

ORIGINAL RESEARCH

Phenotypic Properties of Collagen in Dentinogenesis Imperfecta Associated with Osteogenesis Imperfecta

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Salwa Ibrahim 1,* Adam P Strange^{2,*} Sebastian Aguayo (1)^{2,3} Albatool Shinawi Nabilah Harith Nurjehan Mohamed-Ibrahim 1 Samera Siddiqui² Susan Parekh Laurent Bozec⁴

¹Department of Paediatric Dentistry, UCL Eastman Dental Institute, University College London, London, UK; ²Department of Biomaterials and Tissue Engineering, UCL Eastman Dental Institute, University College London, London, UK; ³School of Dentistry, Faculty of Medicine, Pontificia Universidad Católica de Chile, Santiago, Chile; ⁴Faculty of Dentistry, University of Toronto, Toronto, Canada

*These authors contributed equally to this work

Introduction: Dentinogenesis imperfecta type 1 (OIDI) is considered a relatively rare genetic disorder (1:5000 to 1:45,000) associated with osteogenesis imperfecta. OIDI impacts the formation of collagen fibrils in dentin, leading to morphological and structural changes that affect the strength and appearance of teeth. However, there is still a lack of understanding regarding the nanoscale characterization of the disease, in terms of collagen ultrastructure and mechanical properties. Therefore, this research presents a qualitative and quantitative report into the phenotype and characterization of OIDI in dentin, by using a combination of imaging, nanomechanical approaches. Methods: For this study, 8 primary molars from OIDI patients and 8 primary control molars were collected, embedded in acrylic resin and cut into longitudinal sections. Sections were then demineralized in 37% phosphoric acid using a protocol developed in-house. Initial experiments demonstrated the effectiveness of the demineralization protocol, as the ATR-FTIR spectral fingerprints showed an increase in the amide bands together with a decrease in phosphate content. Structural and mechanical analyses were performed directly on both the mineralized and demineralized samples using a combination of scanning electron microscopy, atomic force microscopy, and Wallace indentation.

Results: Mesoscale imaging showed alterations in dentinal tubule morphology in OIDI patients, with a reduced number of tubules and a decreased tubule diameter compared to healthy controls. Nanoscale collagen ultrastructure presented a similar D-banding periodicity between OIDI and controls. Reduced collagen fibrils diameter was also recorded for the OIDI group. The hardness of the (mineralized) control dentin was found to be significantly higher (p<0.05) than that of the OIDI (mineralized) dentine. Both the exposed peri- and intratubular dentinal collagen presented bimodal elastic behaviors (Young's moduli). The control samples presented a stiffening of the intratubular collagen when compared to the peritubular collagen. In case of the OIDI, this stiffening in the collagen between peri- and intratubular dentinal collagen was not observed and the exposed collagen presented overall a lower elasticity than the control samples.

Conclusion: This study presents a systematic approach to the characterization of collagen structure and properties in OIDI as diagnosed in dentin. Structural markers for OIDI at the mesoscale and nanoscale were found and correlated with an observed lack of increased elastic moduli of the collagen fibrils in the intratubular OIDI dentin. These findings offer an explanation of how structural changes in the dentin could be responsible for the failure of some adhesive restorative materials as observed in patients affected by OIDI.

Keywords: dentin, collagen, dentinogenesis imperfecta, demineralisation, dentistry, atomic force microscopy

Correspondence: Laurent Bozec Faculty of Dentistry, University of Toronto, 124 Edward Street, Toronto ON M5G 1G6, Canada Email L.bozec@utoronto.ca

Introduction

Dentinogenesis imperfecta (DI), or opalescent teeth disease, is an inherited disorder in which dentin quality is affected. DI is a localized mesodermal dysplasia affecting both primary and permanent dentition. The first classification of DI was originally proposed by Shields¹ in 1973 and was further commented by Wiktop² in 1975. A more recent report³ reviewed the evolution in the etiology and nomenclature system of this condition to clarify the relationship between DI and osteogenesis imperfecta (OI). According to Shields, DI is classified into three main groups, Type I (DI associated with OI (osteogenesis imperfecta), an autosomal dominant trait), Type II (also an autosomal dominant trait, but with no OI association), and Type III (thought to have occurred as variable of DI Type II). The differential diagnosis between types I and II is often challenging, possibly due to the rarity of the conditions. The prevalence of DI was recently found to be as few as 1 in 45,000 in Swedish children and adolescents, although ORPHANET (ORPHA:71267) cites the prevalence of DI to be <1/1,000,000. DI type III, also called "Brandywine isolate", is described as an extremely rare form with highest incidence of 1 in 15 in the sub-population of Maryland.⁵

Clinical and radiographic representation of DI varies greatly between, and within types. In DI type I (also denoted in this manuscript as OIDI), primary predecessors are usually more severely affected than permanent teeth. In DI type II, both dentitions are equally affected. Radiographically, DI type II and primary dentition in DI type I, show pulp obliteration, unlike type III which is characterized by thin dentin and large pulp chambers. DI type III teeth are frequently referred to as "Shell Teeth".6 Furthermore, DI affected teeth will often have short roots and bulbous crowns, with a significant constriction at cement-enamel junction.7 Intraorally, teeth are described as being translucent with a discolouration hue ranging from yellowish brown to greyish blue, and an increased susceptibility to wear and fracture. It is believed that the cause of frequent tooth wears and enamel breakage may be due to defective EDJ (Enamel Dentin Junction).8

