**Title:** Broken Childhoods: Rural and Urban Non-Adult Health during the Industrial Revolution in Northern England (Eighteenth-Nineteenth Centuries).

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Abstract

During the eighteenth and nineteenth centuries England underwent a period of rapid urbanization and industrialization. The detrimental effects of urban living conditions and child factory labor on the health of children during this time has been the subject of considerable debate and investigation by historians. It is generally understood that growing up in a rural environment was more conducive to healthy growth and development than within an industrial town. This study presents the first direct comparison of the bioarchaeological evidence for non-adult health from contemporaneous urban and rural sites from the north of England. Rural skeletal assemblages from this period are rare and most published studies are biased towards urban sites in the south of the country. Contrary to expectations, results revealed equal prevalence rates of metabolic and dental disease at both sites, but skeletons from the rural site had greater evidence of growth disruption and respiratory disease. Evidence for specific infectious disease and medical care in response to trauma were also identified. Our interpretations of rural/urban health during this period must take into account the dire consequences of social inequalities and economic migration. There is a tendency for the latter to be characterized as uni-directional – from country to town – without due consideration of rural industry and child migrant workers.
The eighteenth and nineteenth centuries in Britain were a period of rapid urbanization and industrialization; from A.D. 1801 to 1901 the proportion of people living in towns increased from one-fifth to four-fifths (Wohl 1983). By the mid-nineteenth century, almost half of the total population of England was under 20 years of age: “England was awash with children” (Humphries 2013: 115). Traditionally, children from poorer families will always have had to work, either within the domestic context, or as apprentices and laborers. What changed with the industrial revolution was the nature of that work, with a dramatic increase in non-domestic labor by women and children (Goose and Honeyman 2013: 4). Children of the urban poor in the eighteenth and nineteenth centuries have long been conceptualized as the passive, exploited, victims of an insatiable industrial machine, whose wants and needs were overlooked by a laissez-faire society focussed on economy and Empire (Gowland 2015b). In a speech to the House of Lords in 1873, Lord Shaftesbury famously described children leaving the factory gates during the earlier part of the nineteenth century as “a set of sad, dejected, cadaverous creatures... the sight was most piteous, the deformities incredible. They seemed to me, such were their crooked shapes, like a mass of crooked alphabets”. This account is borne out by numerous other descriptions of working class, urban, children at this time (e.g. Engels 1950; Gaskell 1833). By contrast the rural environment during this period was conceptualized as much healthier for children; fresh, unpolluted air, with plenty of sunshine, a more varied diet, less overcrowding and better sanitation. As Gaskell (1833: 200) observed: “A child from an agricultural district, eight or nine years old, blooming with rosy health, full of vitality, is transported to a crowded town ... it is exposed to miasmatic evaporation, shut up in a narrow street, its home is damp and
cold, its food poor and badly cooked: in a few months, from the force of these circumstances, the Hebe-looking child has become pallid”.

These sentiments were echoed by the nineteenth century physician Francis Sharp (in the Reports from the Commissioners 1833), who moved from a rural practice near Scarborough, North Yorkshire, to work in the city of Leeds: “The general appearance of the children in Leeds immediately struck me as much more pallid, and also the firmness of the fibre as much inferior to what I had seen in Scarborough and the adjacent country... If it were not for the individuals who join the mills from the country the factory people would soon be deteriorated”.

There was a general anxiety that urban childhoods, particularly when spent undertaking factory labor, would result in the eventual physiological deterioration of the nation. These concerns sparked a number of enquiries into the health of urban and factory children and a vast amount of qualitative and anthropometric data was commissioned from across the country (Reports from the Commissioners 1833). The results of these endeavors indicated that rural children were, as suspected, on average 3cm taller than their urban counterparts (Kirby 2013). It was concluded, however, that factory labor per se was not responsible for the ‘stinted’ growth and low life expectancy, but rather insalubrious urban conditions more generally, combined with the perceived lowly habits and morality of the poor (Chadwick 1842). There are numerous descriptions of the filthy, overcrowded, insanitary, polluted living conditions in many of the poorer slum areas. So-called ‘back-to-back’ housing in cities such as Bradford and Leeds also allowed very little natural light into the rooms, and this
combined with severe air pollution meant that exposure to sunshine, vital for proper
mineralization of the bones and immune function in growing children, was limited
(Brimblecombe 1978; Hudson 1992). However, some contemporary accounts of rural life
also point to inadequate and over-crowded houses, as well as polluted water supplies
spreading diseases such as typhus (Wilkes 2011). The nineteenth century epidemiologist
William Farr felt that there was an often “fanatical” fixation on the declining conditions of
urban centers, stating that ‘epidemics desolated the country as well as towns, though to less
extent...’ (quoted in Young and Handcock 1956; 771). Rural populations were also highly
susceptible to periods of food shortages and crop failures, and a series of poor harvests
could result in poverty and famine (Bogin 1999). Alfred Ireson, who was born in 1856,
described rural life during this period as “a time of trial and difficulty. The agricultural
workers had long hours, they pay barely enough to keep body and soul together. The
condition of the children in many cases was pitiable. Rough food and clothes; everything
depended on the skill and character of the mother” (quoted in Burnett 1984: 83). While
protective measures regarding child labor in factories were instigated, those working within
agriculture and rural cottage industries were overlooked until the Education Act of 1876,
which decreed that children under 10 years of age could no longer be employed in
agricultural work (Burnett 1984; Pinchbeck and Hewitt 1973).

The vast majority of research concerning childhood health during this period is derived from
historical data and records (Kirby 2013). Whilst these are extremely valuable, some of the
qualitative reports depict partisan viewpoints and political polemics rather than objective
observation (e.g. Engels 1950). Recent excavations of cemeteries from the eighteenth and
nineteenth centuries have highlighted the important contribution of bioarchaeological
evidence for examining the *direct* physiological impact of these environments on children
living at the time (e.g. Brickley and Miles 1999; DeWitte et al. 2015; Geber 2016; Lewis 2002;
Mays et al. 2009; Newman and Gowland 2016; Pinhasi et al. 2006). Childhood growth and
morbidity is multidimensional in etiology, but is widely documented as being a sensitive
indicator of socio-economic environment (Lewis 2007). Recent bioarchaeological analyses
have been strongly biased towards urban population samples, because so few rural sites
dating to this period have been excavated. Mays et al. (2008) provide a valuable comparison
of urban Birmingham with rural Wharram Percy, but the latter site was in use for a
millennium and predominantly medieval, thus lacking chronological resolution. This is
important, because rural lifeways were not static and unchanging across the last
millennium; rural industry also developed apace during the late eighteenth and early
nineteenth centuries.

