Differential diagnosis of a probable case of non-adult thalassaemia from 4th century AD Romano-British Colchester, UK


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Differential diagnosis of a probable case of non-adult thalassaemia from 4th century AD
Romano-British Colchester, UK

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Abstract

Our current understanding of immigration and diasporic disease in Roman Britain has been greatly enhanced by the recent identification of thalassaemia in the non-adult skeletal record. The wide phenotypic variation in the clinical expression of β-thalassaemia, however, means that additional cases may go unrecognised. A probable diagnosis for β-thalassaemia intermedia or a mild form of major in a 1.0-1.5 year old skeleton from Butt Road, Colchester, dating to the 4th century AD is discussed here. The assessment was undertaken using macroscopic and radiographic analysis. Several conditions were apparent, including trauma and probable β-thalassaemia and active vitamin D deficiency. Diagnosis proved difficult due to the challenges that non-adult thalassaemia poses for identification in the skeletal record, as in the absence of the cranium only ‘rib-within-a-rib’ is currently considered as pathognomonic of the condition. This case demonstrates the variations in expression of this type of genetic anaemia and adds emphasis to a more widespread presence of this important condition in Roman Britain.

Keywords

Anaemia – migration – rickets – vitamin D deficiency – fracture
1. Introduction

Thalassaemia has been identified in the Romano-British non-adult skeletal record, however diagnoses remain challenging due to the varied expression of the disease and the low number of published cases (Taher et al. 2006; Lewis 2012). Diagnoses have been hindered by the lack of pathognomonic features and the non-specific nature of cribra orbitalia and porotic hyperostosis (Ortner 2003, 364-365). To date, the only cases of probable non-adult β-thalassaemia in Roman Britain have been recovered from 3rd-5th century AD Poundbury Camp, Dorchester, Dorset (Lewis 2012) (Fig. 1). Since the disease is not endemic to Britain, but the Mediterranean, these cases provide direct osteological evidence for immigration to Roman England.

The alterations of the haemoglobin molecules and red blood cells in β-thalassaemia give the carrier an increased immunity to malaria (Plasmodium vivax and Plasmodium falciparum) (Hershkovitz et al. 1991, 1997; O’Donnell et al. 2009). This type of genetic anaemia is therefore a result of genetic polymorphisms perpetuated by a resistance to malaria. Heterozygous individuals display partial immunity, whereas homozygous carriers are fully immune but suffer from β-thalassaemia intermedia or major (Mitchell 2003; Ustundag 2011), with β-globin protein production partially suppressed due to defective gene function (Taher et al. 2006) (Table 1). Costal osteomas and ‘rib-within-a-rib’ lesions have been noted as suggestive of β-thalassaemia intermedia (Lawson et al. 1981; Tunaci et al. 1999).

Subsequently, Lewis (2012) demonstrated that changes in the thorax, and lesions attributed to anaemia are likely to be skeletal manifestations of non-adult β-thalassaemia. This case study aims to highlight the challenges of a differential diagnosis in incomplete non-adult remains and showcases the range of expressions observable in probable cases of non-adult β-thalassaemia.

2. Materials and Methods

The colonia of Vitricensis Camulodunensium was founded on the oppidum of Camulodunum and is now contained within modern day Colchester, Essex (Wacher 1974, 104; Brooks 2006; Black 2006) (Fig. 1). The burial ground at Butt Road was excavated from 1976-1979, and during 1986-1988 by Colchester Archaeological Trust (Crummy and Crossan 1993, 4). The cemetery is situated outside of the walled town by the main south-west gate. The phases of
the cemetery dating to the 4th-5th centuries hold the remains of 109 non-adults aged 0-17 years old (15.6%), with a total of 700 recovered individuals (Crummy and Crossan 1993, 62; Pinter-Bellows 1993).

G145 was recovered as a single unfurnished inhumation from the eastern edge of the cemetery (Crummy and Crossan 1993) (Fig. 2). The non-adult was aged to a mean dental age of 1.0-1.5 years, based on crown formation of the permanent first mandibular molar and root formation stages of the deciduous first and second mandibular molars (Moorrees et al. 1963a,b). G145 is represented by elements of the trunk, upper limbs and long bone fragments. The skull is absent apart from maxillary and sphenoid fragments (Fig. 3). The individual displayed noteworthy rib lesions and pathological changes to the upper limbs.

