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## Biomimetic remineralization of dentin



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### ABSTRACT

**Objectives.** Remineralization of demineralized dentin is important for improving dentin bonding stability and controlling primary and secondary caries. Nevertheless, conventional dentin remineralization strategy is not suitable for remineralizing completely demineralized dentin within hybrid layers created by etch-and-rinse and moderately aggressive self-etch adhesive systems, or the superficial part of a caries-affected dentin lesion left behind after minimally invasive caries removal. Biomimetic remineralization represents a different approach to this problem by attempting to backfill the demineralized dentin collagen with liquid-like amorphous calcium phosphate nanoprecursor particles that are stabilized by biomimetic analogs of noncollagenous proteins.

**Methods.** This paper reviewed the changing concepts in calcium phosphate mineralization of fibrillar collagen, including the recently discovered, non-classical particle-based crystallization concept, formation of polymer-induced liquid-precursors (PILP), experimental collagen models for mineralization, and the need for using phosphate-containing biomimetic analogs for biomimetic mineralization of collagen. Published work on the remineralization of resin-dentin bonds and artificial caries-like lesions by various research groups was then reviewed. Finally, the problems and progress associated with the translation of a scientifically sound concept into a clinically applicable approach are discussed.

**Results and significance.** The particle-based biomimetic remineralization strategy based on the PILP process demonstrates great potential in remineralizing faulty hybrid layers or caries-like dentin. Based on this concept, research in the development of more clinically feasible dentin remineralization strategy, such as incorporating poly(anionic) acid-stabilized amorphous calcium phosphate nanoprecursor-containing mesoporous silica nanofillers in dentin adhesives, may provide a promising strategy for increasing of the durability of resin-dentin bonding and remineralizing caries-affected dentin.

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## 1. Introduction

Teeth are the most heavily mineralized tissues in the human body. Demineralization and remineralization processes coexist in teeth during the entire life of an individual. In pathological conditions, demineralization outweighs remineralization [1]. Fermentation of dietary carbohydrates by acidogenic bacteria results in the production of acids such as lactic acid, acetic acid and propionic acid that demineralize enamel and dentin. As the carious lesion progresses into dentin, activation of endogenous, bound matrix metalloproteinases and cysteine cathepsins will lead to the degradation of collagen fibrils and decrease in the mechanical properties of dentin [2,3]. Prevention and treatment of dental caries is a major challenge because as many as nine out of ten adults in Western countries suffer from dental caries [4]. In the United States alone, more than 100 million dollars is spent annually on dental service. Despite significant advances in preventive and restorative dentistry, replacement of tooth fillings constitutes a substantial share of this annual dental expenditure, due to limited durability of contemporary resin-based restorative materials, particularly when these materials are applied to damaged dentin in the absence of a superficial enamel layer [5–7].

Apart from caries, resin–dentin bonding is another major reason for dentin demineralization [8]. The formation of resin–dentin bonds is accomplished predominantly by micromechanical retention via resin penetration and entanglement of exposed collagen fibrils in the partially or completely demineralized dentin. This is achieved by etching dentin with acids or acidic resin monomers derived from self-etching primers/adhesives to expose the collagen fibrils [7]. To date, it is impossible for resin monomers to completely displace water within the extrafibrillar and particularly the intrafibrillar compartments of a demineralized collagen matrix, and infiltrate the collagen network completely [8–10]. Even if this may be achieved, the limited intermolecular space (1.26–1.33 nm) between collagen molecules renders it challenging to accommodate even small, extended resin monomer molecules such as triethyleneglycol dimethacrylate ( $\approx 2$  nm long) [11]. This invariably results in the presence of mineral-depleted, resin-sparse, water-rich collagen fibrils along the bonded interface [7,9]. Under the combined challenges of enzymes, temperature and functional stresses, regions of incomplete resin infiltration within the dentin hybrid layer is susceptible to degradation, resulting in damage of interfacial integrity, reduction in bond strength and ultimately, the failure of resin–dentin bonds. Thus, remineralization of demineralized dentin has important consequences for control of dentinal caries as well as improvement of dentin bonding stability [8,10].

Different strategies have been employed for remineralizing demineralized dentin. For instance, fluoride, amorphous calcium phosphate (ACP)-releasing resins or resin-based adhesives containing bioactive glass have been used to improve the resistance of bonded restorations to secondary caries [12–15]. However, most of these studies focused on remineralizing partially demineralized carious dentin, which was based on the epitaxial deposition of calcium and

phosphate ions over existing apatite seed crystallites [16]. With these traditional ion-based strategies, remineralization does not occur in locations where seed crystallites are absent [17]. Thus, the classical ion-based crystallization concept may not be applicable for remineralizing completely demineralized dentin within hybrid layers created by etch-and-rinse adhesive systems or the superficial part of a caries-affected dentin lesion left behind after minimally invasive caries removal, due to the unavailability of seed crystallites in those regions for accomplishing homogeneous nucleation of apatite crystallites [18,19].

Biomimetic remineralization represents a different approach to this problem by attempting to backfill the demineralized dentin collagen with liquid-like ACP nanoprecursor particles that are stabilized by biomimetic analogs of noncollagenous proteins [20–22]. This is achieved by adopting the recently discovered, non-classical particle-based crystallization concept utilized by Nature in various biominerization schemes, ranging from the mineralization of sea-shells (calcium carbonate), siliceous shells of diatoms and sponges (amorphous silica) to the deposition of calcium phosphate salts in fish scales and bone [23,24]. Intrafibrillar mineralization of fibrillar collagen not only significantly increases its mechanical properties [25–28], but also protects the collagen molecules from external challenges, such as temperature, endogenous enzymes, bacterial acids and other chemical factors. Using this biomimetic remineralization strategy, both hybrid layers created by etch-and-rinse adhesives [21,29,30] and moderately aggressive self-etch adhesives [18,31,32], as well as 250–300  $\mu\text{m}$  thick completely demineralized dentin lesions can be remineralized [33,34]. This bottom-up remineralization strategy does not rely on seed crystallites, and may be considered as a potentially useful mechanism in extending the longevity of resin–dentin bonds [35] via restoring the dynamic mechanical properties of the denuded collagen within the hybrid layer to approximate those of mineralized dentin [36]. This paper reviews the changing concepts in calcium phosphate remineralization and the progress in clinical translation of the biomimetic dentin remineralization strategy.

## 2. Changing concepts of calcium phosphate biominerization

Biominerization is the process by which living organisms secrete inorganic minerals in the form of biominerals (e.g. magnetite, silica, oxalates, various crystalline forms of calcium carbonate and carbonated apatite) within cell cytoplasm, shells, teeth and bony skeletons [37,38]. This process exhibits a high level of spatial and hierarchical control as mineralization usually takes place in a confined reaction environment under ambient temperature and pressure conditions. Calcified human tissues consist of the collagen matrix and the hierarchically arranged carbonated apatite inorganic phase; deposition of the latter is regulated by non-collagenous proteins [39,40]. It is generally believed that non-collagenous proteins, along with specific MMPs and other important enzymes secreted by odontoblasts, play critical roles to orchestrate dentin mineralization. They possess carboxylic

acid and phosphate functional groups that act as preferential sites for Ca/P nucleation and subsequent apatite crystallization [41–43]. As the therapeutic use of native or recombinant non-collagenous proteins for *in situ* biominerization is not yet economically viable, research scientists have resorted to the use of polyelectrolyte and poly(acid) macromolecules to mimic the functional domains of these naturally occurring proteins, in biomimetic mineralization [44–47]. In the past few years, this field of research has attracted a lot of attention, resulting in changing concepts of calcium phosphate biominerization.

### 2.1. Particle-based vs ion-based crystallization

Traditional collagen mineralization studies were based on the classical pathway of ion-mediated crystallization, or classical nucleation theory [37,48]. The classical model of crystal formation begins with crystal nucleation, followed by crystal growth. This process starts from primary building blocks like atoms, ions or molecules, forming clusters, which may grow or disintegrate again, depending on the counter-play of surface and crystal lattice energies. Eventually, some clusters reach the size of a so-called critical crystal nucleus. These primary nuclei grow further via ion-by-ion attachment and unit cell replication. While the classical crystallization model was relatively successful in controlling the dimensions of calcium carbonate or calcium phosphate, they achieved limited success in reproducing the structural hierarchy of intrafibrillar apatite deposition [49] within the collagen matrix.

In contrast to the ion-mediated classical crystallization pathway, the non-classical crystallization pathway is particle-mediated and involves a mesoscopic transformation process [22,50]. The term mesoscopic refers to materials of an intermediate length scale, the lower limit may be the size of individual atoms. Whereas macroscopic objects usually obey the laws of classical mechanics, a large number of particles can interact in a quantum-mechanically correlated fashion for mesoscopic objects. Evidence for these pathways is rapidly increasing in the literature. To date, mesocrystals of a wide range of materials including  $\text{CaCO}_3$ ,  $\text{BaSO}_4$ , metal oxides, metal tungstates and chromates,  $\text{NH}_4\text{TiOF}_3$ ,  $(\text{NH}_4)_3\text{PW}_{12}\text{O}_{40}$ ,  $\text{LiFePO}_4$ , metal chalcogenides (e.g. semiconductor cadmium sulphide), noble metals, and organic materials have been successfully synthesized, as described in several excellent reviews [23,51]. The contemporary concept of calcium phosphate biominerization has been advanced by parallel studies on the biominerization of calcium carbonate (calcite and aragonite) from amorphous calcium carbonate [52–54]. In the context of calcium phosphate biominerization, calcium and phosphate ions are sequestered by biomimetic analogs of non-collagenous proteins involved in hard tissue mineralization into nanoparticles that exist in nanoscopic units known as prenucleation clusters [55–57]. These prenucleation clusters ( $\approx 1\text{ nm}$  in diameter) eventually aggregate into larger ( $10\text{--}50\text{ nm}$  in diameter) liquid-like ACP nanoparticles. The latter, on penetration into the intrafibrillar water compartments of a collagen fibril, utilize the latter as a mineralization template, and undergo self-assembly and crystallographic alignment to form a metastable crystalline phase via mesoscale assembly. These mesocrystals probably fused to form iso-oriented

crystal intermediates, and finally to single apatite crystallites within the  $40\text{ nm}$  wide gap zone between the collagen molecules [23,58,59]. This ordered arrangement of apatite crystallites results in the manifestation of a banded appearance in unstained, mineralized fibrillar collagen [60].