The aim of this study was to compare and contrast rural and urban skeletal evidence for the
health and mortality of children from two sites in the north of England in order to assess the
historically accepted idea that rural environments were more conducive to better childhood
health. This research provides a rare insight into a rural post-medieval assemblage (now
reburied) and thus also serves to enrich the existing corpus of bioarchaeological knowledge
for this period, which has so far been largely confined to urban sites. This analysis of sites
from the north of England also provides an important comparative data-set given the
current bias towards sites from the south and midlands. The bioarchaeological data will be
integrated with existing historical evidence to obtain a richer understanding of living and working conditions in both environments.

**Materials**

Skeletal remains were analyzed from the rural churchyard of St Michael and St Lawrence, Fewston, North Yorkshire and the urban site of Coach Lane, North Shields, Tyneside (Fig. 1).

**Fewston, North Yorkshire**

Fewston is a small village located in the Washburn Valley, near Harrogate in North Yorkshire. The skeletal assemblage was excavated from the parish churchyard in advance of building work by John Buglass Archaeological Services in 2009-2010 (Buglass 2009; Caffell and Holst 2010) and was reburied in September 2016. This assemblage is exceptional for several reasons:

1) It is one of only a few predominantly post-medieval rural skeletal collections.

2) Twenty-one of the excavated individuals were confidently identified based on coffin plates and grave monuments.

3) Diaries of John Dickinson, a late nineteenth century resident of the adjacent village of Greater Timble, provide important historical insights into everyday life in the Washburn Valley. The diaries also refer to a number of the identified individuals within the skeletal assemblage, thus providing unique personal insights.
The Church of St Michael and St Lawrence in Fewston was rebuilt in A.D. 1697, most of the
medieval church having been destroyed in a fire (Alred 1997: 67). The graveyard would have
been in use from at least the fourteenth century, and some of the excavated skeletons could
date to the medieval period. Only a relatively small area (approximately 300 m²), however,
was subject to excavation and coffin plates and headstones indicated that the better-
surviving burials in this location were predominantly of nineteenth century date. The
cemetery closed in A.D. 1896, although two individuals who died in the early 20th century
did gain permission to be buried there. A total of 154 individuals were excavated, including
50 non-adults, a category defined here as less than 20 years of age. The more usual cut-off
age for non-adults in bioarchaeological studies is either 17 or 18 years; however, this was
extended for the purposes of this study in recognition that early life adversity (as evident in
many of the skeletons within this sample) is known to cause delays in the chronological age
of attainment of skeletal maturation. In the absence of a more reliable age from dental
development, therefore, some of the individuals within the sample may have been slightly
older than the age that their epiphyseal union suggested.

The use of the term non-adult in this context does not, of course, reflect nineteenth century
social norms regarding the chronological age of attainment of adult status, but denotes the
skeletal age only. Parish records indicate that the social status of those buried within the
cemetery was mixed, including well-off landowners as well as those of lower status, e.g.
laborers. Agriculture was the predominant occupation in Fewston during this period, but
from the late eighteenth century there was also a textile industry, with Westhouse Mill
established at nearby Blubberhouses in A.D. 1791 (Alred 1997: 30-31).
Coach Lane, North Shields

Coach Lane was a former Society of Friends burial ground (c. A.D. 1711-1857) located in North Shields, just east of Newcastle-upon-Tyne. North Shields in the late eighteenth and early nineteenth centuries was a heavily populated shipping, fishing, and coal-mining community, undergoing rapid development (Proctor et al. 2014). The town was heavily industrialized and in a Government report, the environmental conditions were considered derisory, with narrow streets described as “more justly compared to wells filled with noxious emanations...” (Second Report of the Commissioners 1895: 19). Numerous epidemics, including cholera occurred during this period, and air and water pollution from the various industries was extensive (Proctor et al. 2014). The social status of those buried at the Coach Lane cemetery was mixed, with Quakerism attracting those from different spheres of life. The burial records occasionally document the profession of the deceased and these included: “junior painter”, “grocer”, “shopkeeper”, “master mariner” and “linen weaver”. There were many prominent, high status Quakers within the north-east during this period, some of whom were buried at Coach Lane, along with servants (Proctor et al. 2014). The effects of a heavily polluted environment, however, would have been detrimental to all social classes (Gowland and Newman 2018). There is direct evidence that at least some of the children buried at the site had been engaged in factory labor, with the skeleton of one individual (aged 12-14 years) possibly exhibiting the notorious occupational condition “Phossy Jaw” – osteonecrosis of the mandible typically caused by exposure to phosphorus during match-making (Roberts et al. 2016). In contrast to Fewston, the Coach Lane cemetery
was excavated in its entirety (Procter et al. 2014) and consisted of 236 individuals, 81 of
whom were non-adults (under 20 years of age).

**Methods**

Age-at-death of the non-adults was estimated using dental development and eruption, long
bone growth, and epiphyseal fusion (see Table 1 for details). Following convention, dental
age was considered more representative of the chronological age of the child, as tooth
development is less affected by environmental adversity than skeletal development (Elamin
and Liversidge 2013; Saunders 2008—although see Discussion). When dental age could not
be established due to missing teeth, age was instead estimated from epiphyseal fusion and
metric analysis. Long bone diaphyseal lengths were measured and compared to age-at-
death estimated from dental age as well as Maresh’s (1955) known age long bone reference
data. Skeletal growth profiles were constructed from diaphyseal lengths of the major long
bones using the mid-point of the dental age-range, whereby an individual with a dental age
estimate falling between 0.5 to 1.5 years would be categorized as ‘1 year’. This follows the
practice of previous growth studies (see Mays et al. 2008). These age categories act as a
proxy for ‘known age’, and it is not intended that the age midpoint should be taken as a
genuine reflection of the precise chronological age.

Recent methods to detect growth disruption were applied, including measurements of
vertebral body heights (Newman and Gowland 2015) and the left femur was selected for
cortical thickness (CT) measurements (substituted with the right side when necessary)
(Mays et al. 2009). Radiographs were taken using a MPX 10 portable radiography unit set at 70kVp, 0.500mAs and with a Carestream Point-of-Care digital CR reader. Measurements of the total bone width (T) and the medullary width (M) were taken from the mid-shaft of the femur (Mays et al., 2009). Cortical thickness was determined as T-M, and plotted against the dental age (see Newman and Gowland 2016). Finally, skeletal indicators of puberty were recorded following Shapland and Lewis (2013, 2014).

