and one maxillary) observable teeth make treponematosis unlikely. Tuberculosis (TB), while rarely affecting the mandible, does so more commonly in infants and children than in adults (Aufderheide and Rodriguez-Martin, 1998; Lewis, 2018; Roberts and Buikstra, 2019). TB lesions on non-adult skeletal remains are usually multifocal, rather than solitary, and predominately osteolytic. On some elements, such as the mandible, lesions may mimic those of Langerhans cell histiocytosis (Lewis, 2018; Roberts and Buikstra, 2019), making differential diagnoses difficult. In this case, the solitary nature of the lytic alveolar lesion, lack of destructive foci elsewhere on the skeleton, and apparent absence of the disease from Hokkaido until the mid-1600s (Kondo and Aono, 2016) all argue against a diagnosis of TB.

Langerhans cell histiocytosis (LCH) is a reticuloendothelial disease in which an overproduction of histocytes called Langerhans cells causes destructive granulomatous lesions in skeletal tissues and other organs (Willman et al., 1994; Lewis, 2018; Grauer, 2019). Its peak occurrence is among children under the age of 10, but its presentation can vary widely depending on distribution, severity of skeletal involvement, and age of onset (Lewis, 2018; Grauer, 2019). In the paleopathological literature, LCH is generally classified into one of three diseases (Grauer, 2019). Eosinophilic granuloma is the most common and least severe form of LCH, characterized by solitary lytic lesions that often involve the cranial vault or facial skeleton. In the mandible, where the body or angle is typically affected, alveolar destruction can cause radiographic ‘floating’ teeth, tooth loss, and disrupted dental development (Watzke et al., 2000; Can et al., 2005; Mitomi et al., 2005; Lewis, 2018; Hwang et al., 2019; Grauer, 2019). Hand–Schüller–Christian Disease and Letterer–Siwe Disease are more severe versions of LCH, together comprising about 30% of cases. They are characterized by multifocal, coalesced lesions and diffuse skeletal and multi-organ involvement, respectively (Lewis, 2018; Grauer, 2019), none of which appear to be present in this case. LCH lesions can also have sclerotic borders and/or be associated with periosteal new bone formation (Grauer, 2019). However, these are uncommon in children with eosinophilic granuloma and, when present, are usually discontinuous (Yu et al., 1995). The solitary alveolar defect and tooth/crypt loss of Moyoro 165 is consistent with eosinophilic granuloma and, while the adjacent periosteal new bone formation is atypical for the condition, it is not inconsistent. Therefore, LCH, specifically eosinophilic granuloma, is a possible diagnosis for this case.

OM, a bone and marrow infection most commonly affecting children, can be difficult to diagnose from archaeological human remains (Tavares et al., 2017; Lewis, 2018; Roberts, 2019). The condition may present with diverse and even nonspecific skeletal lesions reflecting variation in its manifestation (e.g., acute/subacute versus chronic, or supplicative versus non-suppurative) or infection source (Aufderheide and Rodriguez-Martin, 1998; Ortner, 2003; Suei et al., 2005; Baltensperger and Eyrich, 2009a, b; Lewis, 2018; Roberts, 2019). For example, OM stemming from open wounds or adjacent soft tissue contamination is often localized and restricted to the cortex and periosteum, without clear involvement of the medullary cavity (Ortner, 2003; Roberts, 2019). In fact, characteristic lesions reflecting tissue necrosis (sequestra), periosteal elevation (involucra), and pus drainage (cloaca) are frequently absent, even in cases of acute supplicative infection. In children, this can reflect the ability of exudate to seep through porous immature bone without increasing vascular pressure (Lewis, 2018). Pediatric OM is usually hematogenous in origin, most often initiating in the metaphysis of a single long bone, especially the tibia or femur; it is rare in the skull and jaws (Aufderheide and Rodriguez-Martin, 1998; Ortner, 2003; Lewis, 2018; Roberts, 2019).