The primary cause of the autosomal dominant DI type I (OIDI) is mutations in *COL1A1* or *COL1A2* genes. These genes encode for the α -1 and α -2 polypeptide chains of collagen type I. These polypeptides chains are the basic building blocks of the collagen molecules, and therefore any alterations in the structural integrity of the molecules may trigger abnormal collagen fibril formation. Genetic mutations causing OIDI are often linked to the substitution of the glycine (Gly) residue present in the α -1 and/or α -2 polypeptide chains by other larger residues such as serine. Being the smallest amino acid within the α -1 and α -2 polypeptide chains, Gly

often sits within the centre of the triple helical structure of collagen, allowing the helices to wind together tightly. The Gly residue is also involved in the stabilization of the triple helix through the formation of hydrogen bonds with the internal bound water. ¹⁴ Thus, any changes in identity of the amino acid replacing Gly, and the location of that substitution, will alter the ability of the collagen triple helix to form correctly, and impact the pathology of osteogenesis imperfecta (OI). For example, substitutions of glycine with serine (polar amino acid) in the α -1 collagen chains have been found to cause severe types of OI in bones. ¹⁵ Similarly, it has been established that patients with glycine substitutions will have some degree of OIDI comorbidity. ^{16–19}

While the effect of OI on bones has been well reported and studied, there is a lack of reported outcomes for OIDI in teeth. In this study, we present recent findings in the structure and properties of OIDI-affected dentin and especially in its collagen component. Using a combination of imaging and mechanical approaches, we present a quantitative and qualitative report of the phenotype of DI associated with OI in teeth.

Materials and Methods

Tooth Collection and Preparation

Ethical approval for this study was obtained from the National Health Services Research Ethics Committee (registration reference number 11/LO/0777). Following written informed consent, 8 primary first and second molars from patients with a confirmed genetic diagnosis of OI (OIDI), and 8 primary first and second molars from healthy control individuals were collected over a period of 24 months, at the University College London Eastman Dental Hospital, London, UK. All patients with OI underwent genetic testing at Great Ormond Street Hospital (London UK). Those included in this study had all a confirmed diagnosis of mutations in COLA1 gene and had not been treated with bisphosphonate therapy. Figure 1 shows the vertical bitewing radiographs of one of the patients with OI Type I and OIDI in the primary dentition, showing characteristic features of pulpal obliteration and bulbous crowns. All teeth were extracted as part of the patient's treatment plan. It is worth noting that due to the rarity of the condition, accessing OIDI teeth is one of the major limitations in any study of this disease, and as a result, this study could not be powered.

Following extraction, the teeth were initially stored in a 70% ethanol solution for up to 5 days at room temperature, before being debrided from the remaining soft tissues, and finally stored in a 0.1% thymol solution at 4°C until required

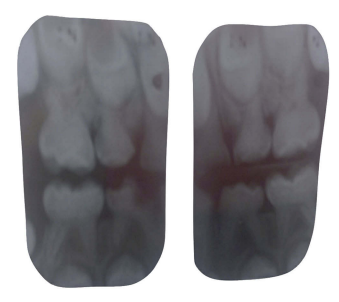


Figure 1 Vertical bitewing radiographs of patient with OIDI in the primary dentition, showing characteristic features of pulpal obliteration and bulbous crowns.

for the study (storage did not exceed 2 months). The teeth were then sectioned longitudinally (~1–1.5 mm thick) using a diamond microtome (Struers, Accutom-50, Struers Ltd., Solihull, West Midlands, UK). Finally, the sections were then sonicated in Ultra-High Quality (UHQ) water for 30 s.

Dentin Hardness

To evaluate the hardness of the dentin, the sections were then mounted in a Wallace indenter (H.W. Wallace, Croydon, England). The sections were held in position on the sample platform using a malleable putty to prevent any sliding of the sections during the measurements. By measuring the indentation depth of the indenter (40µm pyramidal shape), one can obtain the Vickers Hardness Number for the samples using the following equation:

$$VHN = \frac{applied\ force\ (F)}{surface\ area} = 1854.4 \times \frac{F}{d^2}$$

where F is applied load (300 g) and d is the mean diagonal of indentation (in μ m). All measurements were performed by maintaining the load for 15 s before release. For each tooth, a total of 10 individual measurements were taken within the crown region of each section. The VHN values were plotted as boxplots focusing on the two groups, control vs OIDI.

Controlled Exposure of the Collagen Network

Following hardness measurements, sections were then demineralized in a solution of 37% phosphoric acid placed in an

ultrasonic bath. The exposure time of the section to the acid was closely monitored by Attenuated Total Reflectance-Fourier Transform Infrared Spectroscopy (ATR-FTIR) using a Tensor 27 spectrometer (Bruker, Coventry, UK) equipped with a GladiATR (PIKE Technologies, Madison, USA). Following time-controlled acid-exposure, the sections were then washed with UHQ water, before being placed in 6.5% sodium hypochlorite (ReAgent, Liverpool, UK) for 5 s to remove any smear layer resulting from the acid etching process. Finally, the sections were washed again in UHO water before being air-dried for 24 hrs prior to any measurements. This approach was successfully applied to other studies.¹⁷ All infrared spectra were recorded at 128 coadditions and 16 cm⁻¹ spectral resolution over the spectral range 4000-700 cm⁻¹. The spectra were baseline corrected before being normalized using the Amide I band at 1650cm⁻¹. The potency of the demineralization protocol was assessed by calculating the normalised ratio of spectral intensities of the phosphate band over the Amide I band.