A range of conditions have to be considered for the nature of the lesions observed, including β-thalassaemia (Table 1). Diagnoses are based on macroscopic assessment and digital radiography undertaken at the University of Reading.

3. Results

Upper limbs show microporosity of the cortical bone, with enlarged trabecular structure and cortical thinning suggestive of osteopenia, the latter also apparent in the ilium (Figs. 4, 5). Thickening towards epiphyseal ends is evident in the long bones (Figs. 6, 7). The right radius appears thickened and angled at the distal third of the diaphysis with a transverse band of remodelled woven bone, indicative of a fracture callus (Figs. 7). Radiographically, the affected area presents as a fracture of the metaphysis which resulted in angulation of the distal portion and widespread new bone deposits (Fig. 8). Radiographs reveal a radiolucent band at the distal metaphysis of the left radius (Fig. 8).

Bilaterally, ribs appear flared and thickened at the costo-chondral end with increased porosity (Figs. 9-11). Several ribs display localised thickening of the shafts, reminiscent of ‘swelling’ with one expanded focus per element (Fig. 10, 11). These areas of hyper-marrow bone occur bilaterally, although primarily on the right (five right ribs, one left rib). Macroscopically, the expanded foci do not appear porous, or as healed sub-periosteal new bone formation, but rather as localised masses of dense compact bone sitting on top of the original cortical bone (Fig. 11). Radiographs indicate osteopenia, and reveal a radio-opaque appearance of the foci of lamellar bone deposited on the original cortex (Figs. 12, 13).
4. Discussion

Due to the incomplete preservation of the remains and the ambiguous nature of the lesions, several probable causes have to be considered (Table 1). Changes to rib ends, cortical microporosity throughout and a radiolucent band at the left radius strongly suggest active vitamin D deficiency at the time of death (Ortner and Mays 1998; Ortner 2003). It may be suggested that given the fact that this child was sickly, it may have been kept indoors for a prolonged period which exacerbated its vitamin D deficient status. Recent considerations of the disease have demonstrated that skeletal changes attributed to rickets/osteomalacia represent the most extreme end of the disease spectrum, often associated with a range of comorbidities (Snoddy et al. 2016).

Since G145 suffered from active rickets at the time of death, the child may have also been affected by respiratory disease (Pettifor and Daniels 1997; Pettifor 2003). A non-specific respiratory infection may cause visceral new bone formation which has to be discussed as a possible aetiology of the observed rib lesions, alongside rib fractures as a result of violent coughing (Lovell 1997). Proliferation in respiratory infections is most commonly apparent towards the vertebral end and often occurs bilaterally (Santos and Roberts 2001), the location of foci on the ribs of G145 is however inconsistent and isolated. In expansive visceral lesions, porous new bone blends into the existing shape of the rib shaft which is difficult to differentiate macroscopically. However, radiographs confirm a layer of compact bone on the original cortex rather than a blending of original and newly formed bone (Matos and Santos 2006; Lagia et al. 2007). Fracture lines are absent on radiographs of all ribs, and new cortical bone is deposited at the mid-shaft, primarily anteriorly and on top of the original cortex, appearing as localised masses rather than fracture callouses (Lewis 2012).

Although G145 does not display the extreme skeletal alterations described by Lagia et al. (2007) and Lewis (2012) for β-thalassaemia intermedia, there is wide phenotypic variation in clinical expression which may account for the less extreme skeletal changes observed (Weatherall and Clegg 1999). Although the pathognomonic ‘rib-within-a-rib’ is absent, the cortical swelling in the ribs conforms with a diagnosis for costal osteomas. The cortical porosity and swelling observed throughout the skeleton are apparent in β-thalassaemia intermedia and mild forms of major. Additionally, osteopenia has been observed in cases of non-adult β-thalassaemia (Lewis 2012). Yet, in the absence of cranial and facial bones, the characteristic ‘rodent face’ (Almeida and Roberts 2005; Tyler et al. 2006) and ‘hair-on-end’
as thalassaemic traits cannot be assessed for, urging for a consideration of sickle-cell anaemia in G145. The most common complication in sickle-cell disease is vaso-occlusive crisis. Blood vessels are congested by sickle-cell red blood cells, hindering circulation and leading to arthritic changes, dactylitis, or vertebral collapse (Almeida and Roberts 2005). Bone marrow infarction has not been observed which makes sickle-cell anaemia unlikely, although the incomplete skeleton of G145 does not allow for this type of genetic anaemia to be fully ruled out.