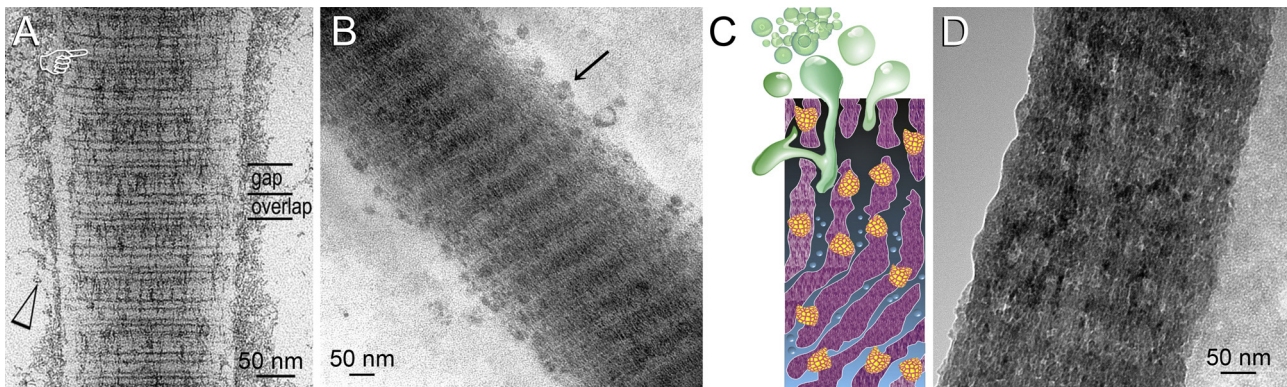
### 2.2. Polymer-induced liquid precursor

Formation of amorphous nanoprecursors is the fundamental step in many forms of biominerization. Existence of transient ACP nanoprecursors has been identified from enamel and bone [61–63]. In biomimetic mineralization of type I collagen, Dr. Gower and colleagues pioneered a process based on formation of a polymer-induced liquid-precursor (PILP) system [54,64]. The plasticity of liquid amorphous mineral precursors allows them to take the shape of their containers, resulting in a variety of biominerals with different hierarchical structures [65,66]. Using the PILP concept, Gower and colleagues were successful in mineralizing a variety of organic matrices with both calcium carbonate and calcium phosphate, including intrafibrillar mineralization of collagen matrices by carbonated apatite [47,67,68].

Based on the aforementioned new concepts of biominerization introduced over the past decade, biomimetic mineralization of type I collagen may be envisaged to proceed in several stages [59]: (1) the collagen fibril serving as an active template for hierarchical intrafibrillar mineralization; (2) calcium and phosphate ions in the calcifying medium self-assemble into stable particulate units known as pre-nucleation clusters. In the presence of a polyanionic analog of matrix proteins such as poly(aspartic acid), these pre-nucleation clusters further condense into larger particles of fluidic amorphous ACP precursors (in the range of  $10\text{--}30\text{ nm}$ ) that are capable of diffusing into the intrafibrillar compartments of type I collagen; (3) the negatively charged polyanion-stabilized ACP precursors interact with positively charged sites along the collagen molecules, inducing solidification and nucleation of the ACP inside collagen; (4) these nucleated amorphous mineral precursor phases further growth and maturation into apatite nanocrystals along the intrafibrillar space of collagen via non-classical crystallization pathways; (5) Growth of the intrafibrillar apatite results in heavy intrafibrillar mineralization by apatite crystallites, with concomitant extrafibrillar mineralization between adjacent collagen fibrils.

### 2.3. Models of collagen mineralization

In our early studies, totally or partially demineralized dentin collagen matrices were employed for remineralization of hybrid layers and caries-like dentin lesions [17,20,28–36]. However, it is not known if the remineralization demonstrated in those studies was caused by the presence of remnant phosphoproteins that remain bound to collagen matrices after demineralization [69]. As self-assembled purified collagen fibrils do not contain bound matrix proteins, the authors employed a two-dimensional model of biomimetic mineralization of collagen based on reconstituted collagen fibrils deposited on grids used for transmission electron microscopy [70–72]. Briefly, bovine skin-derived type I collagen dissolved



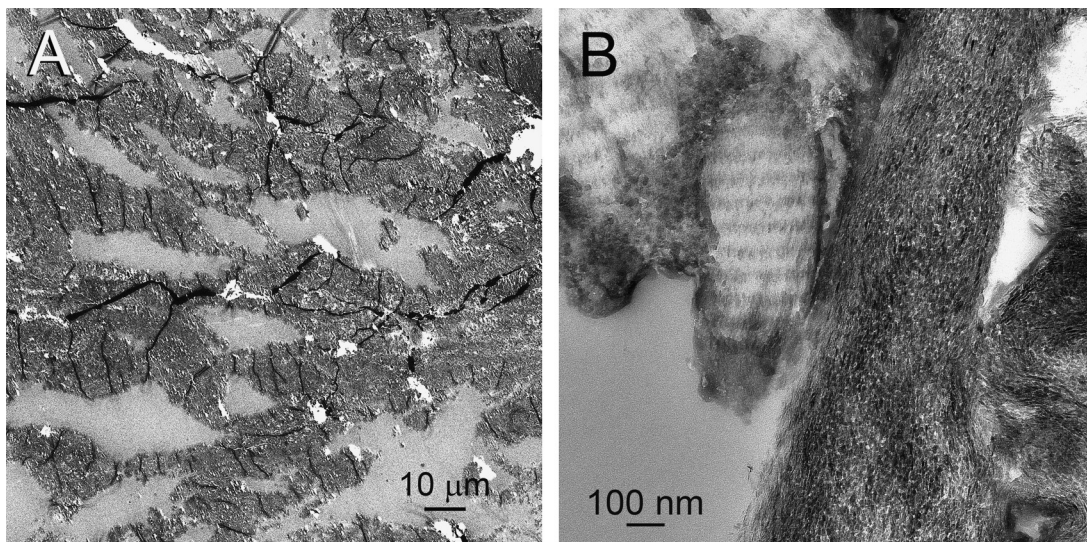
**Fig. 1 – A two-dimensional model for examining mineralization of reconstituted collagen fibrils, using a dual biomimetic analog mineralization protocol. Transmission electron microscopy (TEM) images of unsectioned reconstituted collagen fibrils. (A)** After 24 h of mineralization. The fibril was briefly stained with uranyl acetate to highlight the gap and overlap zones. Amorphous calcium phosphate (ACP) prenucleation clusters (open arrowhead) could be seen along the periphery of the fibril. Pointer: initial needle-shaped intrafibrillar apatite. **(B)** Unstained collagen fibril after 48 h of mineralization. Prenucleation clusters have coalesced into ACP droplets (arrow) that continued to infiltrate the fibril. Denser intrafibrillar apatite deposition could be seen but the banding periodicity caused by hierarchical arrangement of the apatite crystallites was still present. **(C)** Schematic depicting replacement of intrafibrillar water with fluidic polyacrylic acid-stabilized ACP nanoprecursors. **(D)** Unstained fibril showing heavy intrafibrillar mineralization with apatite platelets after 72 h. Banding characteristics was obscured by the heavy mineralization.

in acetic acid was reconstituted on formvar-coated grids by ammonia diffusion and cross-linked with 0.3 M carbodiimide. Then, the grid with a single layer of collagen was floated over a drop of a prospective biomimetic analog-containing biominerization medium. The grid was retrieved at different time points (usually within 72 h) and rinsed with water. They were examined unstained for mineral deposition or stained for correlation of mineral deposition with collagen cross-striation patterns. With this model, we demonstrated a highly hierarchical assembly of intrafibrillar apatite crystallites in reconstituted collagen mineralized ex situ with biomimetic analogs of extracellular matrix proteins [73] (Fig. 1). This 2-D single-layer collagen mineralization model is useful as a rapid screening tool for potential biomimetic analogs for collagen mineralization [71]. Moreover, the technique may be modified for creating intrafibrillarily mineralized collagen coatings on the surfaces of orthopedic and dental implants as a more biocompatible alternative to the use of hydroxyapatite coatings [74].

Although the single-layer collagen model is a convenient and rapid method for studying biomimetic mineralization of individual collagen fibrils, the simple 2-D structure of the collagen model limits its further application. To further mimic natural collagen matrices, 3-D models of reconstituted collagen (Fig. 2) were used by different research groups to evaluate the ability of the biomimetic mineralization scheme to mineralize collagen bundle assemblies within a three-dimensional scaffold [68,75]. Briefly, commercially available type I collagen sponges derived from bovine tendon were cultured in analog-containing biominerization medium for 3–14 days. The mineralization assembly was kept in a 37 °C oven to emulate physiological conditions. At predetermined reaction times, mineralized samples were removed from the solution, copiously washed with deionized water and further

prepared for different analysis, including transmission electron microscopy, thermogravimetric analysis, attenuated total reflection-Fourier transform infrared spectroscopy, nuclear magnetic resonance spectroscopy and X-ray diffraction, which cannot be easily achieved with the 2-D collagen model [68,75,76]. The formed collagen-hydroxyapatite composites indicated that polyaspartic acid-stabilized ACP fluid droplets can diffuse into collagen bundles and mineralize a relatively thick collagen scaffold [76].

