X-ray Study of Human Dental Tissues Affected by Erythroblastosis Fetalis

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DOI: 10.1177/0022034515580987
License: Other (please specify with Rights Statement)

Document Version
Peer reviewed version

Citation for published version (Harvard):

Link to publication on Research at Birmingham portal

Publisher Rights Statement:
Version of Record available at: http://dx.doi.org/10.1177/0022034515580987
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Eligibility checked October 2015

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| Keywords:         | Biomaterial(s), X-ray Crystallography, Microscopy |

**Abstract:**

Numerous diseases are known to cause microstructural alteration of dental tissues structure. One type, in particular, is associated with neonatal jaundice and circulation of bilirubin in blood at high concentration due to increased haemolysis (e.g. in conditions such as erythroblastosis fetalis (EF), septicemia, biliary atresia and other causes of hyperbilirubinaemia). In those conditions, the products of the catabolism of haemoglobin end up deposited in various tissues, including teeth, where they can present clinically as visibly stained brown/green teeth. There is almost no information on the nature or extent of the structural changes taking place in these conditions. Here, advanced non-destructive wide angle synchrotron X-ray scattering (WAXS) techniques combined with scanning microscopy methods were used to investigate for the first time the ultrastructure of the dental hard tissues in an archival case of intrinsically pigmented green teeth. Despite no obvious elemental variation across the pigmented tissue region, the high resolution crystallographic properties probed by WAXS revealed an ultrastructural variation (orientation, particle size and lattice parameter of hydroxyapatite crystallites) associated with a pigmentation line in dentine, and with a distinct neonatal line in enamel.
X-ray study of human dental tissues affected by *erythroblastosis fetalis*

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Word count:

Abstract word count: 182
Total word count: 3147
Total number of figures: 5
Number of references: 18

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Keywords:
bilirubin, tooth, pigments, dental enamel, dentine, neonatal jaundice
Abstract

Numerous diseases are known to cause microstructural alteration of dental tissues structure. One type, in particular, is associated with neonatal jaundice and circulation of bilirubin in blood at high concentration due to increased haemolysis (e.g. in conditions such as erythroblastosis fetalis (EF), septicemia, biliary atresia and other causes of hyperbilirubinaemia). In those conditions, the products of the catabolism of haemoglobin end up deposited in various tissues, including teeth, where they can present clinically as visibly stained brown/green teeth. There is almost no information on the nature or extent of the structural changes taking place in these conditions. Here, advanced non-destructive wide angle synchrotron X-ray scattering (WAXS) techniques combined with scanning microscopy methods were used to investigate for the first time the ultrastructure of the dental hard tissues in an archival case of intrinsically pigmented green teeth. Despite no obvious elemental variation across the pigmented tissue region, the high resolution crystallographic properties probed by WAXS revealed an ultrastructural variation (orientation, particle size and lattice parameter of hydroxyapatite crystallites) associated with a pigmentation line in dentine, and with a distinct neonatal line in enamel.
Introduction

Dental appearance and structure can be affected by substances circulating in blood during odontogenesis, e.g. fluorine (dental fluorosis), tetracycline, porphyrins (congenital erythropoietic porphyria) and bilirubin (in erythroblastosis fetalis (EF) (Marsland and Gerrard, 1953), septicemia (Swann and Powls, 2012), biliary atresia or hyperbilirubinaemia (Amaral et al., 2008)). Sometimes, these changes are accompanied by enamel hypoplasia (e.g, in kernicterus cases (Forrester and Miller, 1955)).

Intrinsic bilirubin tooth staining without obvious enamel hypoplasia presents clinically as a green/brown/yellow crown discolouration and a history of neonatal jaundice (especially in EF). Green teeth associated with neonatal jaundice were first reported in 1912 (Langmead, 1912; Thursfield, 1912). Since then, few reports were published on the histology of “green teeth” (Carrillo et al., 2011; Marsland and Gerrard, 1953). Typically, unstained sections reveal accumulated pigment in dentine forming a chronological line running parallel to the incremental lines. In EF, the pigmentation marks the time when the dentition was formed, therefore appearing at some distance from the dentine-enamel-junction (DEJ), depending on the tooth developmental stage. The chemical composition of these pigmented bands and the local structure of dentine and enamel have not been thoroughly characterized and very few papers discuss the possible processes involved (Bevis, 1956). Here we investigate dental ultrastructure to clarify to what extent tissue formation is affected by bilirubin (or biliverdin) (Bevis, 1956) in blood during the jaundice episode.