Imaging the Collagen Network (by AFM)

No further sample preparation was required for atomic force microscopy (AFM) imaging. Individual sections were mounted onto glass-slides using double-sided tape and kept in ambient conditions. AFM imaging was performed using a Dimension 3100 Atomic Force Microscope (Bruker, Santa Barbara, CA, USA). Imaging was carried out with an MNSL AFM probe (k=0.03-0.1 N/m) (Bruker, Santa Barbara, CA, USA) in contact mode in air. Set point and gain values were adjusted in-situ during scanning for image optimization. Images were obtained at 512×512pixels (image size: 10×10μm² or 2×2μm²) with an average scanning rate of 1 Hz. Images were obtained and analysed using Nanoscope Analysis Software (Version v14, Bruker, Santa Barbara, CA, USA) in order to measure both fibrils diameter (cross-section profile analysis) and D-banding periodicity (line-profile analysis along the fibrils length).

Single Fibrils Collagen Nanomechanics (by AFM)

For the mechanical measurements carried out on the exposed collagen matrix, a Nanowizard AFM (JPK, Berlin, Germany) mounted on an inverted optical microscope (Olympus IX71, Olympus, Japan) was used. Mechanical measurements were performed using REFSPA tips (k=3 N/m) (Bruker, Santa Barbara, CA, USA) in ambient conditions (prior to the SEM sample processing). For all measurements, both the

indentation load (5 nN) and loading rate were kept constant. Prior to acquiring any force-distance curves (FD), a lowresolution image (256 \times 256 pixels -5×5 µm²) was acquired in contact mode with the same probe to ensure that the area being characterized mechanically presented sufficient exposed collagen matrix. FD measurements were then conducted on individual collagen fibrils presenting a D-banding periodicity by choosing the centre of the D-banding (overlap region) as indentation site on the low-resolution AFM image in both the intra-tubular and peri-tubular dentin. At least three series of indentation curves were taken in each location (7) for each sample, with at least three sections per condition analysed, for a minimal of 350 force curves per location. FD curves were analysed on proprietary software (JPK, Germany) and the Young's modulus was calculated using the Sneddon model.²⁰ All force curves were analysed using the JPK Data Processing Software v.5.1.8 (JPK Instruments, Germany).

Imaging the Collagen Network (by SEM)

Following AFM characterization, the sections were fixed for 24 hrs in 3% glutaraldehyde (Agar Scientific, UK) and were then dehydrated using ethanol series, before being thinly coated with gold-palladium using a Polaron E5000 Sputter Coater (Quorum Technologies Ltd, East Sussex, UK). Scanning Electron Microscopy (SEM) imaging was performed using an FEI XL30 FEG-SEM (FEI, Eindhoven, Netherlands) using a 5 kV accelerating voltage. A total of 7 locations were selected at random for each section and imaged at different magnifications.

Statistical Analyses

One-way ANOVA (Kruskal–Wallis test) was conducted to assess the significance (p<0.05) for the mechanical and structural parameters presented as boxplots (median and 95% CI) between the control and OIDI groups. For the nanomechanics analyses, histograms and median values were obtained for the Young's moduli. All data were plotted and processed using Origin software (OriginLab Corp, Northampton, MA, USA).

Results and Discussion

Systematic Demineralization of Dentin Surface

Dentin is the mineralized matrix situated within the core of the tooth. Structurally, it can be characterized by multiple tightly packed dentinal tubules found along its entire thickness. After dentinogenesis is completed, 70% by weight of

dentin is mineralized hydroxyapatite, with the organic content accounting for 20% of the matrix, and the remaining 10% being water. Fibrillar collagen makes up the organic component of dentin, 85% type I collagen, and 15% types III and V collagen. To date, efforts in demineralising dentin surfaces to expose collagen have had mitigated success, in terms of repeatability. This can be explained by the fact that dentin is a heterogeneous tissue, and as such, it reacts differently when exposed to demineralisation agents. However, to access the ultrastructure of the collagen scaffold present within such a mineralized matrix, it is essential to demineralize this matrix, but without altering the morphology and chemistry of the collagen. To control the degree of demineralisation of the dentin surface, and to maintain the clinical relevance of our approach, we decided to use 37% phosphoric acid as it is commonly used in most clinical dental etching protocols. Other acids such as citric, lactic or nitric acids, and chelation solutions such as EDTA, have all been used to remove minerals from bone or dentin samples. Unfortunately, collagen crosslinks are very susceptible to acid attack and will be cleaved when exposed to a sustained acidic environment. Although chelation agents would not affect these crosslinks, their demineralization action is a slow process and is therefore not clinically relevant.