Although the mechanical properties of 3-D collagen sponges increase significantly after functional mineralization, they are still far less than those of the natural mineralized tissues because of their highly porous nature. Thus, attempts have been made by various groups to produce biomaterials that structurally mimic bone and dentin [68]. However, the hierarchical complexity of natural composites is usually far beyond contemporary bioengineering processing capability. One approach is to utilize the hierarchical structure of natural materials as a template for the development of high-performance engineering materials using in vitro biomimetic methods [77]. Thus, researchers revisited the use of natural collagen model, including rat tail tendon (a soft tissue collagen 3D model; Fig. 3), demineralized manatee bone and demineralized dentin collagen [68,78,79]. It was found that the amount of mineral present in the rat tail tendon samples represented the maximum amount of mineral available in the PILP solution [68]. Conversely, for demineralized manatee bone specimens, mineral distribution and penetration across the bulk of the bone matrix appears to be a challenge for the PILP process. The mineral penetration depth was limited to 100 μm [78]. The mineral content for the remineralized manatee bone specimens (about 45 wt%) was below the theoretical value for cortical bone, even though enough calcium and phosphate ions were available in the PILP solution to restore the



**Fig. 2 – A three-dimensional reconstituted collagen model of biomimetic mineralization.** TEM images of an unstained section taken from a highly porous collagen scaffold that had been mineralized with a calcium and phosphate-containing medium containing dual biomimetic analogs for 14 days. (A) Low magnification of the mineralized collagen scaffold, showing interconnecting mineralized collagen bundles. (B) High magnification taken from a part of a mineralized collagen bundle. Collagen fibrils on the left are less heavily mineralized, with hierarchical deposition of apatite platelets in the gap zones, producing a periodic banding pattern. Collagen fibrils on the right are heavily mineralized with apatite platelets. Banding characteristic can no longer be identified.

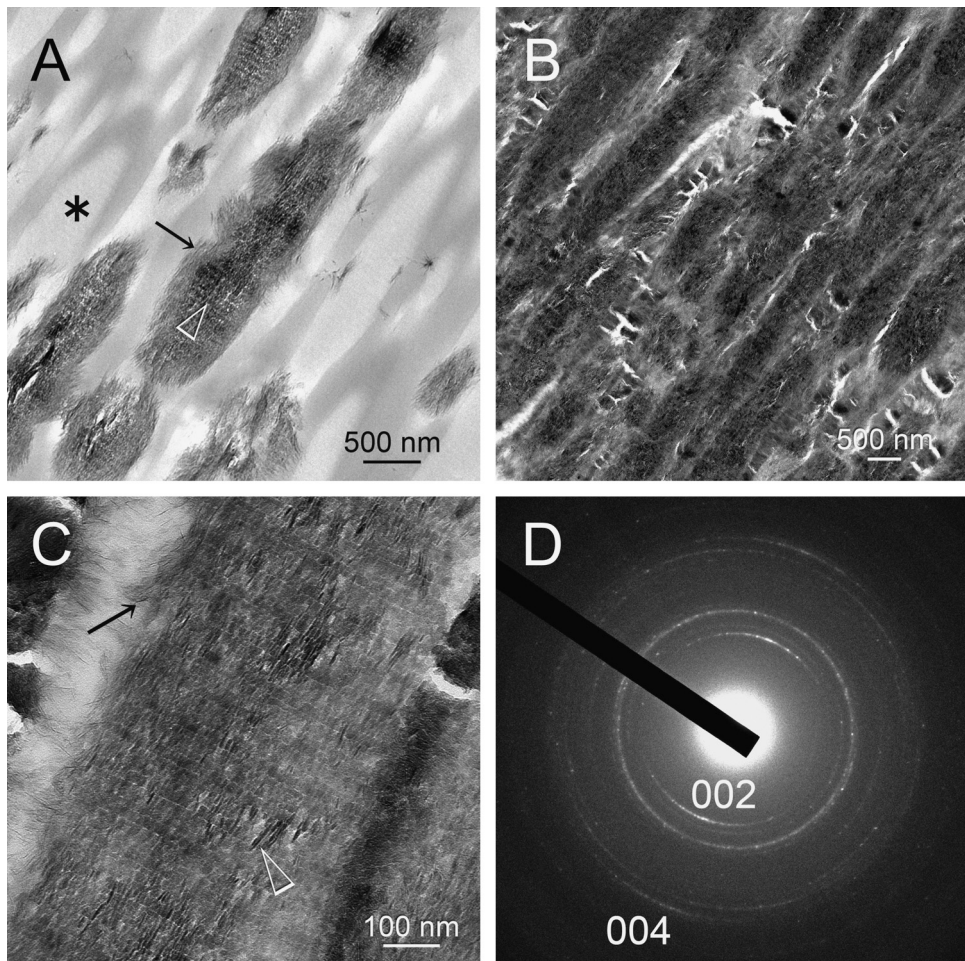
mineral value of bone. It was proposed that auto-transformation and solidification of the surface-infiltrated ACP nanoprecursors into crystalline apatite with time could potentially block further infiltration of the additional nanoprecursors into the bulk of the demineralized bone matrix, thereby limiting their ultimate depth of penetration.

#### 2.4. The need for using phosphate-containing biomimetic analogs in biomimetic mineralization of collagen

The functional role played by collagen in calcium phosphate biominerization has been conjectural based on the results of the past several decades of research. Earlier studies indicated that fibrillar collagen does not initiate biominerization on its own, but serves as a passive depot for the housing of apatite crystallites [69,80,81]. This led to a plethora of studies that examined the biological control of intrafibrillar collagen mineralization by noncollagenous proteins [82–87]. Noncollagenous proteins are believed to play a crucial role in the mineralization of bone and dentin. This assertion is supported by studies demonstrating that mutations in genes that code for these proteins result in abnormal bone and dentin mineralization [88–91]. The extracellular matrix of bone and dentin contains small quantities of noncollagenous proteins such as osteopontin, bone sialoprotein, dentin matrix protein 1 (DMP1), and dentin sialophosphoprotein (DSPP) [39,40]. The latter is further cleaved into dentin sialoprotein and dentin phosphoprotein (DPP; aka phosphophoryn) [92]. These proteins are highly anionic due to the prevalence of carboxylate groups on the polyaspartic acid residues that comprise the protein backbone. Post-translation phosphorylation of the

serine residues produces phosphoserines, which further augments their anionic character [93,94]. For example, dentin phosphoprotein, being the most abundant NCP in dentin, contains a large number of aspartic acid (Asp) and phosphoserines (Pse) in the repeating sequences of  $(\text{Asp-Pse})_n$  and  $(\text{Asp-Pse-Pse})_n$  [95]. Dentin matrix protein 1, another member of the Small Integrin-Binding Ligand Interacting Glycoproteins (SIBLING) family, also contains high levels of Asp and Pse [96,97]. A unifying feature of the SIBLING proteins is that they all contain an acidic serine aspartate-rich MEPE (ASARM)-associated motif. The ASARM motif in SIBLING genes and the released ASARM peptide play roles in mineralization of bone and teeth [98,99]. The highly anionic nature of noncollagenous proteins involved in biominerization enables them to sequester and bind calcium ions and presenting them to collagen fibrils at the mineralization front during the formation of bone and dentin.

In the authors' laboratory, biomimetic mineralization of collagen was performed using a dual biomimetic analog strategy [70–72,100–102]. Polyacrylic acid or polyaspartic acid was used as an analog for sequestering calcium ions released by set calcium silicate cements or a supersaturated calcium phosphate mineralization solution. These analogs function by acting as surfactants to prevent fluidic ACP nanoparticles from aggregating into larger particles, and to inhibit auto-transformation of the ACP nanoparticle into apatite (i.e. apatite nucleation inhibitor) prior to their entry into the intrafibrillar water compartments of the collagen fibril [70,71]. In addition, a polyphosphate-containing biomimetic analog such as polyvinylphosphonic acid, sodium trimetaphosphate or sodium ascorbyl phosphate was employed as a templating biomimetic analog of matrix phosphoproteins. These



**Fig. 3 – A three-dimensional natural soft tissue collagen model of biomimetic mineralization.** TEM images of unstained sections taken from mineralized rat tail tendon with parallel collagen fibrils at different time periods. (A) Incomplete mineralization at 7 days. Asterisk: Unmineralized collagen fibrils; Arrow: extrafibrillar mineralization; Open arrowhead: Intrafibrillar mineralization recapitulating the D-spacings of fibrillar collagen. (B) Heavy mineralization of the parallel collagen fibrils after 14 days. (C) High magnification of B, showing a collagen fibril with extrafibrillar (arrow) and intrafibrillar mineralization by discrete apatite platelets (open arrowhead). (D) Selected area electron diffraction of the mineralized fibril in (C) produced arc-shaped diffraction patterns in the 0 0 2 and 0 0 4 plane of apatite. The C-axis of the apatite platelets is arranged almost parallel to the longitudinal axis of the collagen fibril.

phosphoprotein analogs were allowed to bind to the collagen fibrils prior to immersion of the fibrillar matrices in the poly(anionic) acid-containing mineralization medium [100–102]. Using the aforementioned collagen mineralization models, attachment of ACP nanoprecursors to the D-spacings of non-mineralized collagen fibrils was observed within 24 h. After 48 h, hierarchical apatite deposition was observed from unstained collagen fibrils that resulted in banded mineral arrangement. After 72 h, highly mineralized collagen fibrils could be seen in which the periodicity of the mineral arrangement was obscured as the fibrils were completely mineralized [70–73].