2. Materials and Methods

The green tooth sample used here is an unstained ground section of an upper left
first deciduous molar from the Oral Pathology slides archive, the School of Dentistry, University of Birmingham (existing holdings, HTA license #12313). The sample originates from Case 2 of a previous study (Marsland and Gerrard, 1953) with a known history of EF and neonatal jaundice lasting 8 weeks. At least 7 teeth were pigmented at eruption and several extractions were done on two occasions over 14 months. All extracted teeth showed dentinal staining, with varying intensities.

To facilitate manipulation, the sample was mounted between two plastic plates leaving a 5mm diameter window exposing the region of interest (ROI) (Fig. 1a) whilst preventing other unwanted interference. Light microscopy inspection revealed a brownish dentinal line (~25µm thickness, Fig. 1b) running at 80µm to 220µm from the DEJ. Another band in enamel (30-40µm thick) appeared in places as a double line coincident with the so-called neonatal line. SEM imaging using back-scattered electrons detector was performed on the ROI (Fig. 1c). Previously reported SEM studies of pigmented enamel in shrews’ teeth revealed structural changes (Dumont et al., 2014). However, no obvious structure change (electron density or elemental composition) in the enamel or dentine lines were detected.

Chronologically, the change in metabolism leading to formation of the neonatal line (i.e. at birth) coincides with the time of neonatal haemolysis and jaundice in EF, posing uncertainties over the nature of structural enamel features. For this reason, the, sample was examined using a Zeiss LSM700 confocal microscope (10x objective) and 555nm monochromatic laser light excitation. The sample showed auto-fluorescence in the form of a bright line (Fig. 1f) corresponding to the pigmented band in dentine (Fig. 1e), while no obvious bright line was visible in enamel.

<< Fig.1 here >>
2.1. Micro-focus X-ray diffraction analysis

Wide angle X-ray scattering (WAXS), an X-ray diffraction technique requiring minimal sample preparation, allows non-destructive analysis of the crystal lattice structure, microstructure and strain. The position, amplitude and width of the diffraction peaks contain statistical information about the sample structure. Since each peak corresponds to a certain family of lattice planes within a certain crystallographic phase, quantitative diffraction pattern analysis can be used to identify crystallographic phases and quantify structural parameters.

Micro-beam X-ray diffraction was performed on B16 beamline at Diamond Light Source (Oxford, UK) using monochromatic X-rays at 18 KeV. As illustrated in Fig. 2a with the z-axis oriented along the beam direction, the “X-ray Eye” detector (Photonic Science X-ray MiniFDI) was used to align the sample and collimation slits. Figure 1d shows radiographic images of the sample. The slits were narrowed down to obtain a small beam impinging on the sample ROI, and the 2D WAXS detector (Photonic Science Image Star 9000) was used to collect the WAXS patterns in the x-y plane in the global coordinates. Additionally, the beam spot was focused using a Kirkpatrick-Baez (KB) mirror pair down to 0.53×0.39µm. When evaluating the positional beam stability, a time-dependent drift of 2-3 µm was observed over 30min.

A high resolution acquisition line scan over 800 µm at 2.6 µm steps was performed along the y-axis, as illustrated by the yellow line in Fig. 1d.