Figure 2A shows the normalized FTIR data for control (dotted line) and demineralized dentin (full line) after 10 s exposure to phosphoric acid. In the FTIR spectrum of the native control dentin, the presence of mineral phase of the dentin, hydroxyapatite (HA), dominates the spectral fingerprint. The most characteristic chemical groups in the FTIR spectrum of HA are PO₄³⁻, OH⁻ and CO₃²⁻. PO₄³⁻ group presents intensive IR absorption bands at 560 and 600 cm⁻¹ (not shown) and at 1000–1100 cm⁻¹. The Amide A band is relatively wide, from 3600 to 2600 cm⁻¹, with an explicit peak at 3570 cm⁻¹. CO₃²⁻ group forms weak peaks between 870 and 880 cm⁻¹ and more intensive peaks between 1460 and 1530 cm⁻¹. In contrast, in the FTIR spectrum of the demineralized control dentin, the presence of protein phase of the dentin, collagen, dominates the spectral fingerprint. Collagen infrared fingerprint is defined by several specific absorption bands defined as Amide A, I, II and III. The Amide A band is the broad peak present in the spectrum between 3600 and 3200 cm⁻¹ which relates to N-H stretching vibrations. The amide bonds present in the collagen amino-structure contributes significantly to the FTIR spectrum through both stretching and bending vibrations of the carbonyl (C=O) at 1650 cm⁻¹ (amide I band),

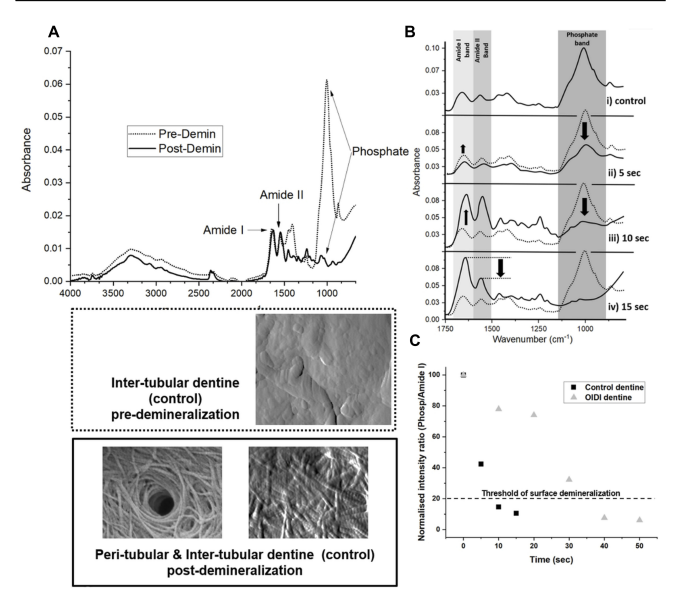


Figure 2 FTIR spectra (A) averaged composite FTIR spectra of control teeth pre-demineralisation (dotted line) and post-demineralisation (solid line). Images (SEM and AFM) present the control dentin surface pre- and post-demineralisation. (B) Infrared spectrum of the control (i) dentine, after 5 s (ii), 10 s (iii) and 15 s (iv) demineralisation. (C) Plot of pf the normalized intensity band ratio (Phosphate/Amide I) as function of the demineralisation time for control and OIDI dentin; dotted line marks the empirical threshold below which the surface is considered fully demineralised.

the N–H and C–N bonds at 1550 cm⁻¹ (amide II band) and at 1230 cm⁻¹ (amide III band). By comparing the relative absorbance intensity of the PO₄³ band relatively to that of the Amide I and II, it is possible to assess the level of demineralization in the dentin following an acid exposure.

Figure 2B shows a series of ATR-FTIR spectra (notnormalized) recorded on the same control dentin section exposed to an increased exposure of phosphoric acid from 0 s up to 15 s. In this figure, one can appreciate the constant decrease of the PO₄³ band already after 5 s exposure to phosphoric acid. After 10 s, this band has almost completely disappeared whereas the collagen Amide I and II doublet become more and more apparent. If the exposure to phosphoric acid is increased up to 15 s, the intensity of the Amide II starts decreasing suggesting that the acid may be starting to degrade the collagen itself. The gradual decrease in intensity of the PO₄³ band confirms that the mineral phase present within the first 2 µm from the surface of the dentin section is being removed by acid exposure. One of the reasons for choosing ATR-FTIR over other spectroscopic techniques, such as Raman for example, is that the depth of sampling in ATR is governed by the evanescent infrared waves emerging from the diamond window. The depth of penetration of these waves in

the sample is only a few microns (<2 µm) moderated by the refractive indices of both the sample and diamond window. Thus, our approach ensures that we are only sampling a surface demineralization rather than a full thickness demineralization. Finally, it is difficult to assess the impact of this protocol onto the phosphorylated non-collagenous proteins as their individual fingerprints have not yet been established using ATR-FTIR spectroscopy. These are likely to combine both the Amide I and the PO₄³⁻ bands but further studies are required to prove this.

By calculating the normalised ratio between the intensities of the phosphate band over the Amide I band as a function of acid-exposure time, one can appreciate the increase in the exposure time (10 s for control and 35 s for OIDI) required to observe a net reduction of the phosphate band presence in the infrared spectra. To date, there have been no reports on the effects of mineralization of OIDI on dentin which may explain for the longer required demineralisation time. However, studies have shown increased mineralization to be a characteristic feature of OI bone achieved by densely packed mineral particles as a result of defective collagen, leading to high fragility and a brittle glass-like nature. 21-23 As a result, we empirically defined a threshold value (20%) for the calculated ratio of the two fingerprint bands to mark when surface demineralisation has been achieved when monitored by ATR-FTIR. The demineralisation of all the sections (control and OIDI) were carried up to the point where the normalised ratio of the two fingerprints (Phosphate and Amide I) fell below this threshold value. This ensured that our systematic demineralization of the control and OIDI dentin surface became reliable even when using 37% phosphoric acid as shown in Figure 2C.