In contrast to the earlier work, more recent studies indicate that type I collagen plays an active role in biominerization by acting as templates for physicochemical (electrostatic) attraction of ACP nanoparticles and direct formation of intrafibrillar apatite within the gap zones, without the intervention or

mediation of other noncollagenous proteins present in vertebrate calcifying tissues [103,104]. Silver and Landis reported specific sites in the e2 band of the gap zones (corresponding to the a, e and d bands) of type I collagen that have the potential to sequester and bind calcium ions [105]. Nudelman et al. [58] calculated that there are net positive charges close to the C-terminal end of the collagen molecules that favor infiltration of the collagen fibrils with negatively charged ACP nanoprecursors. The arrangement of charged amino acids within the gap and overlap zones produce nucleation sites that control the conversion of the ACP nanoprecursors into oriented apatite crystals [58]. Indeed, several research groups have demonstrated that it is possible to produce intrafibrillar mineralization of type I collagen using poly(aspartic acid)-stabilized ACPs alone as an apatite nucleation inhibitor, without the adjunctive use of phosphorylated DDP, phosphorylated DMP1 or their polyphosphate analogs [58,78,106,107].

Although this simplified biomimetic collagen mineralization strategy is more economical and reduces time in preparing mineralized collagen scaffolds from a tissue engineering perspective, it begs the provision of a rationale for the existence of highly phosphorylated noncollagenous proteins in bone and dentin. The use of the simplified biomimetic mineralization strategy also challenges earlier biological studies that reported critical roles played by the phosphorylated forms of these noncollagenous proteins in the formation of mineralized collagenous tissues [93,96,108,109].

A recent study compared the effects of phosphorylated vs non-phosphorylated forms of DPP and DMP1 on collagen mineralization using a 2D model [110]. Although differences existed between DPP and DMP1 in the locations in which collagen fibrils were mineralized (intrafibrillar vs extrafibrillar), both phosphorylated proteins facilitated highly organized intrafibrillar mineralization of collagen fibrils. Conversely, the use of non-phosphorylated forms of these proteins resulted in randomly oriented intrafibrillar crystallites, with no particular organization of their crystallographic axes to the longitudinal axis of the collagen fibrils. Nevertheless, this study failed to explain why non-phosphorylated biomimetic apatite nucleation inhibitors such as polyaspartic acid or fetuin-A [111,112] are capable of producing highly organized intrafibrillarily mineralized collagen. It must be emphasized that while the concepts of particle-based ACP prenucleation clusters and the PILP phenomenon of fluidic ACP infiltration are new in collagen biomineralization research, the theoretical basis for interpretation of ultrastructural results of intrafibrillar apatite deposition is not [58,106], and is based upon the classic model of collagen molecular packing proposed by Petruska and Hodge [113]. This straight and rod-like model of steric arrangement of collagen molecules has since been replaced by a synchrotron X-ray diffraction-derived model in which collagen molecules are arranged in a right-handed helically twisted, discontinuous manner along the length of the microfibrils [114]. In the latter model, interdigitation of adjacent microfibrils placed geometric constraints on the availability of lateral intermolecular spacings between collagen molecules, with no room to accommodate apatite platelets outside the gap zones.

A recent study examined the effect of using single (polyacrylic acid) vs dual biomimetic analogs for mineralization of collagen fibrils (polyacrylic acid and sodium trimetaphosphate) [73]. In that study, polyacrylic acid was employed as an analog to inhibit apatite nucleation, and sodium trimetaphosphate was used as a templating analog for guiding intrafibrillar apatite deposition. The use of polyacrylic acid without a templating analog resulted only in intrafibrillar mineralization with continuous apatite strands. Conversely, the use of both analogs resulted in intrafibrillar mineralization with discrete apatite crystallites. While both methods resulted in intrafibrillar mineralization of collagen fibrils, the authors opined that in the absence of polyphosphate as a templating analog, infiltration of poly(anionic) acid-stabilized ACP nanoprecursors via a PILP process into the interconnecting water-filled volume within a collagen fibril appeared to have resulted in molding of ACP nanoprecursors into a continuum. This, in turn, resulted in crystallization of the carbonated apatite into a monolithic crystalline structure. Such a

crystallization mechanism produces mineralized collagen entities that resemble the monolithic single-crystal structure in sea urchin spines or siliceous bio-skeletons. It is possible that release of poly(anionic) acid into the intrafibrillar milieu results in osmotic swelling and relaxation of the helical collagen microfibrillar arrangement, that facilitates continuous apatite deposition or growth from the gap zones into overlap zones into continuous strands. Conversely, electrostatic binding of polyphosphate analogs to discrete sites along the collagen molecules may have caused the bound analogs to act as inhibitor to discourage continuous growth of the apatite crystallites along the overlap zones, thereby resulting in constraining apatite platelets to the gap zones of the collagen fibril. This hypothesis is speculative and awaits further validation.

Nevertheless, the aforementioned hypothesis is consistent with the timely report comparing the ultrastructure of mineralized collagen in bone with the use of ion-milled sections vs sections prepared by conventional ultramicrotomy [115]. In ion-milled sections, the authors reported that approximately 70% of the minerals in bone are extrafibrillar, confirming previous models proposed by Lees et al. [116] and Hellmich and Ulm [117]. With respect to intrafibrillar apatite, crystallite platelets were only identified in the gap zones, without extension into the overlap zones, in contrast to what was previously proposed by Landis et al. [118]. Similar results were reported in another study using steric modeling to estimate the packing density of apatite within the gap zones [119]. Those modeling results were further confirmed using electron energy loss spectroscopy associated with scanning transmission electron microscopy. Taken together, these novel findings highlight that mechanisms in the control of discrete intrafibrillar apatite platelet deposition in the gap zones by phosphorylated NCPs are not completely understood. It is possible that the current success of intrafibrillar collagen mineralization with a single poly(anionic) acid analog may not result in truly biomimetically mineralized collagen that resembles those present in bone and dentin. These challenging issues require further in-depth investigations to decipher the riddles.

### **3. Adaptation of collagen biomineralization strategy for remineralization of resin–dentin bonds**

Apart from increases in mechanical properties [120,26], a major role played by the presence of intrafibrillar apatite in mineralized collagen is the exclusion of molecules larger than water (ca. 18 Da) from the mineral–protein biocomposite [24]. Whereas even ethanol (ca. 46 Da) cannot solvate air-dried mineralized dentin, demineralized, water-filled collagen is susceptible to penetration by larger molecules. Molecules larger than 40 kDa are completely excluded from the multiple internal water compartments [121] of type I collagen, while molecules smaller than 6 kDa can diffuse into all of the water compartments within the collagen fibril [122]. The physical exclusion of exogenous collagenolytic enzymes (bacterial collagenase – 68–130 kDa; activated MMP-2 – 67 kDa; activated MMP-9 – 85 kDa; activated cathepsin K – 27 kDa) by apatite forms the tenet of the “enzyme exclusion” mechanism that

protects archeological collagen from degradation [123,124]. Experiments on collagenase hydrolysis of dentin have also confirmed the protective role played by the mineral phase on collagen degradation [125]. Endogenous MMPs and cysteine cathepsins, as well as other growth factors become “fossilized” by the intrafibrillar apatite minerals within dentin [126], but retain their biologic activities after they are incorporated into the mineralized dentin matrix. As collagen mineralizes, free and loosely bound water is progressively replaced by apatite. This physiologic dehydration mechanism [127] ensures that the internal environment of the mineralized fibril remains relatively dry to preserve the integrity of the entrapped bioactive molecules [128].