2.2. Data interpretation

Quantitation of WAXS patterns provides insights into the lattice parameters, particle size, and preferred orientation of HAp crystallites. Interpretation involves converting 2-D Debye-Scherrer diffraction images \( I(q,\varphi) \) \( (q \) is scattering vector
related to the radial position, $\varphi$ is azimuthal angle) (Fig. 2b) into 1-D intensity profiles e.g. using Fit2D software package (Hammersley, 1997). Figure 2c illustrates the results of azimuthal integration given by $\hat{I}(q) = \int I(q, \varphi) d\varphi$, and radial integration given by $\hat{I}(\varphi) = \int I(q, \varphi) dq$ for the (002) peak in Fig. 2 d as described in detail previously (Sui et al., 2013; Sui et al., 2014a; Sui et al., 2014b).

Crystallographic analysis of wide angle peaks is performed with the help of Bragg’s law which establishes the relationship between crystal interplanar spacing and the scattering angle of intense reflection:

$$n\lambda = 2d_{\text{hkl}} \sin \theta$$  \hspace{1cm} (Eq. 1)

where $\lambda$ is the wavelength; $d_{\text{hkl}}$ is the strain-free spacing between planes with Miller indices $(hkl)$, $2\theta$ is the scattering angle, and $n$ is the reflection order.

The reflection of (002) plane gives information on the orientation of the hexagonal $c$-axis of HAp crystallites (CIF file 1011242, Ca$_5$ (PO$_4$)$_3$(OH), hydroxylapatite/pentacalcium tris(phosphate) hydroxide). Quantitative analysis of 1-D intensity profile of (002) peaks in Fig. 2c, (e.g. Gaussian curve fitting) reveals the variation of the lattice parameter $c$ within the line scan. The transition from enamel to dentine at the DEJ is readily identified by the steep gradient.

The peak broadening in the 1-D radial diffraction pattern (Fig. 2c) is primarily related to two microstructural factors: the average size of HAp crystallites, and the root-mean-square (RMS) microstrain. The Williamson-Hall (W-H) method (Williamson and Hall, 1953) can be used to interpret e.g. the $(00n)$ family of peaks using the equation: $B \cos \theta = \frac{k\lambda}{L} + C <\varepsilon> \sin \theta$, $(k \equiv 1, C \equiv 4)$, where $B$ is is the full-width at half maximum (FWHM) of the peak analysed, $<\varepsilon>$ is root-mean-square strain, $L$ is the particle size in the direction defined by the scattering vector, and $\lambda$ and
\( \theta \) have their usual meanings in *Bragg's law*. The (004) reflection is relatively weak in the dentine. Nevertheless, careful analysis shows that the crystal size contribution (value) dominates over the RMS microstrain effect (slope): the intercept of the Williamson-Hall plot based on the two (002) and (004) peaks equals 0.11 degrees, whilst the single (002) peak value of the W-H parameter equals 0.12 degrees. Our observations are strongly supported by the dental research literature (Egan et al., 2013). This reasoning leads us to the conclusion that the *Scherrer Equation* (Patterson, 1939) (Eq.2) can be used to quantify the mean crystallite size and how it changes using the (002) peak width analysis, as follows

\[
L = \frac{k \lambda}{B \cos \theta} \tag{Eq.2}
\]

where \( k \) is a constant close to unity weakly dependent on crystallite shape, and other symbols have their usual meanings. Due to the alignment of the \( c \)-axis in HAp in enamel and dentine, \( L \) is often referred to as *length* (i.e. the longest dimension) of HAp crystallites. Since other parameters (\( k, \lambda \) and \( \theta \)) are almost constant, the variation of the length of HAp crystallites can be derived by monitoring the variation of FWHM of (002) peak.

The DEJ transition always manifests itself as a distinct change of crystallite sizes between dentine and enamel. Interestingly, significant crystallite length changes were also found near the EF lines of interest in the dentine and enamel. Thus, WAXS pattern parameters display strong sensitivity to the structural features of dental tissue, and may shed light on the processes caused by the underlying clinical events.

WAXS is also sensitive to the orientation distribution of crystallites (texture). Instead of a complete, homogeneous circle, diffraction rings may consist of circular arcs spanning limited angular ranges, with strong intensity variation as a function of
the azimuthal angle (Fig. 2d). The azimuthal centre position of the pronounced peaks is used to define the preferred orientation direction of the (002) family of planes of HAp crystallites. By tracing the azimuthal centre position of (002) circular arcs in the 2-D patterns, the HAp c-axis orientation variation with respect to the global coordinates is obtained, as shown in Fig. 2.