The pathological lesions recorded include rickets (vitamin D deficiency), scurvy (vitamin C deficiency), cribra orbitalia, dental enamel hypoplasia, dental caries, non-specific periosteal reactions on the major long bones, endocranial and ectocranial lesions, maxillary sinusitis, rib lesions and evidence for tuberculosis (Table 1). Rickets was diagnosed through the presence of bowed long bones, flared and/or porous metaphyses and sternal rib ends, and the presence of medial tilting of the distal tibia, while coxa vara and flattening deformities in the proximal femur were also considered (Brickley and Ives 2008; Mays et al. 2006; Ortner and Mays 1998;). Scurvy was diagnosed through the presence of woven bone in the orbits, porosity on the mandible and/or maxilla (especially focused on the internal ramus and coronoid process of the mandible, and around the infraorbital foramen of the maxilla), porosity on the greater wing of the sphenoid, and porosity in the supraspinous fossa of the scapula (Ortner 2003; Stark 2014). The pathological lesions associated with metabolic disease may be very subtle (Brickley and Ives 2008) and this is confounded further by the porous appearance of the rapidly growing bones of infants and children. All pathologies were therefore recorded by two bioarchaeologists working side-by-side (Caffell and Gowland) and the results compared to those of an observer working independently (Newman). Rickets and scurvy were only diagnosed as present when both sets of independent observations were positive. If one independent observer identified indicators
of rickets or scurvy, but the other did not, then these skeletons were re-examined and
agreement reached. Due to the multiple skeletal elements that can display signs of vitamin
C or vitamin D deficiency, only crude prevalence rates (per individual) for these pathologies
were calculated; however, individuals for whom the diagnostic skeletal elements were
absent were excluded from the prevalence calculations.

All long bones were examined for the presence of non-specific periosteal new bone growth
in the form of woven bone, lamellar bone, and/or abnormal porosity. Likewise, the
endocranial and ectocranial surfaces and maxillary sinuses were examined for new bone
formation (woven and/or lamellar) and porosity. All deciduous and permanent teeth
(including unerupted but observable tooth crowns) were examined for defects in enamel
formation (lines, grooves or pits), and the prevalence rates for dental enamel defects were
expressed as a percentage of all of the teeth present (including observable unerupted
teeth). Cribra orbitalia was recorded using the Stuart-Macadam (1991) system, but only
presence or absence is reported here and results are provided as true prevalence rates
(percentage of preserved orbits affected). The minimum number of ribs present on the left
and right side for each individual was counted, as was the minimum number of ribs showing
new bone formation (woven or lamellar). The number of un-sided rib fragments was
counted, along with those exhibiting pathological lesions. Crude prevalence rates were
calculated for the minimum number of ribs affected as a proportion of the minimum
number of ribs present.

Results
Age-at-death

The age distribution of the non-adults from Coach Lane follows the pattern frequently observed at archaeological cemetery sites, with a high proportion of deaths in those under 5 years of age and relatively few deaths during adolescence (Fig. 2). By contrast, Fewston presents with a more unusual non-adult age distribution, with higher numbers of adolescent deaths. Only two individuals at Fewston were aged less than 1 year, one of whom was a named individual. The lack of infants is likely to have resulted from the common practice of spatial segregation of this age group within cemeteries and the fact that only a relatively small area was excavated.

The difference in age structure between the two sites with respect to the excess of adolescents at Fewston, is important to consider. Is this unusual age profile simply an artefact of the incomplete excavation of Fewston, or could it reflect genuine differences in age-structure and mortality risks between the sites? If the latter, then what was the driving force behind this? We hypothesize that this difference reflects genuine mortality risks in this section of the cemetery, which is dominated by nineteenth century burials, and that it is driven by the migration of adolescents from poor urban areas seeking employment in rural industry. This will be explored further with reference to the results for the pathological lesions below. Comparison of the various pathologies between the two sites will also explicitly consider the different age-at-death profiles when inferring aspects of the lived experiences of these children.
Growth

Diaphyseal lengths, when plotted against dental age, were similar between Fewston and Coach Lane; unfortunately only relatively few measurements were possible from the former site due to poorer preservation (Fig. 3). These values aligned with Maresh’s (1955) data up until approximately 10 years of age, after which most adolescents drop below the modern values. This could relate to the relatively delayed pubertal growth spurt in these individuals compared to modern children (Fig. 4). In keeping with the findings of Mays and colleagues (2009), cortical thickness appears to be a more sensitive indicator of adversity and was compromised in children from both sites. Growth retardation was observed particularly in children older than 4 years of age (Fig. 3). While only small numbers of femora were available for measurement from Fewston, cortical thickness measurements of the humeri and tibiae from these sites were similarly reduced when compared to modern norms.

Vertebral measurements were taken following the protocol outlined by Newman and Gowland (2015), which identified measurements from vertebral groupings of C5-6, T6-8, and L2-4 as being reliable indicators of potential population differences in growth patterns. Due to the C5-6 group presenting a more robust sample within this study for both sites, these measurements were plotted against dental age (Fig. 5). Values for several adolescents from Fewston were anomalously low and an ANCOVA test revealed that vertebral body height was statistically different \( [F(1,35)=13.551, \ p=0.001] \) between the two sites, whilst adjusting for dental age. These adolescents also exhibited a range of skeletal indicators of poor health. For example, Skeleton 262 had extensive new bone formation and severe enamel hypoplasia; Skeleton 338 had severe marrow hyperplasia, woven bone on multiple
skeletal elements and evidence for scurvy; Skeleton 208 had sinusitis and cuspal enamel hypoplasia as defined by Ogden et al. (2007), and Skeleton 331 had evidence of scurvy, rickets, enamel hypoplasia, and new bone formation on the ribs.

Pathology

A wide range of pathological lesions were observed in the non-adult skeletons from both sites. Dental enamel defects were common in both the deciduous and permanent dentitions. A greater proportion of non-adults at Fewston expressed dental defects than at Coach Lane (72% compared to 50%), but the true prevalence rate for defects was similar at both sites (Table 2), with no statistically significant difference between them (two sample t-test, p > 0.1).

There were some differences in the type of the dental enamel defects, however, with plane-form and cuspal enamel defects more frequent at Fewston (4% of all teeth compared to 0.6% at Coach Lane). Deciduous as well as permanent teeth were affected by enamel hypoplasia at both sites, including pitting defects, but cuspal enamel defects in deciduous teeth were not common (Supplementary tables 1-4). In such instances, the period of stress that produced these lesions would have commenced in utero and have occurred during the formation of both the deciduous and permanent dentition.

The crude prevalence of rickets was similar between sites and is very high when compared to contemporaneous sites in London (Lewis 2002). Likewise, the prevalence of scurvy was
high and there was no statistically significant difference between Coach Lane and Fewston (Table 3). Again, this prevalence was higher than the London sites of Chelsea Old Church, St Benet Sherehog, and Bow Baptist (ranging from approximately 4-10%), but were more comparable with rates seen at Cross Bones (37%), a site that was located in a particularly impoverished parish of London during this period (Newman and Gowland 2016). Developments in the diagnostic criteria for both rickets and scurvy since some of these reports were published are partly responsible for the disparity, along with the exclusion of poorly preserved skeletons from the prevalence calculations in this study. The north of England, however, was a renowned focal point for rickets in the nineteenth century (British Medical Journal 1889). The prevalence rate of cribra orbitalia was, again, high at both sites, but Fewston exhibited significantly higher frequencies (Table 3).