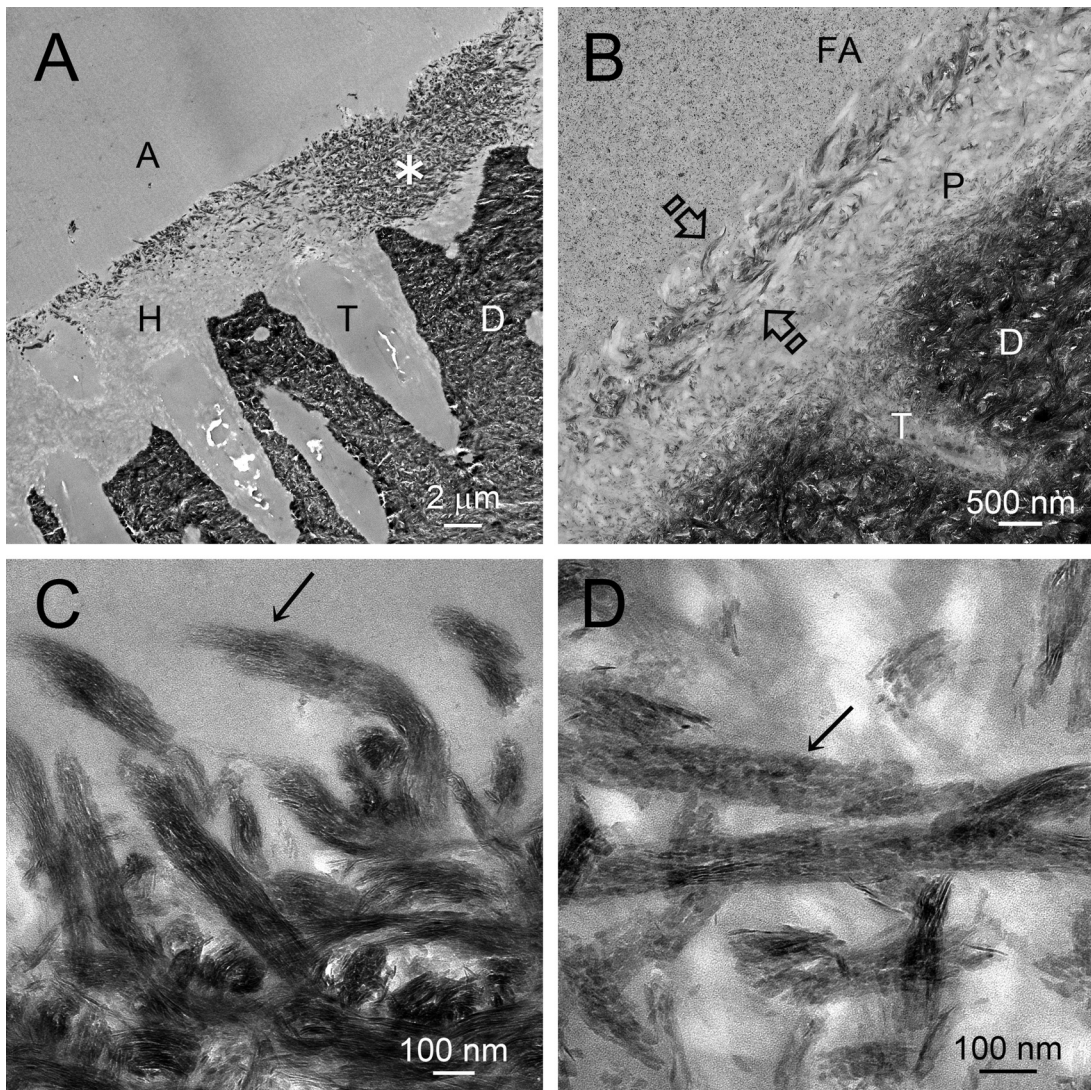
In dentin bonding, the mineral phase of dentin is intentionally removed by acids, chelating agents or acidic resin monomers to expose the collagen for micromechanical retention of resins. As dentin demineralizes, bioactive molecules such as growth factors, matrix metalloproteinases and cysteine cathepsins are activated, provided that the demineralization agent is not strong enough to denature these molecules [3]. Contemporary etch-and-rinse and self-etch adhesives are incapable of completely replacing water from the extrafibrillar and intrafibrillar collagen compartments with resin monomers [21,129,130]. This can be seen by the number of publications on nanoleakage and micropermeability within hybrid layers. Biomimetic remineralization of resin-dentin bonds is a post-bonding technique that replaces intrafibrillar water as well as resin from water-rich, resin-sparse regions of the hybrid layer with intrafibrillar and extrafibrillar apatite crystallites [29]. By restoring the enzyme exclusion and fossilization properties of mineralized dentin, this proof-of-concept strategy is capable of preserving the longevity of resin-dentin bonds [35].

Both hybrid layers created by etch-and-rinse adhesives (Fig. 4A) and those created by moderately aggressive (Fig. 4B) and aggressive self-etching adhesives are amenable to the biomimetic remineralization approach. For etch-and-rinse adhesives, apatite crystallites can be detected in both extrafibrillar and intrafibrillar spaces after remineralization, as denuded collagen fibrils are present within those hybrid layers. Transmission electron microscopy of remineralized hybrid layers revealed that not the entire hybrid layer was remineralized [29,30]. Regions within hybrid layers that were well-infiltrated by adhesive resin monomers did not undergo remineralization. While the exact pattern of remineralization differed from specimen to specimen, the remineralized regions of poorly resin-infiltrated hybrid layers that (Fig. 5A) in general corresponded well to water-rich, resin sparse regions of the hybrid layers that exhibited extensive silver nanoleakage when specimens were examined immediately after bonding (Fig. 5B), or regions in which denuded collagen had been degraded after *in vivo* aging (Fig. 5C). Along the surface of the hybrid layer, collagen fibrils that were unraveled due to the simultaneous action of cutting and phosphoric acid etching were remineralized in the form of continuous intrafibrillar strands, even when dual biomimetic analogs were employed (Fig. 4C). This is possibly due to loosening of the microfibrillar arrangement and enlargement of the water compartments along the overlap zones of the collagen molecules, thereby enabling the ingress of poly(anionic)

acid-stabilized ACP nanoprecursors into those regions. Conversely, collagen fibrils that were remineralized within the middle part of the hybrid layer contain discrete intrafibrillar apatite platelets (Fig. 4D). These observations are in support of the hypothesis brought forward in Section 3.1 [73].

For self-etch adhesives, remineralization was not usually apparent in mild versions of these adhesives in which partially demineralized dentin was present in the entire thickness of thin (<0.5 μm thick) hybrid layers. Mild self-etch adhesives that contain the functional monomer 10-methacryloydecyldihydrogen phosphate (MDP) are capable of nano-layering with hydroxyapatite [131,132], thereby chemically interacting with the partially demineralized apatite platelets to increase bond durability via the AD-concept [133]. For other mild self-etch adhesives that do not contain MDP, or those containing hydroxyethyl methacrylate that inhibit nanolayering of MDP [134], calcium phosphate deposition on remnant apatite platelets via epitaxial growth may have prevented the ultrastructural aspects of remineralization to be accurately depicted [22] without the use of a high resolution-transmission electron microscope. For moderately aggressive versions of self-etch adhesives, apatite deposition is almost exclusively identified from the completely demineralized part of the hybrid layers. For the aggressive versions of self-etch adhesives, remineralization had been observed within the surface, middle or bottom parts of those thick hybrid layers, depending on the status of resin infiltration and the amount of water present within those hybrid layers. Apart from the use of transmission electron microscopy, the authors have employed a water-soluble fluorescent dye to backfill the water-rich regions within hybrid layers created by an aggressive self-etch adhesive and examined the dye-impregnated resin-dentin interface using confocal laser scanning microscopy [31,32]. In non-remineralized hybrid layers, water-rich regions within those hybrid layers and within the adhesive layer exhibit strong fluorescence of the dye. Interestingly, the intense fluorescence identified from those water-rich regions was quenched after biomimetic remineralization. These observations provide the evidence that biomimetic remineralization of hybrid layers is a process that progressively dehydrates water-rich hybrid layers. Unlike the conventional remineralization approach that proceeds rapidly via epitaxial growth over existing seed crystallites, biomimetic remineralization is a slower process as it involves at least two kinetically driven pathways. Thus, it is important to prevent denuded collagen fibrils from degrading while they are being remineralized. Polyvinylphosphonic acid, a biomimetic analog used to simulate phosphorylated non-collagenous proteins, has been shown to possess matrix metalloproteinase-inhibiting properties that prevent collagen degradation during remineralization [135].

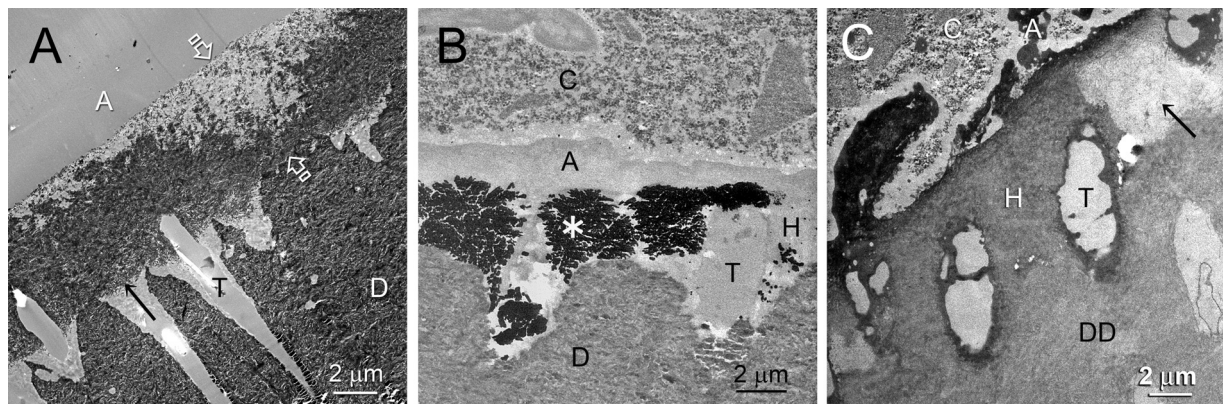
A critical problem encountered with the use of the other methods of matrix metalloproteinase and cysteine cathepsin inhibitors in extending the longevity of resin-dentin bonds is that hydrated, denuded collagen fibrils within the hybrid layer remain flaccid and exhibit weak mechanical properties even with preservation of their integrity [28]. The modulus of elasticity of resin-infiltrated dentin beams increased by 55–118% after biomimetic remineralization as a result of intrafibrillar mineralization of the collagen matrices [136]. A limitation