<< Fig.2 here >>

3. Results

3.1. Microscopy characterization

The energy-dispersive X-ray spectroscopy (EDX) analysis of the ROI is shown in Fig. 3a. Figures 3b-e illustrate the Ca, P, O and Fe EDX mapping, with the corresponding energy spectrum. The mass fraction and atomic concentration of those elements is shown in Fig. 3f. The Fe concentration in the stained material, if present, was too low to be detected. Both bilirubin and biliverdin molecules (the likely responsible candidates for the dentine pigmentation (Bevis, 1956)) contain C, H, N and O but not Fe, (the enzyme heme-oxygenase converts the heme into Fe²⁺ and biliverdin which is subsequently converted into bilirubin).

<< Fig.3 here >>

3.2. DEJ centre determination

Given the small beam size used, the line scan resolution depends on the sample thickness (~300µm). Long time line scans (~11h, ~2min per point) can lead to drift. DEJ can be identified from the sharp transition from enamel to dentine (Fig. 4a), but its position needs to be reliably to serve as a point of reference.
To determine the precise DEJ position we employed differentiation of the transition curve (a edge detection technique commonly employed in image processing and analysis). The point of the maximum gradient (peak of the derivative curve) can be identified with the centre of the transition region. It is also fairly common to use the FWHM of the derivative curve as a measure of the transition width. This method is used routinely to calculate the cross-sectional dimensions of micro-focused beams at synchrotron beamlines (using a knife edge cross-scan), providing an accurate quantitative measure. It is important not to confuse this with the FWHM of “real” diffraction peaks discussed earlier.

By fitting the differentiated curves with a Gaussian function, the FWHM size of the DEJ was found to be 37µm (Fig. 4b and 4c). The centre of the differentiated curves was also calculated to be 487µm and 491µm (average 489µm). Thus, 489µm in Fig.4 was defined zero reference position with respect to DEJ centre. Due to difficulties in precise sample positioning and its internal geometry, the DEJ plane may not be perfectly perpendicular to the sectioning plane, nor parallel to the incident beam, causing a contribution to the apparent DEJ broadening. Taking these considerations into account, scan positions below ~425µm can be safely identified as 'pure enamel', and beyond ~550µm as 'pure dentine'.

<< Fig.4 here >>

3.3. Ultrastructural characterization

The 'peak centre', 'preferred orientation' and particle size ('peak width') derived from the diffraction patterns in enamel are shown in Fig. 5a,c,e, superimposed on Fig. 1b. The yellow horizontal line refers to the scanning line in Fig. 1d. There is visible and clear nonlinear region approximately -330µm from the DEJ centre. This appears
consistent with the optical microscope image (Fig. 1b) showing the distinct banded enamel feature ~200-350µm from the DEJ. Note that the time of high bilirubin exposure in EF coincides with the neonatal line development.

The diffraction pattern parameters labelled 'peak centre', 'preferred orientation' and 'particle size' (derived from the peak width) in dentine are plotted in Fig. 5b,d,f, also superimposed on Fig. 1b, with the horizontal scanning line shown in yellow in Fig. 1d. The stained band in dentine lies 220µm from the DEJ (Fig. 1b). The structural changes are mostly reflected in the plots of preferred orientation and peak centre position around the scan location at 211-236µm (on the stained line).

Structural parameter plots across banded features allow making deductions about the effect of high bilirubin. In Fig. 5, the DEJ lies between the two image columns, with the direction of enamel deposition to the left, and dentine to the right. In both dentine and enamel, the position of the bands corresponds to the time of disruption of tissue formation. Peak centre plots in Fig. 5a, b reveal that the overall systematic trend is for the lattice spacing of HCP lattice basal planes to decrease as the tissues grow away from the DEJ.

Fig. 5c-d show plots of the preferred orientation angle as a function of position. Both enamel and dentine exhibit localised changes in the orientation angle of crystallites.