Non-adults at both sites showed evidence of maxillary sinusitis and rib lesions, but the prevalence of both was significantly higher at Fewston (Table 4, rib lesions p <0.05; sinusitis, p < 0.1) and most particularly in children of working age (12-20 years). By contrast endocranial and ectocranial new bone was slightly higher at Coach Lane, although this was not statistically significant (Table 4). At Coach Lane, infants from 1-12 months were much more likely to exhibit endocranial new bone growth (70%) than other age groups, whereas at Fewston higher prevalence rates were observed again amongst the adolescents. Care must be taken when interpreting cranial lesions in such young infants, because rapid bone growth during infancy may be mistaken for pathological lesions (Lewis 2007). Periosteal new bone formation (NBF) was recorded for all the major long bones and was much more prevalent amongst the Coach Lane population compared to Fewston. Periosteal new bone
formation at this site was most frequently expressed in the tibiae (TPR = 55%), whilst at Fewston the femur was more frequently affected (TPR = 22%) (Table 5).

In terms of specific diseases amongst the non-adults, there were three individuals of particular note. The first was skeleton 69 from Coach Lane, who had severe osteonecrosis of the mandible accompanied by woven bone formation. This has been interpreted as likely due to phosphorous poisoning, which was a notorious occupational condition associated with the match-making industry at this time. A detailed description of this individual was provided by Roberts and colleagues (2016).

Skeleton 223 (14-15 year old) from Fewston had multiple lytic lesions accompanied by new bone formation in the skull. Unfortunately, the skull of this individual was highly fragmented and only parts of the skull, vertebrae and upper ribs were preserved. Both orbital roofs had thick deposits of lamellar bone, with woven bone also present in the left orbit. A circular lytic lesion approximately 14mm in diameter, with sharp clear margins, had penetrated the thickness of the right orbital roof (Fig. 6). Four further lytic lesions were present elsewhere on the cranium. The surrounding external surfaces of these generally had scalloped roughened areas, with deposits of partially remodeled woven bone and lamellar bone (Fig. 7). The body of the vomer was enlarged and covered with well-remodeled lamellar bone, with a smooth circular depression in both lateral walls and the perpendicular plate was absent (apparently remodeled). A large shallow lytic lesion was present on the internal surface of the right mandibular ramus and the margins were again bounded by thick deposits of partially remodeled woven bone. Deposits of lamellar bone extended along the
inferior surface of the right mandibular body and a plaque of lamellar bone was present on
the external surface of the right ramus. The axis had a lytic lesion on the external surface of
the left lamina surrounded by partially remodeled woven bone. An upper right rib had a
deposit of porous lamellar bone on the visceral surface of the neck. None of the other rib
fragments had any evidence for lesions, but few of the rib heads and necks survived.

Tuberculosis (TB) manifested in the skull, whilst uncommon in adults, can occur more
frequently in children (Dawson and Robson-Brown 2012; Ortner 2003: 247-248). Multiple
cranial lesions, usually circular, less than 20 mm in size, which penetrate the thickness of the
vault, as observed here, are all features of childhood TB. While bone formation around the
lesions is not typical, it can occur (ibid.). Chaudhary et al. (2004) and Sheikh et al. (2012)
state that involvement of the jaw is more likely in older individuals (but can occur in
children) and typically manifest as destructive lesions in the mandible, with potential
involvement of the temporo-mandibular joint. Tuberculosis of the orbit is more common in
children and is typically unilateral (Dalvin and Smith 2016), as expressed here. Blindness is a
frequent complication of ocular tuberculosis (ibid.; Shameem et al. 2009) and, given the
severity of the lesions, it seems likely that Skeleton 223 experienced a similar outcome.

Endocranial new bone formation and rib lesions have also both been associated with
tuberculosis (Lewis 2004, 2007; Matos and Santos 2006; Roberts and Manchester 2005, 190;
Santos and Roberts 2001, 2006).

Finally, the skeleton of a 12-14 year old from Coach Lane (Skeleton 157) exhibited direct
evidence of medical care in the form of cranial surgery in association with trauma. The
individual had suffered a peri-mortem blunt-force injury to the left frontal squama, located
44 mm superior to the left orbit rim and 20 mm medial to the coronal suture (Fig. 8). On the ectocranial surface a smoothly curved fracture line extended superiorly and posteriorly, terminating in the coronal suture 45 mm from bregma. The lateral segment was not detached, but displaced internally slightly at the inferior/medial end. At this inferior part, the fracture extends at an angle through the bone to the internal surface, where a fracture line extends superiorly for 13.6 mm. A second bevelled peri-mortem fracture line extends inferiorly and laterally to the temporal crest. A neat circular peri-mortem trepanation is located in the left frontal squama 11.7 mm medial to the curved fracture line, 5.2 mm from the coronal suture and 12.8 mm from the metopic suture. The external diameter (16.7 mm) is slightly larger than the internal diameter (14.1 mm) (Fig. 8). While the external margins are slightly crushed, the internal margins are sharp, and the walls are smooth. This represents an unsuccessful attempt to provide life-saving medical treatment in response to the peri-mortem injury.

Discussion

The results initially appeared to contradict the sentiments espoused at the time-i.e. that urban environments were much more damaging to childhood health than a rural upbringing. Skeletal remains from both places show very high levels of metabolic disease and other non-specific indicators of physiological disruption. The higher prevalence of many of the pathological conditions, together with poorer growth, amongst some of the children from Fewston compared with Coach Lane was contrary to expectations and historical records. Two questions arise from these results: why did the children at Fewston suffer so much chronic health stress and why were there so many adolescents at this site? Usually,
the adolescent age category is one of the least well-represented within archaeological
cemetery populations (as observed at Coach Lane); a factor lamented in recent studies of

A direct interpretation of this skeletal evidence would lead to the conclusion that the rural
environment was at least as deleterious to health as the urban environment. Historical
evidence for the Washburn valley in which Fewston is located does in fact point to
inadequate, overcrowded and insanitary housing for some rural-dwellers, along with
polluted water supplies. For example, local resident John Dickinson noted in his diary entry
for 1st October 1881: “There is part [some] talk about the deficient water supply and
drainage of the village, and it is thought the sanitary authority will take the matter in hand
themselves and compel the proprietors to alter things. The proprietors fear the cost and so
it happens that we drink water highly polluted with sewage and our sinks [drains] are simply
beastly”. In A.D. 1840, Elizabeth and William Scaife lost three children (John aged 2 years,
Hannah aged 9 years and Christiana Eliza aged 6 years) within just one week to typhus fever,
spread via contaminated water during an epidemic that lasted two years from A.D. 1838.