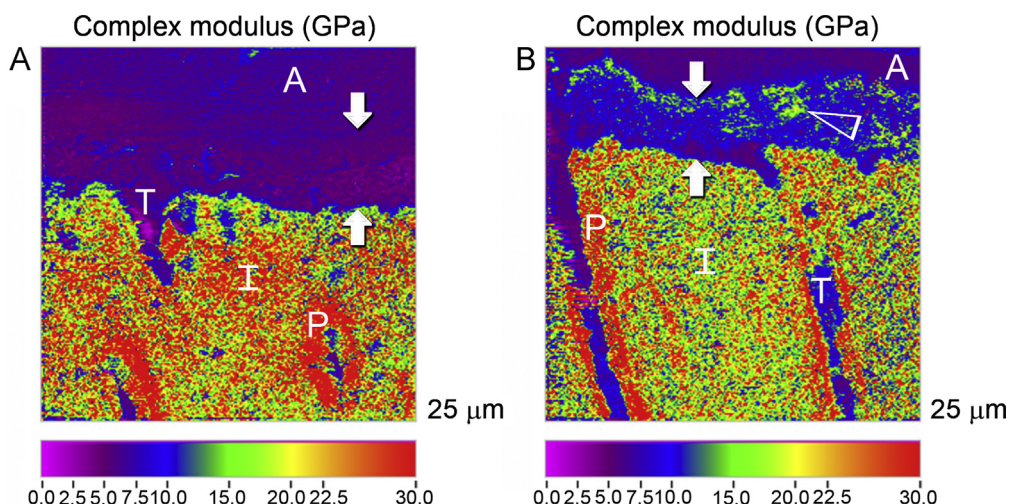
**Fig. 4 – Unstained TEM images illustrating biomimetic remineralization of hybrid layers created by (A). An etch-and-rinse adhesive. A: adhesive; H: apatite-depleted hybrid layer created by phosphoric acid etching for 15 s; T: dentinal tubule; D: mineralized dentin; Asterisk: water-rich part of the hybrid layer remineralized by apatite crystallites. (B) A moderately aggressive self-etch adhesive. FA: filled adhesive; P: partially demineralized base of the hybrid layer; T: dentinal tubule; D: mineralized dentin; between open arrows: water-rich surface of the hybrid layer remineralized by apatite crystallites. (C) High magnification of the surface of a remineralized hybrid layer showing partially unraveled surface collagen fibrils in which their intrafibrillar compartments were filled with continuous strands of apatite mineral. (D) High magnification image taken from the middle part of a remineralized hybrid layer showing collagen fibrils that were remineralized with discrete intrafibrillar apatite platelets.**

of that study, however, is that the resin-infiltrated dentin beams (macro-hybrid layers) were evaluated en masse using three-point bending. As remineralization does not occur in locations of the collagen matrices that are occupied by resins, three-point bending is insensitive in evaluating the localized increases in modulus of elasticity of the remineralized collagen fibrils. Due to the complexity of these regions, the authors have resorted to the use of nanoscopic dynamic mechanical analysis for characterizing the viscoplastic mechanical behavior of resin-infiltrated dentin before (Fig. 6A) and after biomimetic remineralization (Fig. 6B). This is achieved using scanning probe microscopy attached to a triboindenter to

performed rasterized imaging and nanoindentations across an area of interest [137]. In addition, a method of scanning these hybrid layers under hydrated conditions was used to prevent water from evaporating from the specimens during scanning. This involves the application of a layer of ethylene glycol over the specimen surface to prevent water evaporation during a typical 25- to 30-min scanning period for a  $25 \mu\text{m} \times 25 \mu\text{m}$  area of the resin-dentin interface [36]. The combined use of nanoscopic dynamic mechanical analysis and scanning probe microscopy for mapping the resin-dentin interface indicates that the complex modulus ( $E^*$ ), loss modulus ( $E''$ ) and storage modulus ( $E'$ ) of pre-aged hybrid layers



**Fig. 5 – Similarities in the ultrastructural manifestations of biomimetic remineralization; nanoleakage and degradation of aged hybrid layers created by etch-and-rinse adhesives.** Abbreviations: C: resin composite; A: adhesive; H: hybrid layer; D: mineralized dentin base; DD: demineralized dentin base; T: dentinal tubule. (A) Unstained mineralized section of a hybrid layer that had undergone biomimetic remineralization prior to aging for 12 months. Regions within the hybrid layer that were remineralized appear as electron-dense regions (arrow) within the unstained hybrid layer (between open arrowheads). (B) Unstained mineralized section showing highly electron dense, extensive silver deposits (asterisk) within a hybrid layer that had been aged for 12 months prior to immersion in ammoniacal silver nitrate for nanoleakage evaluation. (C) Stained demineralized section of the resin–dentin interface that had been aged for 12 months prior to laboratory processing. Part of the hybrid layer was degraded (arrow) and did not take up the staining observed in the non-degraded part of the hybrid layer.



**Fig. 6 – Nanodynamic mechanical analysis and scanning probe microscopy of the resin–dentin interface created by an etch-and-rinse adhesive (A) before and (B) after biomimetic remineralization.** Only the complex modulus is shown. For Figure B, remineralized regions (open arrowhead) within the hybrid layer exhibit increases in the complex modulus. A: adhesive; between solid arrows: hybrid layer; T: dentinal tubule; I: intertubular dentin; P: peritubular dentin.

created by an etch-and-rinse adhesive ( $E^*$ ,  $3.86 \pm 0.24$ ;  $E''$ ,  $0.23 \pm 0.05$ ;  $E'$ ,  $3.85 \pm 0.24$  GPa) are much lower than those properties of the underlying mineralized dentin ( $E^*$ ,  $19.20 \pm 2.42$ ;  $E''$ ,  $6.57 \pm 1.96$ ;  $E'$ ,  $17.39 \pm 2.0$  GPa). After aging of the resin–dentin bonds for 6 months, the water-rich, resin-sparse regions of the hybrid layers degraded, and the dynamic mechanical properties of those regions further declined ( $E^*$ ,  $0.83 \pm 0.35$ ;  $E''$ ,  $0.88 \pm 0.24$ ;  $E'$ ,  $0.62 \pm 0.32$  GPa). Conversely, when aging of those bonds were performed concomitantly with biomimetic remineralization, those resin-sparse regions became remineralized by intrafibrillar and extrafibrillar apatite crystallites.

The dynamic mechanical properties of those remineralized regions ( $E^*$ ,  $19.73 \pm 3.85$ ;  $E''$ ,  $8.75 \pm 3.97$ ;  $E'$ ,  $16.02 \pm 2.58$  GPa) were restored to those values characteristic of intact mineralized dentin.

#### 4. Biomimetic remineralization of carious dentin

Caries is a dynamic process caused by imbalance between demineralization and remineralization. Carious dentin can

be classified into outer caries-infected dentin and inner caries-affected dentin [138]. Mineral distribution of caries-affected dentin is highly variable and the lesion depth can extend hundreds of micrometer below the excavated surface [139–142]. Contemporary caries management is based on a conservative and preventive approach [143,144]. This minimum invasive philosophy avoids unnecessary tooth sacrifice and leaves caries-affected dentin as the clinical bonding substrate [145]. The bond strengths of dentin adhesives to caries-affected dentin substrate have been reported to be significantly lower than those bonded to noncarious dentin [146–149]. This is attributed to the obliteration of dentinal tubules by acid-resistant mineral crystals, a thicker zone of exposed collagen after the application of the adhesive and the lower stiffness and increased water content of the caries-affected dentin [150–154]. Unlike caries-infected dentin that is denatured, the collagen matrix of caries-affected dentin is not structurally and biochemically different from that found in sound dentin and is physiologically remineralizable [150,155,156].