Finally, Fig. 5e,f illustrate the variation in particle size (derived from the peak width) with the distance from the DEJ. Two opposing trends are observed: the particle size increases with distance from the junction in the enamel. At the same time, the particle size in dentine becomes smaller further away from the DEJ. However, a reversal of the respective trends is observed in both enamel and dentine within the EF bands. These result in the appearance of a local maximum of crystal size. Once again,
the trend is reversed subsequently.

In summary, detailed analysis of the variation of structural parameters across the stained band reveals localised changes consistent with other aspects of the structure. It appears that during pigment deposition, a change in the tissue mineralization process leads to temporary increase of the c-axis lattice spacing, accompanied by larger HAp crystallites produced. These changes are temporary, and the prevailing growth trends in dentine and enamel are eventually restored.

<< Fig.5 here >>

4. Discussion

A number of key conclusions can be drawn from the results.

There is a clear overall decreasing trend for the lattice spacing of HCP lattice basal planes with the distance from the DEJ. Different hypotheses to explain this phenomenon have been put forward, and a controversy regarding the underlying mechanisms persists. Analysing calcite in seashell nacre, (Pokroy et al., 2006) state that in biogenic calcite the unit cell parameters $a$ and $c$ were systematically larger than those in the non-biogenic calcite. They suggest that it is caused by lattice swelling from organic macromolecules incorporated within the crystals during biomineralization. However, there are arguments against this mechanism being active for HAp crystallites, due to their extremely small size. The incorporation of citrate and carbonate ions may take place, but presently no firm proof of this hypothesis had been found. In terms of crystal perfection, the maturation of hard tissues (Arnold et al., 2001) is thought to lead to the decrease in the crystal lattice fluctuations in bone, dentine and enamel that may be associated with changes in the content of ions such as hydroxyl, carbonate, and sodium. The formation of more perfect inorganic crystallites
is accompanied by the reduction in the lattice parameter.

Given the maturation trends described above, our results reveal that in the EF-affected bands the maturation is temporarily retarded or even reversed: local maxima of HAp interplanar lattice spacing are observed in both dentine and enamel. We surmise that disease led to a temporary decrease in the extent of mineralisation, leaving the HAp crystal lattice less perfect and more widely spaced during the haemolysis episode, followed by subsequent recovery, when the expected tissue formation and mineralisation rates were restored.

Regarding the change in the preferred orientation of crystallites and the local maximum that it exhibits at the pigmented bands, we note that underlying systematic curvature of the tubules reflecting the continuous structural make-up of dentine or the consequence of sectioning, might result in monotonic variation of orientation, but would not lead to the appearance of local maxima. We interpret such localised changes as consequence of a disruption in the normal dentinal growth and associated with the haemolysis episode. Biliary pigments in blood are a likely cause of cellular injury to odontoblasts and ameloblasts, affecting the rate of secretion and the composition of the organic matrix. This, in turn, changes the rate of advance of the tissue mineralisation front, resulting in altered in crystal growth, packing and orientation.

Our observations suggest an interesting explanation for the influence exerted by haemolysis on hard mineralized tissues growth. Analysis of multiple samples would be highly preferred, but was not possible due to the combination of extreme sample scarcity ('green teeth' are rarely available for analysis: discolouration alone causes only an aesthetic concern, and tooth extraction is not indicated) and limited availability of facility beamtime. Although these considerations preclude
generalization, we feel nevertheless that the quantitative nature of our measurements merits the presentation of the results as a hypothesis deserving further evaluation.

While no elemental composition changes were detected in dentine or enamel by SEM or EDX analysis (Fig. 1b), X-ray diffraction, revealed morphological differences at the (sub)nano-scale: particle orientation, size and lattice parameter changes in the HAp crystallites.

The morphological information from enamel reveals a correlation between the visible band and the underlying ultrastructural variation, even though there was no detectable autofluorescence after laser excitation. We conclude that the enamel band is not pigmented, but corresponds to changes in the enamel refraction index due to the local variation of crystallite morphology.