OM of the jaws is a unique condition that continues to present diagnostic challenges, even for modern clinicians (Suei et al., 2005; Baltensperger and Eyrich, 2009b). Like OM elsewhere in the body, hematogenous spread is a known route of infection, especially in the absence of modern therapeutic treatment. However, direct extension from an odontogenic focus is frequently implicated and today accounts for most cases of jaw OM. Among infants and young children, such as Moyoro 165, developing tooth germs represent an additional source of (hematogenous or direct mucosal) infection (Ortner, 2003; Baltensperger and Eyrich, 2009a; Roberts, 2019). Because of its architectural similarities to long bones (i.e., thick cortex and abundant medullary tissue) and more restricted blood supply, the mandible is ten times as likely to develop OM than is the maxilla in individuals over the age of two years (Baranoff, 1934; Ortner, 2003; Baltensperger and Eyrich, 2009a; Roberts, 2019). The mandibular body is most often affected, followed by the symphysis and angle. While mandibular OM can present with pathognomonic lesions, it may also be clinically silent and/or non-suppurative, characterized by localized nonspecific lesions such as diffuse sclerosis, cortical thickening, and little or no osteolysis (Aufderheide and Rodriguez-Martin, 1998; Ortner, 2003; Baltensperger and Eyrich, 2009a; Lewis, 2018; Roberts, 2019). Furthermore, OM stemming from tooth germ infection can result in the necrosis and exudation of dental tissues, with subsequent tooth loss and developmental facial deformities (Natvig and Dingman, 1957; Ortner, 2003; Baltensperger and Eyrich, 2009a; Roberts, 2019). Both lesion features of the Moyoro 165 mandible are consistent with a diagnosis of OM, making OM a more likely diagnosis than LCH in this case.

Discussion

In paleopathology, OM diagnosis is limited by lesion specificity, with etiology generally focusing on the route of infection rather than pathogen identification. Unfortunately, establishing the cause of OM from archaeological human remains is not usually possible, except in rare cases of compound fractures (Santos and Suby, 2015; Lewis, 2018). OM of the jaws is distinct in this regard, as its etiology is most often linked to odontogenic sources (Oulis et al., 2000; Belli et al., 2002; Ortner, 2003; Suei et al., 2005; Baltensperger and Eyrich, 2009a; Khudaverdyan, 2011; Waters-Rist, 2012; Chang et al., 2015; Fukuda et al., 2017; Roberts, 2019).
Direct extension from an odontogenic infection is implicated for Moyoro 165 by the clear evidence of alveolar involvement and tooth loss, and by the localized nature of the lesion. Because developing and erupting teeth are particularly susceptible to infection (Huang, 2009), the Moyoro child’s young age is compatible with OM development secondary to an infected tooth germ (e.g., tooth 34 and/or tooth 35). However, a periapical infection from caries or direct trauma, or a periodontal infection associated with a recently erupted tooth (e.g., tooth 74 and/or 75), are also possible etiological mechanisms (Oulis et al., 2000; Belli et al., 2002; Chang et al., 2015; Fukuda et al., 2017).

The pathogenesis of OM reflects the complex interplay between infection virulence and the effectiveness of the host’s immune response (Aufderheide and Rodriguez-Martin, 1998; Baltensperger and Eyrich, 2009a). Acute OM (<4 weeks) that is eradicated quickly indicates a strong response relative to pathogen number and/or virulence, whereas acute OM that is fatal, occurring in 20–30% of pre-antibiotic era cases (Bancroft, 1921; Bass, 1928; Amberg and Ghormley, 1934; Lacey and Engel, 1939; Aufderheide and Rodriguez-Martin 1998), suggests the opposite. Chronic OM (4+ weeks) usually reflects the host’s failure to resolve the acute stage of the disease, with the infection invading deep into the cortical bone and medullary cavity. Its manifestation, especially in the jaws, can be quite varied (e.g., Oulis et al., 2000; Belli et al., 2002; Ortner, 2003; Suei et al., 2005; Baltensperger and Eyrich, 2009a; Khudaverdyan, 2011; Waters-Rist, 2012; Chang et al., 2015; Fukuda et al., 2017). In the case of Moyoro 165, the chronic nature of the condition is demonstrated by lamellated periosteal bone formation, partial remodeling, and dense trabeculae comprising much of the mandible’s interior. Furthermore, the infection appears to have been active at the time of death, as the porous and slightly roughened cortical bone is inconsistent with a partially or completely healed lesion. Thus, infection likely persisted for at least several months, and possibly longer than a year, prior to death. While local immunological and microbial factors would have played a considerable role in the condition’s pathogenesis (Baltensperger and Eyrich, 2009a), at this point it is not possible to link these to aspects of Okhotsk cultural practice or diet. Skeletal lesions pathognomonic of OM reflect the combined effects of vascular obstruction (tissue necrosis) and pus formation (periosteal elevation and drainage cloacae). The anatomical structure of the mandible makes it particularly vulnerable to infection, and thus these changes. For example, most of the blood supply to the mandibular body is delivered via the small inferior alveolar artery, passing through the mandibular canal (Castelli, 1963; Baltensperger and Eyrich, 2009a). The artery is easily damaged, such as through acute inflammation, and subsequent vascular disruption is critical in the establishment of mandibular OM. In addition, the canal acts as a corridor for the rapid dissemination of pus through the mandibular body, which further compresses the neurovascular tissues contained within it (Baltensperger and Eyrich, 2009a). Thus, for Moyoro 165, the inferior alveolar artery and mandibular canal may have contributed to and exacerbated the infection. Other than bone loss and evidence of dental tissue necrosis, no other characteristic signs of OM are evident on the mandible. However, given the young age of this individual, the unique manifestation of OM in the jaws, and the chronic nature of the condition, this is not unexpected (Aufderheide and Rodriguez-Martín, 1998; Ortner, 2003; Baltensperger and Eyrich, 2009a; Lewis, 2018; Roberts, 2019).