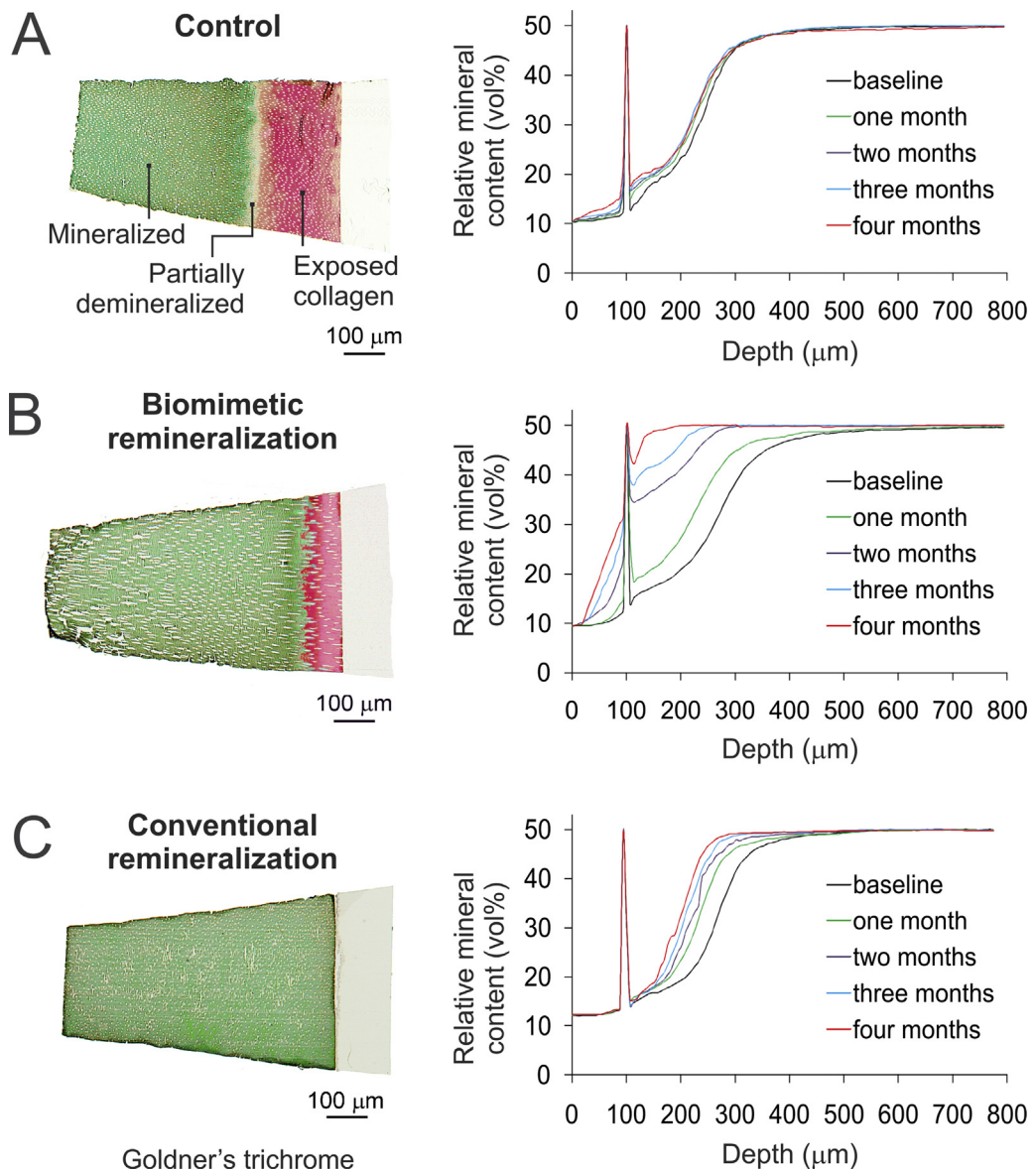
Remineralization of partially demineralized dentin is not new. Reports on remineralization of carious dentin appeared in the dental literature more than half a century ago. The dental literature also contains reports on the use of fluoride-releasing glass ionomer cements and calcium phosphate releasing dental materials [157–160] for remineralizing partially demineralized carious dentin. Intact, non-denatured collagen is remineralizable as long as seed crystallites are present as niduses for heterogeneous nucleation of calcium phosphate phases [161]. Remineralization by epitaxial growth is a thermodynamically favorable process that overcomes the energy barriers of homogeneous nucleation [162]. In nanotechnology terminology, remineralization techniques currently employed in dentistry represent a top-down approach [163]. This approach creates materials using scaled down versions of a bulk material that incorporates nanoscale details of the original material. Partial demineralization of a mineralized collagen matrix by acids derived from bacteria or dentin bonding procedures creates the seed crystallites necessary for this top-down remineralization approach. The orientation of those remineralized crystalline lattices is predetermined by the lattice of the original seed crystallites. However, remineralization does not occur in locations where seed crystallites are absent, as demonstrated with the use of a strontium-based glass ionomer cement on phosphoric acid-etched dentin [16]. This limitation severely restricts apatite-sparse superficial part of caries-affected dentin to be remineralized with a conventional top-down approach.

In biominerization that occurs in nature, there are no seed crystallites present in an organic scaffold. Consequently, biominerization has to proceed via an alternative pathway that involves homogeneous nucleation. One of the mechanisms of homogeneous nucleation involves sequestration of amorphous mineral phase by polyanionic extracellular matrix protein molecules. Fluoride, which has been shown to enhance conventional remineralization via the top-down approach, is not employed by nature as a functional motif for biominerization. In addition, fluoride-based dentin remineralization strategies also result in hypermineralization of the lesion surface [164,165] and prevent effective

remineralization of the deeper parts of the carious lesion [166,167]. Homogeneous nucleation is not as thermodynamically favorable and involves alternative kinetically driven protein/polymer-modulated pathways for lowering the activation energy barrier for crystal nucleation via sequential steps of phase transformations [18,168]. In nanotechnology terminology, such pathways are examples of a bottom-up approach [163], wherein nanoscale materials are created using a particle-based self-assembly process [37,47].

To evaluate the efficacy of a bottom-up approach to remineralize carious dentin, the authors created, via pH cycling, 250–300 μm thick artificial carious dentin lesions, each with a gradient of demineralization from the surface to the base of the lesion (Fig. 7A) [17]. The artificial carious lesions were randomly divided into two groups and remineralized for 4 months using a classical top-down approach (Fig. 7B) or a non-classical bottom-up approach (Fig. 7C). In the bottom-up biomimetic remineralization strategy, two analogs were used to mimic the functions of matrix proteins in biominerization. Polyacrylic was employed as an apatite nucleation inhibition agent to stabilize ACP nanoprecursors derived from set Portland cement or mineral trioxide aggregate (MTA) and simulated body fluid (SBF). Polyvinylphosphonic acid was used as a template to recruit the ACP nanoprecursors to the gap zones of the collagen fibrils, where they nucleate and self-assemble into hierarchically arranged apatite nanocrystals within the gap zones of the collagen fibril. No analogs were used in the control top-down approach. To quantitatively assess the changes of mineral density before and after mineralization, micro-computed tomography was used to examine the mineralized collagen scaffold non-destructively in three dimensions, over a period of 4 months. To delineate between intrafibrillar and extrafibrillar apatite deposition within the collagen fibrils, transmission electron microscopy was used to examine the dimension and hierarchy of apatite deposition within the mineralized collagen matrix. It was found that the top-down approach could only mineralize the base of the partially demineralized scaffold where remnant seed crystallites were abundant. Minimal mineralization was observed along the surface of the scaffold, wherein extrafibrillar mineralization was predominantly observed. Conversely, the entire partially demineralized scaffold including apatite-depleted collagen fibrils was mineralized by the bottom-up approach, with evidence of both intrafibrillar and extrafibrillar mineralization. Similar results were obtained when polyvinylphosphonic acid was replaced by sodium trimetaphosphate as a biomimetic analog of phosphorylated noncollagen proteins to recruit polyacrylic acid-stabilized ACP nanoprecursors [33].

In a more recent study, Burwel et al. applied the PILP system (polyaspartic acid only as the apatite nucleation inhibition agent) to remineralize 140 μm deep artificial caries lesions for 7–28 days [79]. The authors examined the nanomechanical properties of the hydrated artificial lesions before remineralization and at different time points after biomimetic remineralization. Prior to remineralization, the hydrated artificial carious-like lesions had a low reduced elastic modulus of 0.2 GPa extending about 70 μm into the lesion, with a sloped region to about 140 μm where the reduced elastic modulus reached that in normal dentin (18–20 GPa). After 7 days of biomimetic remineralization, specimens recovered



**Fig. 7 – Comparison of the status of remineralization of artificial caries-like lesions created in dentin over a 4-month period.** For each figure, the light microscopy section on the left was stained with Goldner's trichrome; mineralized collagen was stained green while exposed collagen was stained red. The chart on the right represents the changes in overall mineral profile over time for the 250–300  $\mu\text{m}$  thick artificial caries lesion, as determined by micro-computed tomography. (A) Control (no remineralization after immersion in a non-mineralizing medium). (B) Conventional remineralization (top-down approach) using a medium consisting of 1.5 mM CaCl<sub>2</sub>, 0.9 mM KH<sub>2</sub>PO<sub>4</sub>, 130 mM KCl, 20 mM HEPES and 5 mM NaN<sub>3</sub>. (C) Biomimetic remineralization (bottom-up approach) using a medium consisting of 136.8 mM NaCl, 4.2 mM NaHCO<sub>3</sub>, 3 mM KCl, 1 mM K<sub>2</sub>HPO<sub>4</sub>·3H<sub>2</sub>O, 1.5 mM MgCl<sub>2</sub>·6H<sub>2</sub>O, 2.5 mM CaCl<sub>2</sub>, 0.5 mM Na<sub>2</sub>SO<sub>4</sub>, 5 mM NaN<sub>3</sub>, supplemented with 500  $\mu\text{g}/\text{mL}$  polyacrylic acid and 200  $\mu\text{g}/\text{mL}$  of polyvinylphosphonic acid. Set white Portland cement was used as the source of sustained calcium and hydroxyl ions release for initial formation of amorphous calcium phosphate nanoprecursors, in the presence of phosphate ions derived from the liquid medium.

mechanical properties in the sloped region by 51% compared to the artificial lesion. Between 7 and 14 days, recovery of the outer portion of the lesion continued to a level of about 10 GPa with 74% improvement. After 28 days of PILP remineralization, there was 91% improvement of the reduced modulus compared to the artificial lesion. During remineralization,

intrafibrillar mineral increased and crystallinity improved with intrafibrillar mineral exhibiting the orientation found in normal dentin or bone. These researches generated important information to address the critical barrier to progress in remineralization of caries-affected dentin and shifted existing paradigms by providing a novel method of remineralization

based on a nanotechnology-based bottom-up approach. All these results support the translation of the proof-of-concept biomimetic strategy into a clinically relevant delivery system for remineralizing caries-affected dentin created by micro-organisms in the oral cavity.