While such infectious diseases undoubtedly took a toll on the morbidity and mortality of the
Fewston population, they would have proven more fatal to younger children, rather than
adolescents, as the dreadful suffering of the Scaife family attests. These diseases are also
acute; they will kill quickly and leave no visible sign on the skeleton. They are not, therefore,
responsible for the evidence of chronic disease observed amongst many of the adolescent
skeletons. When interpreting the prevalence of pathological lesions, the younger age-at-
death profile of the children at Coach Lane should be considered, particularly with regard to
the osteological paradox (Wood et al. 1992). However, given that many of the lesions
recorded and interpreted at both sites were at least initiated during the first few years of
life (e.g. enamel hypoplasia, rickets, scurvy and cribra orbitalia) the comparison of the two
different age-at-death profiles continues to have merit.

The presence of documented adults at Fewston has made it possible to gather the
associated birth and death certificates, together with census records. This provides a known
record of individuals who were born, lived and died in Fewston and its environs – i.e. the
survivors. The skeletons of these adults do not show comparable levels of pathological
lesions (e.g. cuspal enamel hypoplasia) and their stature does not appear to have been
compromised; indeed for males it was above the average stature for post-medieval England
(177.3 cm for Fewston males and 160.4 cm for Fewston females) (Caffell and Holst 2017).
One could argue that the children in the assemblage were non-survivors and that this is why
they show a higher prevalence of pathology compared to the adults (Wood et al., 1992).
Another hypothesis (mentioned above) is that these adolescents were not local to the area
and instead represent migrants. This interpretation will be explored further after a more
detailed comparison with Coach Lane.

Coach Lane was widely known to be a heavily industrialized, polluted and insalubrious town
in the nineteenth century (Proctor et al. 2014). The prevalence of metabolic, respiratory and
non-specific indicators of health stress at this site was high, as expected. In A.D. 1889 the
British Medical Association estimated that vitamin D deficiency in the form of rickets in
childhood was concentrated in the heavily industrialized areas of the northeast, where
Coach Lane was located (British Medical Journal 1889). Vitamin D deficiency at this site is
likely to have been exacerbated by the more northerly latitude, in conjunction with air
pollution from the associated industries.

The high levels of vitamin D deficiency observed at Fewston, however, were not in keeping
with historical records. John Snow, the famous nineteenth century spatial epidemiologist,
best known for his work on cholera, began his medical career as a doctor in a North
Yorkshire village close to Fewston. Snow (1857) observed that rickets was a relatively rare
disease in Yorkshire when compared to London. He attributed this to people from Yorkshire
baking their own bread due to cheaper fuel sources, in contrast to Londoners who almost
universally ate shop-bought bread, heavily adulterated with alum. Snow, like his
contemporaries, believed that vitamin D deficiency was primarily dietary in origin, rather
than a lack of sunlight, and he thought that alum was responsible for ‘weakening’ the bones.
The connection between sunlight and vitamin D deficiency was not made until the late
nineteenth century and diet was still thought to be the predominant cause of the disease
even in the 1920s (see Findlay and Mellanby 1922). The key point here is that the
prevalence of rickets at Fewston is contrary to Snow’s observations in North Yorkshire.

The prevalence rates of scurvy were also equally high in both the rural and urban sites
studied here. It was expected that scurvy would be less prevalent in rural environments due
to the increased availability of fruit and vegetables. A regional survey of diet and nutrition
highlights that in rural Yorkshire the working classes ate largely bread, oatmeal, some sugar
and weak tea, rather than vitamin C rich food such as potatoes (Horrell and Oxley 2012).

Historical records have also documented that the diet of women and children was more monotonous and less likely to include meat compared to men (Horrell and Oxley 2012; Humphries 2010).

Dental enamel hypoplasia (DEH) prevalence was comparable between the sites. DEH has multiple etiologies (e.g. infectious disease, trauma, malnutrition), and is generally regarded as a ‘non-specific’ indicator of disease; however, it is worthwhile focusing on the type of hypoplastic defects present. While linear enamel hypoplasia is by far the most common form exhibited in skeletal samples generally, at Fewston there are a number of individuals that have ‘plane-form’ and ‘pitted’ defects, of the type described by Ogden et al. (2007), as ‘cuspal enamel hypoplasia’ (Fig. 9). This term describes the combination of pitted and plane-form lesions, together with a disruption in the pattern of the molar cusps, such that multiple small cusps form and develop. Ogden and colleagues (2007: 960) noted a relatively high number of individuals with such defects (on deciduous and permanent teeth) from the post-medieval site of Broadgate in London and explored a range of reasons for these defects, including vitamin D deficiency. Pitting and cuspal enamel hypoplasia is also observed on the deciduous teeth at Fewston, indicating an in utero insufficiency. Purvis and colleagues’ (1973) clinical study noted an association between very similar hypoplastic lesions in the deciduous teeth and neonatal tetany, arising from deficiency in vitamin D in the mothers during the third trimester of pregnancy. Their sample was from Scotland, a northerly latitude, and they observed an increase in frequency during winter and spring when hours of sunshine were lowest. They also noted an association with increased maternal parity and
low social status. Similar instances of modern cases of enamel hypoplasia associated with nutritional rickets have been reported in the more recent medical literature (e.g. Davit-Béal et al. 2014). The presentation of these defects are identical to those observed amongst the Fewston skeletons in terms of the teeth affected and location. Vitamin D deficiency in pregnant women is known to result in adverse maternal health and birth outcomes impacting on the birth weight, bone mass and immunity of the infant (Karras et al. 2016; Paterson and Ayoub 2015).

An early twentieth century medical book by Dick (1922) focusing on the geographical distribution and clinical manifestations of rickets, also describes dental lesions, which he argued to be pathognomonic of vitamin D deficiency as follows: “There may be only a pitting, producing a honeycombed appearance of the enamel, or the enamel covering is slight, and the cutting edge of the tooth presents sharp points, giving a characteristic appearance to the tooth.” (Dick 1922: 191-192). This description is again consistent with the dental lesions observed in some of the Fewston individuals. While it is easy to dismiss these early medical texts as out-dated, particularly when published during a period in which the etiology of the condition was only poorly understood, it is important to give credit to the fact that these observations were made during a time when the disease was frequently encountered.