Morphological Assessment of Demineralized Dentin Surface

Although the clinical presentation of OIDI varies from patient to patient, OIDI-affected primary teeth will often have short roots and bulbous crowns, with a significant constriction at the cement-enamel junction (CEJ) and pulpal obliteration. The primary dentition is normally more severely affected than the permanent teeth.²⁴ Although these teeth also present an increased susceptibility to wear as a result of lower enamel hardness and resistance to attrition, it is understood that structural changes in the dentin may explain the failure of some adhesive restorative materials.⁸ However, there is currently no reports detailing

such structural changes. In dentin, it is the composite nature of the mineralised collagen scaffold that provides the tooth with its mechanical integrity. Forming an ordered structure, the collagen fibrils present in dentin are arrayed in a consistent distribution of interrupted striations and thinner fibrils, thus forming an overall collagen network. The remaining non-collagenous part of dentin is composed of non-structural proteins. Any alteration of this ordered array at the mesoscale, or fibrillar ultrastructure at the submicroscale, may result in an increased susceptibility to fracture as reported elsewhere.⁸

Mesoscale Ultrastructure of Demineralised Dentin

Figure 3 shows representative images of both the demineralised control dentin a) and demineralised OIDI dentin at various magnifications. In the case of the control sample, one can observe the regular distribution of dentinal tubules across the surface of the samples (at the lowest magnification image 2-a-i). These tubules are also all unblocked as expected following the NaOCl step in our protocol. Imaging the demineralized control dentin sample at higher magnification, it is possible to observe the exposed collagen matrix on the surface of the sample, both around and inside the tubules as presented in Figure 3A-ii, iii and iv. In these images, clear interstitial gaps between the fibrils can be observed suggesting that the demineralization protocol used was effective in removing the mineral matrix that surrounds the collagen matrix. At the higher magnification (2-a-iii & iv), it is possible to evaluate whether the demineralization protocol may have affected the topology of the collagen fibrils. As one can clearly identify the collagen D-banding periodicity without any signs of fibrils unwinding or swelling, it is likely that our controlled demineralisation has not affected the collagen fibrils morphology.

In the case of the OIDI, as presented in Figure 3B-i, very few tubules are present at the surface of the sample. Those present are more ovoid in shape which could be due to the angle at which the sections were cut. Figure 3B-ii shows a "nesting" effect of tubules, which can only be observed in OIDI samples. This nesting or branching of the tubules was also observed histologically by Andersson et al.²⁵ In their study, they also reported variations in the width of the dentin tubules, presence of hyaline dentin void of dentin tubules, layering of dentin, and presence of cell lacunae and duct-like structures in the body of the dentin. This corroborated with our findings, including the diameter of tubules which appears to be also less uniform in our images. We

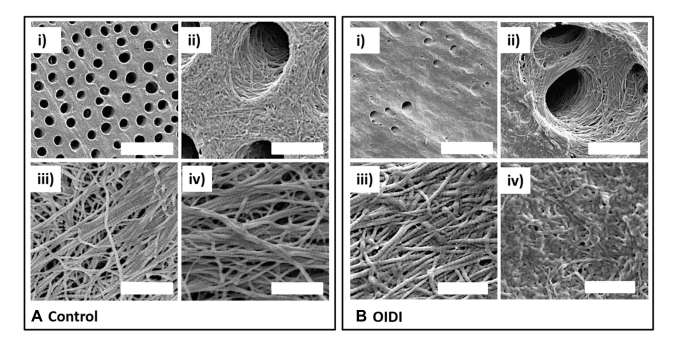


Figure 3 (A) SEM images of control dentin: i) pre-demineralisation (scale bar 20μm): regular dentinal tubule arrangement, tubules are open and uniform in appearance; ii) post demineralisation (scale bar 2μm): higher magnification image of tubules, collagen scaffold is visible inside the tubules and in the surrounding space; iii) and iv) post demineralisation (scale bar 800nm): higher magnification images showing regular D-banding periodicity of collagen fibrils. (B) SEM images of OIDI dentin: i) predemineralisation (scale bar 20μm): tubules varying in size and partially blocked or occluded; ii) post demineralisation (scale bar 2μm): "inesting" of several tubules can be seen within a larger tubule, collagen can be seen in the tubule (scale bar 2 μm), iii) post demineralisation (scale bar 800nm): higher magnification image showing regular D-banding periodicity of collagen fibril; iv) post demineralisation (scale bar 800nm): higher magnification image showing collagen fibril coalescence.

should, however, be careful with this outcome as tooth sectioning angle may affect the cross-sectional uniformity of the dentine tubules. Although the OIDI samples were subjected to the same sonication protocol as used for the control samples, partial dentinal tubules occlusions can be also observed in Figure 2B-i. We suggest that a tubule is partially occluded when we can observe its outline, but its void is filled with non-specific compounds. At this stage, it is difficult to evaluate if the presence of these partial occlusions is phenotypically relevant to OIDI as this finding has not yet been reported elsewhere to date.