## 5. From proof-of-concept to clinical translation – potential strategies

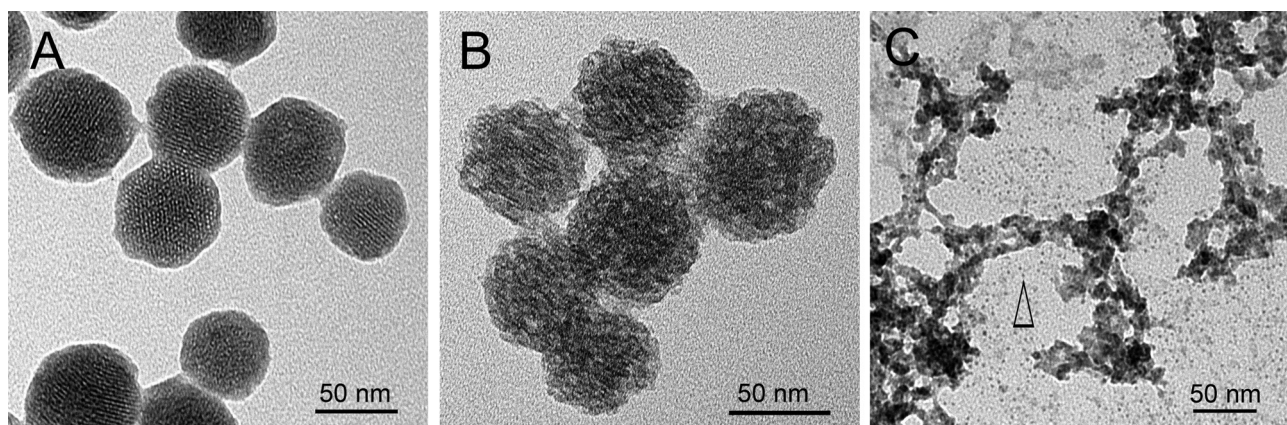
Although the particle-based biomimetic remineralization strategy based on the PILP process demonstrates great potential in remineralizing faulty hybrid layers or caries-like dentin, the strategy remains a proof-of-concept. To-date, studies published by various research groups utilized sectioned dentin slabs of the resin-dentin interface or artificially created carious lesion and not the whole tooth for remineralization. These slabs were immersed in a metastable remineralization solution containing supersaturated calcium and phosphate ions, as well as the biomimetic analogs responsible for generating fluidic ACP nanoprecursors. Whereas poly(anionic acid)-stabilized ACP nanoprecursors can readily infiltrate sideways into the interfacial defect in a sectioned slab, this is not possible clinically. This is because the area occupied by the bonded dentin or carious dentin is segregated from the oral environment by a restorative material and only a very small part of the interface is exposed along the cavosurface margin. It is unrealistic for biomimetic remineralization to be accomplished through the use of a mouthrinse or a topically applied delivery system such as a remineralization paste or gel, wherein the polymer-stabilized-ACP nanoprecursors have to diffuse thousands of micrometers into the adhesive or caries-affected dentin within a restoration. For effective remineralization, the critical components of the biomimetic scheme (i.e. calcium and phosphate source and biomimetic analog/analogs) have to be incorporated into a dentin adhesive or restorative material. These critical components should be able to be released from polymerized resins. This necessitates incorporation of hydrophilic resin monomers in the adhesive or resin composite to facilitate water, ion and nanoparticulate diffusion. Although some calcium phosphate phases may precipitate within the adhesive or restorative material, it is critical that the bulk of the remineralization components diffuse through the adhesive joint to backfill water-rich regions of faulty hybrid layers or the caries-affected dentin with intrafibrillar apatite crystallites. To increase the durability of the restorations, there should also be sustained release of these critical components to remineralize tooth-restoration interfaces that have been subjected to secondary caries. These requirements impose considerable challenges to the translation of a scientifically sound concept into a clinically applicable approach, without adding extra steps into current bonding/restorative protocols.

The idea of incorporating calcium phosphate or bioactive glass particles into resins to develop composites with remineralizing capabilities has been explored by different researchers [14,15,169–174]. One group of composites contained ACP fillers, with the zirconia-stabilized ACP particles having a median diameter of 7.4 µm (ranging from 0.3 to

80 µm) [175]. However, in another study, the zirconia-stabilized ACP particles were larger, with a median diameter of 55 µm (ranging from submicron to 100 µm) [176]. These experimental calcium phosphate-containing composites are promising for remineralization of enamel because ACP is a precursor that forms initially and eventually transforms into apatite [177]. More recently, ACP nanoparticles with a much smaller diameter (116 nm) have been synthesized using a spray-drying technique [178]. Being solid nanoparticles that are used as fillers within a polymerized resin matrix, they are incapable of being released from the composite, or diffusing through the adhesive into faulty hybrid layers or adhesive-bonded caries-affected dentin [179]. Nevertheless, calcium and phosphate ions released from these ACP nanoparticles can diffuse through water channels within the hydrophilic adhesive layer. Amorphous calcium phosphate nanoparticles have also been incorporated into experimental dentin adhesives [180,181]. In this case, the ACP nanoparticles are in closer proximity to the hybrid layer or caries-affected dentin. As the adhesive flows into dentinal tubules after the latter are rendered patent by acid-etching, incorporating ACP nanoparticles into a dentin adhesive may be more efficacious in remineralizing thick, partially demineralized caries-affected dentin, as this shortens the path of diffusion of the calcium and phosphate ions, and reduces the chance of precipitation of calcium phosphate phases along the path of diffusion.

The use of ACP nanoparticle-containing dentin adhesive is innovative. These experimental adhesives are very likely to be able to remineralize partially demineralized dentin by epitaxial deposition of calcium and phosphate phases over remnant apatite seed crystallites. However, the use of solid state ACP nanoparticles is not compliant with the non-classical theory of particle-based crystallization. Firstly, the ACP nanoparticles are still larger than the gap zone (40 nm) between the collagen molecules. Secondly, they do not possess the fluidic characteristics of amorphous mineral particles generated by a PILP process that enable them to infiltrate the intrafibrillar water compartments of collagen fibrils. Thirdly, although they release calcium and phosphate ions, they lack biomimetic analogs to sequester those ions into ACP prenucleation clusters. Thus, it is difficult to envisage how the use of preformed ACP nanoparticles can remineralize apatite-free denuded collagen fibrils within a faulty hybrid layer, or the superficial part of a caries lesion. It is pertinent to recapitulate that intrafibrillar mineralization of seed-crystallite free collagen is the hallmark of contemporary biomimetic collagen mineralization strategies. To date, intrafibrillar remineralization of dentin collagen has not been reported with the use of experimental adhesives containing solid ACP nanoparticles.

Another interesting form of stabilized ACP is casein phosphopeptide-ACP (CPP-ACP) [182]. Tryptic digestion of caseins derived from milk produces phosphopeptides that contain cluster of phosphoserine residues. The phosphorylated residues in CPP-ACP help to stabilize calcium and phosphate ions through the formation of complexes [183]. Theoretically, CPP can serve as a biomimetic analog of phosphorylated dentin noncollagenous proteins for recruiting ACP nanoprecursors into the gap zones of collagen fibrils. Although CPP-ACPs have been reported to be successful in



**Fig. 8 –** Unstained TEM images of (A) mesoporous silica nanoparticles with a hexagonal array of mesopores. (B) Loading of polyacrylic acid-stabilized amorphous calcium phosphate into the mesopores. (C) Intentional dissolution of the silica framework of the loaded mesoporous silica nanoparticles in 0.05 N NaOH results in the release of amorphous calcium phosphate particles and prenucleation clusters (open arrowhead). Although the silica dissolution process is far removed from clinical application, the technique illustrates that the calcium phosphate phase within the mesoporous remains amorphous and has not been converted into apatite (selected area electron diffraction not shown).

the remineralization of non-cavitated demineralized enamel [184,189], evidence is lacking in terms of their ability to introduce intrafibrillar minerals into apatite-depleted dentin collagen fibrils.

A potential delivery strategy is to produce poly(anionic) acid-stabilized ACP prenucleation clusters and store them as “cargos” in mesoporous silica nanofillers (Fig. 8). These mesoporous silica nanofillers may be incorporated in dentin adhesives as controlled release devices for the delivery of the ACP prenucleation clusters, or coalesced ACP fluidic PILP phases, into faulty hybrid layers or caries-affected dentin. The synthesis of mesoporous silica is based on the formation of liquid-crystalline mesophases of amphiphilic molecules (surfactants) that serve as templates for the *in situ* polymerization of orthosilicic acid. The surfactants are then removed to produce nanoparticles with variable patterns of mesoporous channels. The mesoporous silica nanoparticles currently synthesized by the authors have a median particle diameter of 50 nm. They are composed of amorphous silica with a hexagonal system of mesopores, as determined by small angle X-ray diffraction. They have an extremely large specific surface area ( $>1000\text{ m}^2/\text{g}$ ), as determined by nitrogen adsorption (Brunauer–Emmett–Teller method). This value is much higher than the specific surface area of the ACP nanoparticles ( $17.8\text{ m}^2/\text{g}$ ) described previously [178]. The pore volume and pore size of the mesoporous silica nanoparticles synthesized in the authors’ laboratory, as determined by the Barrett–Joyner–Halenda method, is close to  $1\text{ cm}^3/\text{g}$  and 3.5 nm, respectively. As the average diameter of ACP prenucleation clusters  $0.87 \pm 0.2$  nm when they are first formed [55], they can readily enter the mesopores of the silica nanofillers. As the internal surface of the mesopores contains silanol groups that are negatively charged, different methods are available for functionalization of the internal pores to render them positively charged to enhance electrostatic attraction

of the poly(anionic)acid-stabilized ACP nan precursors. The advantage of using mesoporous silica nanofillers for delivery of ACP nan precursors is that silica is only sparingly soluble in an acidic environment. This prevents the silica nanoparticles and their loaded contents to be solubilized by acidic resin monomers present in some dentin adhesives. Another potential strategy currently contemplated by the authors is the use of calcium and phosphate-containing mesoporous bioactive glass nanoparticles synthesized by a sol-gel route for the loading and delivery of biomimetic analogs. The ion-releasing nanoparticles provide the source of calcium and phosphate for *in situ* generation of prenucleation clusters. Research in the development of these novel nanotechnologies for clinical translation of the concept of biomimetic remineralization of dentin is in order.

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