The neonatal line, invariably present in deciduous teeth and first permanent molars, represents mineralisation pattern changes associated with the physiological transition from intra- to extra-uterine life. This event coincides with the possible interference of odontogenesis by bilirubin because haemolysis in EF also occurs immediately after birth. The appearance of green pigmented teeth, in this case, is therefore a consequence of the pigment in dentine, perceived through the enamel translucency, and not enamel pigmentation. Marsland and Gerrard (Marsland and Gerrard, 1953) described a double neonatal line in some (but not all) of their EF tooth samples. Some parts of the enamel in our sample showed this feature. However, these lines in ground sections are again visible because of changes in the enamel refraction index; in reality the 'lines' are finite 3-D layers of enamel, and oblique sectioning planes on a varying thickness sections could create 'double line' effects when the layer forming the “line” is thinner than the section (see Fig 1 of (Eli et al., 1989)). It has been suggested that if bilirubin were temporarily deposited in the organic enamel matrix, it might be lost.
during mineralisation and maturation. In dentine, the presence of structural changes might be attributed to the larger proportion of retained organic content compared to enamel (Marsland and Gerrard, 1953). This is also supported by the absence of auto-fluorescence in the enamel.

5. Conclusions

The properties of distinct bands induced in dentine and enamel in EF were investigated and correlated with ultrastructural variations in crystal lattice parameters, size and orientation determined by WAXS, revealing quantitative changes associated with EF at a level of detail not possible previously. Future investigations will concern elucidating ultrastructural variations within neonatal lines in dentine and enamel and the extent to which they affect tissues’ biomechanical properties.

Acknowledgement:

AMK acknowledges funding received for the MBLEM laboratory at Oxford through EU FP7 project iSTRESS (604646). Diamond Light Source is acknowledged for providing the beam time allocation under experiment number NT5038. The authors have declared no conflict of interest.

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Figure caption

Fig. 1 Basic characterization of green tooth samples. Sample preparation of the green tooth (a) the region of interest (ROI); (b) the optical microscope image of ROI, where the pigmented lines in dentine and a distinct line in enamel are obvious; (c) Back-scattered electron image (BSE) of ROI, with no visible difference compared to (b); (d) “X-ray Eye” image of ROI, used to identify the line scan for X-ray diffraction; (e) the optical microscopy image of the line in dentine; (f) fluorescence of the line in dentine revealed by confocal microscopy of the same region as shown in (e).

Fig. 2 (a) Micro- (Nano-) focus Experiment set-up in B16 beamline, Diamond Light Source. KB mirrors were used to focus the incident X-ray beam. Sample was mounted on the sample stage between the KB mirrors and WAXS detector at the focus point, which was highlighted in the red oval. WAXS data interpretation from (b) A representative enamel 2-D WAXS pattern to 1-D profile by (c) azimuthal and (d) radial integration of (002) peak (Gaussian fit indicated in red, background shown in green).

Fig. 3 EDX analysis of ROI. (a) The SEM image of ROI; (b)-(e) respective element mapping of Ca, P, O and Fe in ROI; (f) the energy spectrum covering all element lines, together with the table showing the mass percent and number density of different elements.

Fig. 4 (a) (002) scattering angle peak centre (blue) and (002) peak width (black) plot from enamel to dentine across the DEJ; Differentiation (blue) of (b) peak width (black) and (c) peak centre (black) superimposed with the original curve; Gaussian peak fitting curves (red) for peak width and peak centre are also shown in (b) and (c) to determine the FWHM.

Fig. 5 Correlations between structural parameter variations in the pigmented tooth section for enamel (-489 to -64µm from DEJ, left column) and dentine (61-311µm from DEJ, right column): (a) and (b) peak centre (red); (c) and (d) preferred orientation (light blue), and (e) and (f) particle size (black). The background contains
magnified optical images of the enamel and dentine microstructure, and the yellow line denotes the position of the X-ray scan. (The standard deviation is about 4 orders of magnitude lower than the fitted value, e.g. the $d$-spacing of the peak centre fit is given by $3.477\pm1.14E-05$ and therefore scatter bars are not included in the plots).
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