The trauma on the right distal radius possibly describes a torus fracture with buckling of the cortex, or a greenstick fracture with the cortex intact on the ulnar aspect, albeit only minimal (Ogden 2000, 617). Looser’s zone has to be considered as a pathological pseudo-fracture in the distal radius due to rickets (Aufderheide and Rodriguez-Martin 1998, 307; Keller and Barnes 2008). However, the fracture presents with angulation and displacement which refutes a pseudo-fracture (Keller and Barnes 2008). The radial fracture may be the result of loss in bone mineral density secondary to genetic anaemia, and probably exacerbated by active rickets (Michelson and Cohen 1988; Keller and Barnes 2008). Generally, children around 18 months old start walking, and given the angle of the fracture a fall may be a likely cause. However, we have to consider that G145 was limited in mobility and the fracture arose due to general weightbearing in crawling or handling.

5. Conclusion

The lesions apparent in this non-adult skeleton provide several diagnostic challenges. The missing cranium prevents from making a conclusive diagnosis of β-thalassaemia major or intermedia, although co-occurrence of thalassaemia and active rickets is likely. This case may be used to illustrate the difficulty in diagnosing the β-thalassaemias in non-adults, and the possibility for less severe lesions which may be easily overlooked during osteological analysis and recording. This case demonstrates the need to look for β-thalassaemia in the Romano-British non-adult skeletal record, and the importance of awareness of the varied expressions this particular anaemia may yield. Evidently, this case also demonstrates that the disease was not unique to Poundbury Camp, and provides direct skeletal evidence for migrants in Roman Colchester.
Acknowledgements

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6. Literature cited


Table 1. Genetic and clinical status of the β-thalassaemias

<table>
<thead>
<tr>
<th>Feature</th>
<th>β-thalassaemia minor</th>
<th>β-thalassaemia intermedia</th>
<th>β-thalassaemia major</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genomic pathology</td>
<td>Heterozygous: one β-globin gene with mutation (2, 3)</td>
<td>Mostly homozygous or rarely compound homozygous: two β-globin genes with mutation, one of which is mild; or one β-globin mutation combined with α-globin genes (1, 2)</td>
<td>Homozygous: two β-globin genes with severe mutation (2, 3, 5)</td>
</tr>
<tr>
<td>Clinical status</td>
<td>Anaemia mild or absent, very limited to no skeletal changes (2, 5)</td>
<td>Anaemia, transfusion-independent, skeletal involvement (2, 5)</td>
<td>Iron overload, severe anaemia, transfusion-dependent, skeletal involvement (2, 3, 5)</td>
</tr>
<tr>
<td>Phenotypic expression</td>
<td>Asymptomatic/silent (2, 3, 5)</td>
<td>On a spectrum from asymptomatic to severe, discernible in osteological record as gross skeletal changes but able to live into later life (2, 3, 4, 5)</td>
<td>Severe, lifelong reliance on blood transfusions and supportive care, probably not have survived past the age of two years in the past (2, 5)</td>
</tr>
</tbody>
</table>

From 1, Galanello and Cao (1998); 2, Rund and Rachmilewitz (2005); 3, Thein (2005); 4, Taher et al. (2006); 5, Lagia et al. (2007)
Table 2. Features of non-adult pathological conditions that may cause rib lesions including the β-thalassaemias