While enamel hypoplasia is generally considered to be a non-specific form of stress; in these instances, it seems likely to have resulted from an insufficiency of vitamin D and/or calcium, beginning in utero and extending until two years of age. Vitamin D deficiency contributes to
hypocalcemia and hypophosphatemia, the latter of which is compounded by hyperparathyroidism brought about by hypocalcaemia (Foster et al. 2014: 6-7). This lack of calcium and phosphorus results in developmental and mineralization defects in teeth via both direct and indirect mechanisms (Foster et al. 2014). It has been observed in clinical studies that vitamin D deficiency can result in inter-globular dentine and this has been identified in archaeological teeth of individuals with residual rickets (D’Ortenzio et al. 2016).

Future analysis of the Coach Lane site may benefit from a histological analysis of the teeth, but the Fewston skeletons have now been reburied. Of interest is that none of the adults from Coach Lane or Fewston exhibited evidence of cuspal enamel hypoplasia, indicating a strong association between these lesions and early life mortality risk.

When interpreting the presence of pathological lesions and growth disruption early in the life course, it is important to consider the significance of maternal health for growth and immune status (Gowland 2015a). Maternal malnutrition, ill health, and exposure to toxins within the living or working environment during pregnancy will adversely affect the developing fetus. Intrauterine growth retardation is strongly associated with shorter height in childhood and adolescence (Chung and Kuzawa 2014). Furthermore, a mother who fails to meet her own genetic potential in growth is more likely to have an infant who is small for gestational age (SGA). Growth deficits early on in life can be difficult to catch up and particularly so in adverse biocultural environments (Chung and Kuzawa 2014). Indeed, the effects of a poor environment on growth could span multiple generations before it is fully reversed despite alleviation of environmental stressors (Barker 2012).
Finally, when interpreting palaeopathological evidence and delayed growth in terms of the local living environments, we must be sure that the pathological conditions observed in the skeletons of the children from Fewston and Coach Lane do in fact relate to the local area. With regard to this period of rapid industrialization and urbanization, migration is often conceptualized as uni-directional – from the countryside to towns. It is important to consider that during this period there was a significant amount of rural industry and in some instances this led to the apprenticeship of urban children, who were removed to the countryside to work as laborers, domestic servants, or in textile mills. These individuals were known as pauper apprentices: children taken from workhouses in cities, usually aged between 10-12 years, and indentured to work far from their original home. The children worked long hours, 12 to 14 hours per day, and for no pay, but were instead ‘taught a trade’ and were housed, fed, and given some education (Honeyman 2013; Levene 2010).

There was a flax mill at Fewston, known as Westhouse mill and historical sources document the employment of pauper apprentices from the workhouses of Lambeth and Shoreditch in London, as well as elsewhere in England (Honeyman 2007). It therefore seems probable that this excess of adolescents at Fewston, with high frequencies of skeletal lesions and stunted growth, reflect a poor and destitute childhood in London, rather than one spent at Fewston. Ives and Humphrey’s (2017) study of diaphyseal growth in children from the poor London parish of Bethnal Green during the nineteenth century also highlights severe growth deficits. Social status is one of the strongest determinants of health and this was particularly evident in post-medieval England (Gowland 2018; Marmot and Wilkinson 2006). As discussed, the evidence for rickets and enamel hypoplasia at Fewston is consistent with the
pathological evidence from the low status post-medieval cemetery of Broadgate in London (Ogden et al. 2007; Pinhasi et al. 2006). The high prevalence of respiratory disease amongst these children at Fewston is also consistent with mill labor in which pulmonary complaints, including byssinosis and tuberculosis, were a known occupational hazard (Wohl 1983). Further isotopic and archival investigation is currently being undertaken in order to verify the non-local origins of these adolescents (Gowland et al. 2017).

**Conclusion**

While the children at Coach Lane undoubtedly showed evidence of adverse living conditions in the form of poor skeletal growth and the presence of physiological insults to the skeleton, it was surprising and contrary to historical evidence that the children from the rural site showed comparable health deficits. One explanation is that we need to re-evaluate our ideas of rural life during this time in terms of it providing a more beneficial living environment and diet. Alternatively, the health stress observed in some of the Fewston children may not relate to the local living environment at all, but instead to an impoverished urban childhood in the workhouses of London and elsewhere. Childhood mobility during this period as well as urban to rural migration should be factored into our interpretations. Industry was not just an urban phenomenon, but also a rural endeavor and we should consider the impact of this on the health and population structure of Fewston. Finally, poverty has heritable biological consequences via poor maternal health status. This biological disadvantage then becomes exacerbated by poor working and living conditions. The industrial revolution was a period of relative social mobility, with a rise in the middle classes, but for the very poor and disenfranchised, the inheritance of biological inequality in
terms of morbidity and mortality risks, in addition to social disadvantage, meant that only
the very tenacious could survive or thrive.

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Washburn Heritage Centre for their contribution and support with the Fewston assemblage
and Jenny Proctor at Pre-Construct Archaeology for access to the Coach Lane collection.

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and Bodies. Springer, New York.


<table>
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<th>Age-at-Death</th>
<th>Dental development</th>
<th>Moorrees et al. 1963a, b; Smith 1991</th>
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<tbody>
<tr>
<td>Long bone growth</td>
<td>Maresh 1955; Scheuer and Black 2000</td>
<td></td>
</tr>
<tr>
<td>Epiphyseal fusion</td>
<td>Scheuer and Black 2000</td>
<td></td>
</tr>
<tr>
<td>Growth</td>
<td>Cortical thickness</td>
<td>Mays et al. 2009</td>
</tr>
<tr>
<td></td>
<td>Vertebral body height</td>
<td>Newman and Gowland 2015</td>
</tr>
<tr>
<td>Pathologies</td>
<td>Metabolic disease</td>
<td>Brickley and Ives 2006, 2008; Mays et al., 2006; Ortner and Mays 1998; Ortner 2003; Stark 2014</td>
</tr>
<tr>
<td></td>
<td>Cribra Orbitalia</td>
<td>Stuart-Macadam 1998</td>
</tr>
<tr>
<td></td>
<td>Dental enamel defects</td>
<td>Hillson 1996; Ogden et al., 2007; Ogden 2008</td>
</tr>
<tr>
<td></td>
<td>New bone formation</td>
<td>Ortner 2003; Roberts and Connell 2004; Lewis 2007</td>
</tr>
</tbody>
</table>

Table 1: Methods of analysis.