Chronic OM can persist for months or years. In the jaws, clinical symptoms such as localized swelling, pain, and/or numbness are usually less pronounced than those of the acute phase of the disease (Oulis et al., 2000; Belli et al., 2002; Baltensperger and Eyrich, 2009a; Chang et al., 2015; Fukuda et al., 2017). In some cases, patients can have no distress and even retain normal function such as mastication. The lytic alveolar lesion, mandibular canal destruction, and dental tissue loss on Moyoro 165 suggest that the infection was not clinically silent and had at least some impact on jaw function, if only temporary. However, the long-term survival of the child for months or even years after the onset of infection, and stable carbon and nitrogen isotope values consistent with other Moyoro individuals (Tsutaya et al., 2015: ID 1072 is Moyoro 165), imply adequate nutrition and care by family and community members even if masticatory function was impaired in some way. Chronic OM may have contributed to the death of the child, but other conditions cannot be ruled out as potential causes or contributing factors, especially considering the generally poor state of skeletal preservation. Indeed, the two linear enamel hypoplastic defects noted on the permanent mandibular canines (Figure S1) indicate physiological stress of some sort within the preceding 12–18 months.

While OM is considered to have been a relatively common condition in the past, especially among non-adult individuals (Baltensperger and Eyrich, 2009b; Santos and Suby, 2015; Roberts, 2019), it is rarely reported in the paleopathological literature (Tavares et al., 2017; Lewis, 2018; Roberts, 2019). Tavares and colleagues (2017) identified only ten published cases of pediatric OM, and a handful of others (e.g., Anderson and Carter, 1995; Ortner, 2003; Waters-Rist, 2012; Lewis, 2018) can be added to this list. OM of the jaws, in particular, has been documented even less frequently from archaeological human remains. Despite several reported infections being possibly attributable to jaw OM and/or odontogenic sources (Wood-Jones, 1910; Hooton, 1930; Roney, 1966; Khudaverdyan, 2011), we have identified only two other paleopathological cases involving non-adult individuals. In the first, Ortner (2003: 197–198) identified possible hematogenous spread of OM from a periapical maxillary abscess to the postcranial skeleton in a six-year-old child from 16th century AD Virginia (USA). In the second, Waters-Rist (2012) documented mandibular OM, likely stemming from an infected deciduous canine germ, in a 9–to 18-month-old infant from Early Neolithic (~7000 cal. BP) Siberia. Lesion appearance in the latter case was considerably different from that presented here: there was evidence of suppuration and new periosteal bone formation, but no osteolytic defects or tooth loss. Thus, Moyoro 165 represents a unique manifestation of mandibular OM and, to the best of our knowledge, only the second pediatric example reported from an archaeological context.
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Conclusion

Moyoro 165 presents an unusual mandibular lesion with two distinct features, periosteal new bone formation and lytic alveolar destruction, that are most consistent with a diagnosis of OM. Direct extension from an odontogenic source, likely an infected tooth germ, is implicated by the lesion’s location and its alveolar and dental involvement. Further lesion features, such as lamellated periosteal bone deposition and partial remodeling, suggest a chronic stage of infection over a period of months or even years that was still active at the time of death. Dental evidence of physiological stress during the previous 12–18 months is consistent with this interpretation. Of the fewer than 20 cases of pediatric OM reported in the paleopathological literature, Moyoro 165 represents only the second involving the mandible, exhibiting features considerably different from those observed in the other case. The unique lesion appearance documented here highlights the diverse manifestation of OM, especially in the jaws and in the absence of modern therapeutic treatment. More research on OM during the pre-antibiotic era—including its variable prevalence, presentation, and distribution—is essential to more fully understand the condition’s evolution and long-term impact on human populations.

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