Contrary to the control samples, two types of morphological collagen scaffold organisation could be observed at higher magnification. Figure 3B-iii presents a very uniform collagen scaffold very reminiscent to that of the control dentin. The D-banding periodicity is very apparent on all collagen fibrils with no obvious sign of structural damage. However, in Figure 3B-iv, the collagen network has been significantly altered and the fibrils network is no longer homogeneous. In fact, the topology of the fibrils suggests a significant level of intro-fibrillar disorganising with notable fibrillar diameter swelling and constriction. Sporadic presence of D-banding periodicity can be observed on a few sparse fibrils, but overall the fibrils coalesce together

amorphously. The prevalence of such morphologically compromised area over the entire OIDI dentin sections was not quantified and therefore should be carefully considered.

Quantitative Assessment of Dentin Tubules

For the control dentin, a total of 1104 tubules were identified over 175 regions (20×20um² each), averaging 6.3 tubules per area. For the OIDI dentin, a total of 223 tubules were identified over 137 regions (20×20µm² each), averaging 1.6 tubules per area. Compared to the OIDI sample, the native control samples presented on average over 5 times more tubules per area investigated although the measurements were performed at the same locations on all teeth. Here we can define dentine density at a ratio between dentine (as a continuous bulk matrix) and the number of tubules present per unit area. This clear difference in the tubules number per unit area suggests that the OIDI dentin is much denser than the control dentin. This could be linked to either a diminished odontoblastic activity or increased mineralization towards tubule occlusion in OIDI patients. From a mechanical point of view, this apparent density should provide the OIDI dentin with a greater resistance to fracture. A significant difference in tubule diameter between the 2 groups could also be recorded as presented in Figure 4A. In the case of the native control Ibrahim et al Dovepress

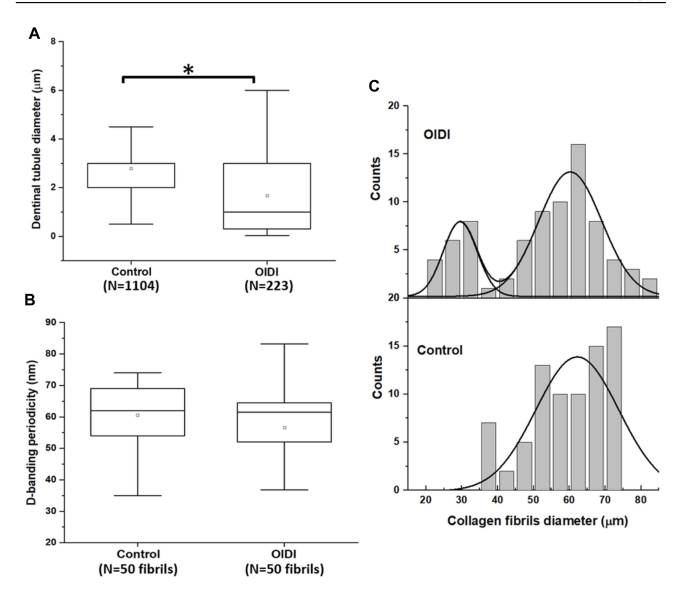


Figure 4 Graphical presentation of the nanoscale phenotypic properties (structural) for control and OIDI dentin: (A) boxplots of the measured dentinal tubules diameter (*significance p<0.05). (B) Boxplots of the measured collagen fibrils D-banding periodicity (no significant differences were observed). (C) Fitted histograms of the collagen fibril diameter with bimodal distributions (Gaussian) for OIDI collagen (R²>0.9) and for control collagen (R²>0.8).

sample, the tubule diameter was found to be narrowly distributed, $2.75\pm0.72~\mu m$ in diameter, whereas in the case of the OIDI samples, not only does the median diameter decrease to $1.67\pm0.11~\mu m$ but the distribution of the diameter is also increased moderately when compared to that of native control dentin. Statistical analysis using Kruskal–Wallis ANOVA shows a significant difference (P<0.05) between the tubule diameters in OIDI and control, with the control tubules being larger. Since the formation of tubules is directly influenced by both the odontoblasts as to form the collagen matrix and the odontoblast processes around which the dentine matrix mineralizes, it is plausible to hypothesize that the variation in dentinal tubules diameter could be directly linked an alteration

to the collagen scaffold deposited. This is yet to be investigated as much needs to be done to elucidate the influence between odontoblastic activity and OIDI prevalence.

Quantitative Assessment of Collagen Fibrils Structure

The ultrastructure of the exposed matrices in both native control and OIDI demineralized dentin was also investigated by using higher magnification electron microscopy and AFM imaging as presented in Figure 3A-iii and B-iii. In both control and OIDI images, collagen fibrils can be clearly observed presenting a well-defined D-banding periodicity, in a regular, ordered state. D-banding periodicity

is an intrinsic topographical feature of native collagen fibrils. 26-28 Together with an overall regular fibrillar topography, this D-banding can be used as a marker for intact collagen as both the shape of the fibrils, and the width of the D-banding. The lack of D-banding may be the result of an internal structural and molecular disorganisation, enzyme-mediated digestion, acid or thermal denaturation. Using these images, it was possible to measure the distribution of D-banding and collagen fibril diameter in both the series of samples. As shown in Figure 4B, both native control and OIDI presented a unimodal distribution with medians of (63.2 ± 0.2) nm (N=50 fibrils) and (62.3 ± 0.3) nm (N=50 fibrils), respectively. Kruskal-Wallis testing confirmed that there were no significant differences in D-banding length (P>0.05). The spread of D-banding value was however larger in the case of OIDI spanning from 50 to 85 nm, whereas this value was 52 to 75nm in the control samples. The average D-banding length found for both the native control and OIDI was also comparable to the commonly reported average of 67 nm. 29-31 This average however often refers to the D-banding periodicity length found in collagen in skin or tendon, and very little data have been reported on teeth until now.