| Age Group | Deciduous Teeth | | | | Permanent Teeth | | | |
|------------|-----------------|------------|-------|------------|-----------------|-----------------|-------|------------|-----------------|-------|
|            | Coach Lane      | Fewston    |        | Coach Lane | Fewston        |        |        | Coach Lane | Fewston        |        |
|            | no. teeth (no. DEH) | % DEH     | no. teeth (no. DEH) | % | no. teeth (no. DEH) | % DEH | no. teeth (no. DEH) | % | no. teeth (no. DEH) | % DEH |
| <36w       | 0               |            | 0               | 0 | 0             |            | 0 |            | 0             | 0 |
| 36w-1m     | 34 (0)          |            | 0               | 0 | 0             |            | 0 |            | 0             | 0 |
| 1-12m      | 128 (13)        | 10         | 7 (0)           | 0 | 3 (1)         | 33         | 8 (0) |            | 0             | 0 |
| 1-5.9y     | 382 (66)        | 17         | 131 (12)        | 9 | 108 (30)      | 28         | 202 (28) |            | 14            | 0 |
| 6-11.9y    | 45 (8)          | 18         | 67 (7)          | 10 | 108 (37)      | 34         | 338 (72) |            | 21            | 0 |
| 12-19.9y   | 0               |            | 3 (0)           | 0 | 243 (114)     | 47         | 475 (139) |            | 29            | 0 |
| Total      | 589 (87)        | 14         | 208 (19)        | 9 | 462 (182)     | 39         | 1023 (239) |            | 23            | 0 |

Table 2: Dental enamel hypoplasia at Coach Lane and Fewston.
<table>
<thead>
<tr>
<th>Age</th>
<th>Coach Lane</th>
<th>Fewston</th>
<th>Coach Lane</th>
<th>Fewston</th>
<th>Coach Lane</th>
<th>Fewston</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Rickets</td>
<td>CPR</td>
<td>Rickets</td>
<td>CPR</td>
<td>Scurvy</td>
<td>CPR</td>
</tr>
<tr>
<td>36w-1m</td>
<td>0(9)</td>
<td>0</td>
<td>0(0)</td>
<td>0</td>
<td>7(8)</td>
<td>87.5</td>
</tr>
<tr>
<td>1-12m</td>
<td>7(12)</td>
<td>58.3</td>
<td>1(1)</td>
<td>100</td>
<td>10(12)</td>
<td>83.3</td>
</tr>
<tr>
<td>1-5.9y</td>
<td>11(25)</td>
<td>44.0</td>
<td>2(4)</td>
<td>50</td>
<td>7(27)</td>
<td>25.9</td>
</tr>
<tr>
<td>6-11.9y</td>
<td>3(7)</td>
<td>42.8</td>
<td>6(9)</td>
<td>66.6</td>
<td>0(6)</td>
<td>0</td>
</tr>
<tr>
<td>12-19.9y</td>
<td>7(10)</td>
<td>70</td>
<td>4(15)</td>
<td>26.6</td>
<td>2(10)</td>
<td>20</td>
</tr>
<tr>
<td>Total</td>
<td>28(63)</td>
<td>44.4</td>
<td>13(29)</td>
<td>44.8</td>
<td>26(63)</td>
<td>41.2</td>
</tr>
</tbody>
</table>

Table 3: Crude prevalence rates of rickets and scurvy by age group and true prevalence rates of cribra orbitalia at Coach Lane and Fewston.

<table>
<thead>
<tr>
<th>Age</th>
<th>Coach Lane</th>
<th>Fewston</th>
<th>Coach Lane</th>
<th>Fewston</th>
<th>Coach Lane</th>
<th>Fewston</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sinus</td>
<td>TPR</td>
<td>Rib</td>
<td>CPR</td>
<td>Ecto</td>
<td>CPR</td>
</tr>
<tr>
<td>&lt;36w</td>
<td>0(0)</td>
<td>0</td>
<td>0</td>
<td>0(1)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>36w-1m</td>
<td>2(2)</td>
<td>100</td>
<td>1(11)</td>
<td>9.1</td>
<td>3(13)</td>
<td>23</td>
</tr>
<tr>
<td>1-12m</td>
<td>1(2)</td>
<td>50</td>
<td>1(19)</td>
<td>5.3</td>
<td>12(20)</td>
<td>60</td>
</tr>
<tr>
<td>1-5.9y</td>
<td>6(24)</td>
<td>25</td>
<td>1(25)</td>
<td>4</td>
<td>5(28)</td>
<td>17.9</td>
</tr>
<tr>
<td>6-11.9y</td>
<td>0(9)</td>
<td>0</td>
<td>0(4)</td>
<td>0</td>
<td>3(6)</td>
<td>50</td>
</tr>
<tr>
<td>12-19.9y</td>
<td>6(19)</td>
<td>31.6</td>
<td>4(9)</td>
<td>44.4</td>
<td>2(10)</td>
<td>20</td>
</tr>
<tr>
<td>Total</td>
<td>15(56)</td>
<td>26.8</td>
<td>7(69)</td>
<td>10.1</td>
<td>25(78)</td>
<td>32</td>
</tr>
</tbody>
</table>

Table 4: Prevalence rates by age and site for maxillary sinusitis (sinus), rib lesions (rib), ectocranial lesions (Ecto) and endocranial lesions (endo). For maxillary sinusitis true prevalence is provided, while for the other pathologies crude prevalence rates only are given.
<table>
<thead>
<tr>
<th>Age Group</th>
<th>Coach Lane (TPR)</th>
<th>Fewston</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Hum</td>
<td>Rad</td>
</tr>
<tr>
<td>36w-1m</td>
<td>10</td>
<td>21</td>
</tr>
<tr>
<td>1-12m</td>
<td>32</td>
<td>54.5</td>
</tr>
<tr>
<td>1-5.9y</td>
<td>16</td>
<td>2.4</td>
</tr>
<tr>
<td>6-11.9y</td>
<td>11</td>
<td>0</td>
</tr>
<tr>
<td>12-19.9y</td>
<td>27.7</td>
<td>22</td>
</tr>
<tr>
<td>Total</td>
<td>20</td>
<td>15</td>
</tr>
</tbody>
</table>

Table 5: True prevalence rates of periosteal new bone formation on the major long bones from each of the sites

Supplementary Tables

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Deciduous</th>
<th>Permanent</th>
<th>All Teeth</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DEH</td>
<td>Total</td>
<td>%</td>
</tr>
<tr>
<td>foetus &lt;36w</td>
<td>-</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>neonate 36w-1m</td>
<td>0</td>
<td>34</td>
<td>0.0%</td>
</tr>
<tr>
<td>infant 1-12m</td>
<td>13</td>
<td>128</td>
<td>10.2%</td>
</tr>
<tr>
<td>young juvenile 1-5.9y</td>
<td>61</td>
<td>382</td>
<td>16.0%</td>
</tr>
<tr>
<td>older juvenile 6-11.9y</td>
<td>8</td>
<td>45</td>
<td>17.8%</td>
</tr>
<tr>
<td>adolescent 12-19.9y</td>
<td>-</td>
<td>0</td>
<td>114</td>
</tr>
<tr>
<td>Total</td>
<td>82</td>
<td>589</td>
<td>13.9%</td>
</tr>
</tbody>
</table>