<table>
<thead>
<tr>
<th>Element</th>
<th>Non-specific respiratory infections</th>
<th>Trauma</th>
<th>Vitamin D deficiency</th>
<th>β-thalassaemia intermedia and major</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ribs</td>
<td>- visceral rib lesions (7, 14)</td>
<td>- complete or partial rib fractures</td>
<td>- flaring, thickening and increased porosity at costo-chondral end (6)</td>
<td>Costal osteoma</td>
</tr>
<tr>
<td></td>
<td>- lesions as layers of dense periosteal new bone, resorption or plaque formation (1)</td>
<td>- rachitic children are susceptible to respiratory disease such as whooping cough leading to rib fractures due to weakened bone (4, 12)</td>
<td>- fractures may arise as a result of weakened bone through continuous coughing, vomiting or overly tight swaddling (3, 4, 12, 19)</td>
<td>- benign neoplastic lesion mostly formed of lamellar bone (11)</td>
</tr>
<tr>
<td></td>
<td>- expansion: new bone blends into existing bone shape (14)</td>
<td>- stress fractures due to continuous coughing and/or vomiting (3)</td>
<td></td>
<td>- swollen aspects of hyper-marrow bone on the original cortex of the rib (17, 21)</td>
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<td></td>
<td>- proliferation most apparent towards vertebral end and often occurs bilaterally (9)</td>
<td>- often result from blunt chest trauma and deliberate injury (8)</td>
<td></td>
<td>- radio-opaque appearance (21)</td>
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<td></td>
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<td>- care must be taken when diagnosing and interpreting these osteomas without the use of radiographs, as they emerge similar to fracture calluses (21)</td>
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<td>Rib-within-a-rib</td>
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<td>- radiographs: radio-opaque band within the affected rib (16)</td>
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<td>- longitudinal marrow expansion within the cortex (16)</td>
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<td>- posterior aspects of the ribs expand posteriorly which may correspond with paraspinal extra-medullary haematopoiesis (16)</td>
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<td></td>
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<td></td>
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<td>- gross macroscopic examination: rib-within-a-rib appearance emerges as pitting and thickening of the ribs with hypertrophic aspects and costal osteomas (21)</td>
</tr>
<tr>
<td>Skull</td>
<td>- endocranial lesions may be observed (20)</td>
<td>- additional if arisen secondary to underlying condition such as vitamin D deficiency</td>
<td>- craniotabes</td>
<td>- porotic hyperostosis, ‘hair-on end’ (2, 11, 16, 17, 21)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- porosity and pitting on ectocranial aspect (6, 15)</td>
<td>- cribra orbitalia (11, 17)</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>- generalised periosteal and/or trabecular thickening (11, 16, 17, 21)</td>
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<td></td>
<td></td>
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<td></td>
<td>- rodent face’ (13, 16)</td>
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<tr>
<td>Long bones</td>
<td>widespread new bone formation may be observed (20)</td>
<td>additional if arisen secondary to underlying condition such as vitamin D deficiency</td>
<td>bending deformities</td>
<td>increased porosity, angulation and swelling at growth plate</td>
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<tr>
<td>General skeleton</td>
<td>widespread new bone formation may be observed (20)</td>
<td>abuse victims may display more widespread trauma throughout the skeleton (10)</td>
<td>thickening, bowing and increased cortical porosity may be observed throughout</td>
<td>osteopenia: thinned cortex and meagre cancellous bone with reinforced vertical trabeculae (11, 21)</td>
</tr>
</tbody>
</table>

From 1, Pfeiffer (1991); 2, Hershkovitz et al. (1997); 3, Lovell (1997); 4, Pettifor and Daniels (1997); 5, Aufderheide and Rodriguez-Martin (1998, 307); 6, Ortner and Mays (1998); 7, Roberts et al. (1998); 8, Ogden (2000, 422-424); 9, Santos and Roberts (2001); 10, Barsness et al. (2003); 11, Ortner (2003); 12, Pettifor (2003); 13, Almeida and Roberts (2005); 14, Matos and Santos (2006); 15, Mays et al. (2006); 16, Tyler et al. (2006); 17, Lagia et al. (2007); 18, Keller and Barnes (2008); 19, Lewis (2010); 20, Lewis (2011); 21, Lewis (2012)
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