Interestingly, the distribution of the fibril diameters in this study presented clear differences between the study groups. The native control sample fibrils' diameter was uniformly distributed over the range 50 to 75 nm where the OIDI fibrils diameter was bi-modally distributed. The large population of fibrils diameter were found to have a median of 62.1 nm, whereas the smaller population of fibrils had a median of 30.7 nm as presented in Figure 4C. This type of bimodal distribution is more reminiscent of tendons than bones or dentin. 31,32 The population of fibrils with the lower diameter is often associated with immature fibrils on tendons. Following this possible reasoning, our results may suggest that there is significant proportion of immature collagen fibrils present in OIDI dentin, which may result in weaker mechanical properties for the dentin structure and the tooth overall. Unlike tendons or bones, there is no direct regeneration through the combined osteoclasts/osteoblast activities in the dentin. Dentin is a calcified tissue formed through dentinogenesis, which results in the conversion of un-mineralized predentin into mineralized dentin.³³ As odontoblast cells move towards the pulp, they form an odontoblastic process, which penetrates the un-calcified predentin shaping the dentinal tubules.³⁴ Thus, once the odontoblast cells have remodelled the dentin, it is accepted that the dentin and its constituent remain unaltered throughout the lifetime of the

tooth, although these can be subjected to exogeneous factors (caries, for example). The presence of this bi-modal fibril diameter population is therefore not a result of fibril growth during the lifetime of the tooth but is already present at the dentinogenesis stage. It is plausible to consider that this variation in collagen fibrils diameter could be one of the key factors affecting the overall dentin mechanical stability and entire tooth prognosis in the case of OIDI.

Mechanical Properties of Dentin vs Dentinal Collagen Dentin Hardness

To evaluate the bulk mechanical properties of OIDI dentin versus control dentin, hardness measurements were performed directly on the sections prior to demineralisation. The hardness of a tissue determines its resistance to deformation. scratching, or abrasion. Additionally, hardness is described as the capability to resist permanent indentation.³⁵ In the case of teeth, it is a valid indicator to assess the tooth or dentin resistance to fracture. As presented in Figure 5A, the hardness of the control dentin was found to be significantly higher (p<0.05) than that of the OIDI. The mean hardness of the control dentin was found to be VHN_{control}=56.4±3.7 (N=80 indentations) while the mean hardness of the OIDI dentin was found to be VHN_{OIDI}=21.9.4±2.0 (N=80 indentations). The hardness of the OIDI is less than half of that of control dentin. This result confirms the expected result of this study that OIDI dentin is more fragile than control dentin as suggested by another study.³⁶ When considering the hardness of dentin, it is particularly important to evaluate its mineral contents. Kinney et al³⁷ measured the mineral content in DI type IIaffected and normal dentin using high-resolution synchrotron radiation computed tomography and found that the mineral concentration was 33% lower on average in the DI type II dentin with respect to normal dentin. Unlike the form of dentinogenesis (OIDI) that is linked with osteogenesis imperfecta, DI type II is not believed to be associated with any known defects in collagen structure. However, as OIDI is caused by abnormal type I collagen synthesis, it is highly likely that the interaction between collagen and minerals may also be affected and as a result, this would lead to poor mechanical properties.

Collagen Fibril Mechanical Properties

To evaluate if OIDI had any mechanical impact on dentinal collagen at the single fibril level, it was essential to ensure that only collagen fibrils with intact morphology would be characterized. As such, we omitted any area

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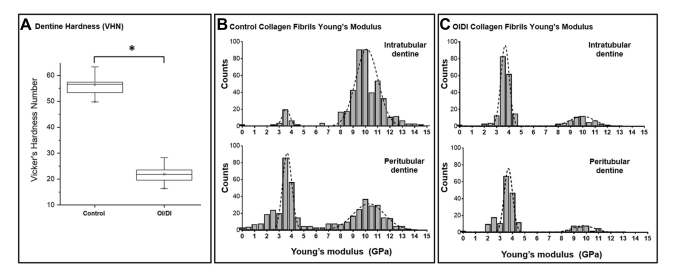


Figure 5 Graphical presentation of the nanoscale phenotypic properties (mechanical) for control and OIDI dentin (A) boxplots of the Vickers hardness value (*significance p<0.05). (**B** and **C**) Fitted histograms of the collagen fibril Young's modulus with bimodal distributions (Gaussian R²>0.9 for all distributions) as a function of measurement site: intratubular and peritubular, respectively.

previously described as weakened from this mechanical analysis as it was not possible to assert whether the topological changes where due to the demineralization protocol or if these are inherently associated with the disease. Thus, the AFM was used in imaging mode to identify collagen present with clear D-banding periodicity. By performing a systematic mechanical analysis, ³⁸ it was then possible to extract mechanical properties of individual collagen fibrils immediately surrounding the tubules (denoted intratubular) and equidistantly between tubules (denoted peritubular). Figure 5B and C shows the profile for the elastic (Young's) modulus of the control and OIDI collagen fibrils, respectively. In Figure 5B, the values of collagen elastic modulus for the control intratubular and peritubular regions followed a reverse mechanical profile. In the case of control intratubular dentin, the bimodal distribution with the greater population is centred at 10.4 GPa whereas a smaller population is centred at 3.6 GPa. In the case of control peritubular dentin, the bimodal distribution with the greater population is centred at 3.8GPa whereas a smaller population is centred at 10.3 GPa. From the process of dentinogenesis, it is known that odontoblasts lay down the primary dentin in the peritubular region first, before reinforcing and withdrawing through the tubules. The reinforcement method of collagen is by nanoscale chemical cross-linking, and previous research from our group has shown that these changes can be detected. 12 The data of the control supports this; the peritubular collagen has not been crosslinked to the extent that the intertubular collagen has, and so it is weaker.