Table S1: Coach Lane: Enamel hypoplasia (pitting, lines and grooves)

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Deciduous</th>
<th>Permanent</th>
<th>All Teeth</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CEH</td>
<td>Total</td>
<td>%</td>
</tr>
<tr>
<td>foetus &lt;36w</td>
<td>-</td>
<td>0</td>
<td>-</td>
</tr>
<tr>
<td>neonate 36w-1m</td>
<td>0</td>
<td>34</td>
<td>0.0%</td>
</tr>
<tr>
<td>infant 1-12m</td>
<td>0</td>
<td>128</td>
<td>0.0%</td>
</tr>
<tr>
<td>young juvenile 1-5.9y</td>
<td>2</td>
<td>382</td>
<td>0.5%</td>
</tr>
<tr>
<td>older juvenile 6-11.9y</td>
<td>0</td>
<td>45</td>
<td>0.0%</td>
</tr>
<tr>
<td>adolescent 12-19.9y</td>
<td>-</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>2</td>
<td>589</td>
<td>0.3%</td>
</tr>
</tbody>
</table>

Table S2: Coach Lane: Cuspal Enamel Hypoplasia
<table>
<thead>
<tr>
<th>Age Group</th>
<th>Deciduous</th>
<th></th>
<th>Permanent</th>
<th></th>
<th>All Teeth</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DEH</td>
<td>Total</td>
<td>%</td>
<td>DEH</td>
<td>Total</td>
<td>%</td>
</tr>
<tr>
<td>foetus &lt;36w</td>
<td>-</td>
<td>0</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>neonate 36w-1m</td>
<td>-</td>
<td>0</td>
<td>-</td>
<td>-</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>infant 1-12m</td>
<td>0</td>
<td>7</td>
<td>0.0%</td>
<td>0</td>
<td>1</td>
<td>0.0%</td>
</tr>
<tr>
<td>young juvenile</td>
<td>1-5.9y</td>
<td>12</td>
<td>131</td>
<td>9.2%</td>
<td>16</td>
<td>112</td>
</tr>
<tr>
<td>older juvenile</td>
<td>6-11.9y</td>
<td>6</td>
<td>67</td>
<td>9.0%</td>
<td>53</td>
<td>247</td>
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<tr>
<td>adolescent</td>
<td>12-19.9y</td>
<td>-</td>
<td>3</td>
<td>-</td>
<td>111</td>
<td>455</td>
</tr>
<tr>
<td></td>
<td><strong>Total</strong></td>
<td><strong>18</strong></td>
<td><strong>208</strong></td>
<td><strong>8.7%</strong></td>
<td><strong>180</strong></td>
<td><strong>815</strong></td>
</tr>
</tbody>
</table>

**Table S3: Fewston: Enamel hypoplasia (pitting, lines and grooves)**

<table>
<thead>
<tr>
<th>Age Group</th>
<th>Deciduous</th>
<th></th>
<th>Permanent</th>
<th></th>
<th>All Teeth</th>
<th></th>
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<tbody>
<tr>
<td></td>
<td>CEH</td>
<td>Total</td>
<td>%</td>
<td>CEH</td>
<td>Total</td>
<td>%</td>
</tr>
<tr>
<td>foetus &lt;36w</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0</td>
</tr>
<tr>
<td>neonate 36w-1m</td>
<td>-</td>
<td>0</td>
<td>-</td>
<td>-</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>infant 1-12m</td>
<td>0</td>
<td>7</td>
<td>0.0%</td>
<td>0</td>
<td>1</td>
<td>0.0%</td>
</tr>
<tr>
<td>young juvenile</td>
<td>1-5.9y</td>
<td>1</td>
<td>67</td>
<td>1.5%</td>
<td>12</td>
<td>271</td>
</tr>
<tr>
<td>older juvenile</td>
<td>6-11.9y</td>
<td>0</td>
<td>3</td>
<td>0.0%</td>
<td>28</td>
<td>472</td>
</tr>
<tr>
<td>adolescent</td>
<td>12-19.9y</td>
<td>-</td>
<td>3</td>
<td>-</td>
<td>28</td>
<td>475</td>
</tr>
<tr>
<td></td>
<td><strong>Total</strong></td>
<td><strong>1</strong></td>
<td><strong>208</strong></td>
<td><strong>0.5%</strong></td>
<td><strong>40</strong></td>
<td><strong>815</strong></td>
</tr>
</tbody>
</table>

**Table S4: Fewston: Cuspal enamel hypoplasia**
Figure Headings

Figure 1: The location of the Coach Lane and Fewston sites.

Figure 2: The age-at-death distribution of the non-adults from Fewston and Coach Lane.

Figure 3: Femoral diaphyseal lengths and cortical thickness from Fewston and Coach Lane plotted against dental age as a proxy for known age and modern comparative data.

Figure 4: Pubertal stages in Coach Lane (n = 11) and Fewston (n = 24) males and females plotted against dental age.

Figure 5: Vertebral body height for C5 and C6 plotted against dental age.

Figure 6: Lytic lesions penetrating the right orbit of skeleton 223.

Figure 7: Lytic lesion and new woven and lamellar bone on the left frontal bone of skeleton 223.

Figure 8: Frontal and parietal bone of skeleton 157 with perimortem trauma (marked B), a radiating fracture (marked A) and evidence for surgical intervention (marked C).

Figure 9: Cuspal enamel hypoplasia on teeth from Fewston.
The location of the Coach Lane and Fewston sites.

295x339mm (72 x 72 DPI)
The age-at-death distribution of the non-adults from Fewston and Coach Lane.

338x190mm (96 x 96 DPI)
Femoral diaphyseal lengths and cortical thickness from Fewston and Coach Lane plotted against dental age as a proxy for known age and modern comparative data.

254x190mm (96 x 96 DPI)
Pubertal stages in Coach Lane (n = 11) and Fewston (n = 24) males and females plotted against dental age.

254x190mm (96 x 96 DPI)
Vertebral body height for C5 and C6 plotted against dental age.

254x190mm (96 x 96 DPI)
Lytic lesions penetrating the right orbit of skeleton 223

274x205mm (96 x 96 DPI)
Lytic lesion and new woven and lamellar bone on the left frontal bone of skeleton 223

488x365mm (96 x 96 DPI)
Frontal and parietal bone of skeleton 157 with perimortem trauma (marked B), a radiating fracture (marked A) and evidence for surgical intervention (marked C)

432x356mm (96 x 96 DPI)
Cuspal enamel hypoplasia on teeth from Fewston

298x224mm (96 x 96 DPI)