In Figure 5C, the values of collagen elastic moduli for the OIDI intratubular and peritubular regions do not follow the same pattern as observed for the control sample. In fact, the collagen fibrils from both the intra and peritubular dentin present the same bimodal mechanical distribution. The largest population of collagen elastic moduli values for both the regions is centred around 3.8GPa whereas the smallest population for the collagen elastic moduli values is centred around 10.0 GPa and 9.8 GPa for the intratubular and the peritubular regions, respectively. This lack of reverse pattern in the collagen elastic moduli in the case of OIDI with respect to the distance from the tubule is indicative that the collagen is mechanically more uniform throughout the dentin thickness, albeit being also mechanically weak.

The formation of dentinal tubules is the direct outcome of the migration of differentiated odontoblasts towards the dental papilla, leaving behind processes around which the dentine matrix mineralizes. Yet, histologically, and microscopically, these tubules appear as tubular voids within the dense mineral matrix that is dentine. For a structural and mechanical point of view, the presence of these tubular voids ought to weaken the composite material properties of dentine. Yet, collagen fibrils immediately surrounding the tubules were found to be mechanically stiffer in the control dentin when compared to the fibrils present equidistantly between dentine tubules. This ought to suggest that the collagen fibrils immediately surrounding the tubules are mechanically reinforced to support the interruption dentine structure created by the presence of tubular

voids (dentine tubules). However, the OIDI samples did not show this region-dependent change in stiffness. This would suggest that the dentine structure is not reinforced around the dentinal tubules, which would lead to overall weaker dentine mechanical and structural properties. A plausible reason for this to occur may be linked to the inability of the collagen present in these intra-tubular regions, the case of OIDI to respond to extra-cellular posttranslational processing such as crosslinking for example.

Although more research would be required to evaluate the crosslinking profiles of regions specific collagen, this finding suggests that the quality of the collagen fibrils is mechanically poorer in the case of dentinogenesis imperfecta associated with osteogenesis imperfecta, which in turn reduce the dentine structural and mechanical properties.

Clinical Impact

Dentinal collagen plays an important role in the stability of dentin but unlike in bones, it is not renewed during the lifetime of the tooth. Thus, any endogenous (genetic, disease) or exogenous alteration²¹ to the collagen network will impact both its structure and function as part of a functional mineralised tissue. In the case of OIDIaffected teeth, we have demonstrated in an exploratory approach how dentinal collagen fibrils exhibit a bimodal distribution in terms of diameter which is not found in control samples. Correlating this finding with the observed lack of increased elastic moduli of the collagen fibrils in the intratubular OIDI dentin may be a first explanation towards the definition of structural changes in the dentin responsible for the failure of some adhesive restorative materials as observed in patients affected by OIDI.

Mild types of OI can be difficult to diagnosis clinically, therefore dental assessment, including clinical and radiographic examination, can be important to aid diagnosis.³⁹ Children with suspected OI should always be screened by a specialist paediatric dentist, as dental management of OIDI children can be complex. Whilst preformed metal crowns are suitable for restoration of primary molars, due to the increased tendency of composite restorations to fail in DI patients; glass ionomer materials are commonly used as means of restoring anterior teeth as they do not depend on mechanical interlocking with collagen. 40 It is unclear whether the mode of restoration and improper etching of teeth prior to applying dental composites, or possible dentinal matrix or collagen alterations, that might be ultra-structurally defective and mechanically weaker thus forming an ineffective hybrid layer during bonding, which can be a causative factor in failure. 40

Conclusion

This study presents a systematic approach to the characterization of collagen structure and properties in OIDI as diagnosed in dentin. A modified demineralization protocol comprising acid sonication resulted in much clearer topographic images of exposed dentinal collagen. In our approach, we propose that demineralization protocols should ideally be sample-specific and tailored to each sample examined. After the analysis of data obtained from AFM and SEM imaging, it was found that to study the collagen ultrastructure within a mineralized matrix, it was essential to reach a balance between mineral removal and collagen exposure, whilst avoiding the denaturation of the collagen matrix itself. Structural markers for OIDI at the mesoscale and nanoscale were found and supported by a mechanical analysis. D-banding periodicity average values were consistent with the literature, but were statistically insignificant when compared with OIDI samples. However, dentinal tubule number and collagen fibril diameter, and the overall fibril meshwork arrangement, were seen to vary considerably. One of the main outcomes of this study was to explore the topography of the collagen network and its mechanical properties in the case of OIDI. New questions and challenges have been raised regarding the quality of the collagen network in the case of DI associated with OI, its impact upon treatment strategies in dentistry, and ultimately the correct treatment protocol to most benefit patients.

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Author Contributions

All authors contributed to data analysis, drafting or revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

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Disclosure

The authors report no conflicts of interest in this work.

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