DENTINE MATRIX METALLOPROTEINASES AS POTENTIAL MEDIATORS OF DENTINE REGENERATION

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Abstract

Matrix metalloproteinases (MMPs) have been implicated not only in the regulation of developmental processes but also in the release of biologically active molecules and in the modulation of repair during tertiary dentine formation. Although efforts to preserve dentine have focused on inhibiting the activity of these proteases, their function is much more complex and necessary for dentine repair than expected. The present review explores the role of MMPs as bioactive components of the dentine matrix involved in dentine formation, repair and regeneration. Special consideration is given to the mechanical properties of dentine, including those of reactionary and reparative dentine, and the known roles of MMPs in their formation. MMPs are critical components of the dentine matrix and should be considered as important candidates in dentine regeneration.

Keywords
Dentine; enamel; periodontal ligament; tooth; dental regenerative repair; matrix metalloproteinases; dental physiology; signalling molecules/growth factors

Tissue structure and function in regeneration

The tightly correlated structure-function relationship in physiological dentine requires that regeneration of the tissue approximate primary dentine as closely as possible. In other words, dentine regeneration necessitates the preservation of the mechanical properties of the tissue rendered by its ultrastructure. Dentine is a mineralised, but vital, tissue housing the cellular processes of odontoblasts within dentinal tubules and responsible for dissipating the stresses of mastication (Goldberg et al., 2011). These forces must be transferred from a stiff (enamel, 96 % mineral, by weight) to a more elastic (dentine, 70 % mineral, by weight) material. Dentine matrix is composed of collagenous (86 % type I, as well as types III, V and VI) and non-collagenous proteins.

Dentine is capable of limited repair following pulpal insult from carious lesions, cavity preparations, erosion and restorative dental materials. Dentine repair through tertiary dentine
deposition results in inward growth of the circumpulpal dentine layer constricting the pulp chamber and root canals. Tertiary dentine does not preserve the precise tubular system of physiological dentine. Minimal trauma that spares the underlying odontoblast layer or Hoehl’s cells results in reactionary dentine formation, in the form of tubular orthodentine or atubular, bone-like osteodentine (Goldberg et al., 2015). Reactionary dentine has reduced hardness and lower elasticity, despite being heavily mineralised (Charadram et al., 2013; Senawongse et al., 2006). More severe insults that result in cellular death require progenitor cells to repair the defect by depositing reparative dentine in the form of a dentineal bridge. It is unclear whether the amount of tertiary dentine present in a tooth affects its function.

Dentine hardness is directly proportional to the mineral density of the tissue and has a linear correlation with lack of elasticity. The mineral density varies across the tissue, with the most mineral dense, hardest and least elastic dentine near the dentineal tubule edge, called peritubular dentine (MacDougall et al., 1992). Between tubules lies the more elastic intertubular dentine. Hardness is lowest near the DEJ, peaks in circumpulpal dentine and again decreases towards the pulp (Wang and Weiner, 1997). Dentineal tubules become more abundant, closer together and larger as they approach the odontoblastic cell body (Pashley et al., 1985). Changes in dentineal tubule density correlate with changes in the ratios of intertubular and peritubular dentine, thus affecting the hardness and the mechanical properties of the tissue.

To the authors’ knowledge, no studies have shown whether MMP activity affects tubular density and the physical and mechanical properties of tertiary dentine. Due to the importance of these factors to dentine function, future regenerative models should consider using them as indicators of success.

**What’s dentine regeneration and what does it entail?**

Dentine repair and regeneration invariably depend on pulp vitality. Regenerative endodontics has used this concept in the treatment of immature teeth with open apices. The dental pulp is a loose connective tissue composing the core of the tooth structure. It consists of cellular (fibroblasts, odontoblasts, immune cells, neurovascular networks) and extracellular (collagen, fibronectin, glycoproteins) components that exist in close relationship with dentine (Goldberg et al., 2011). Vital pulp therapies promote the regeneration of vascularised, innervated dental pulp able to support odontoblast differentiation and dentine neogenesis for the completion of root formation (Huang, 2011). Dentine regeneration in such a form fails to address the loss of clinical crown due to caries. Studies do not clarify the amount of dentine that can potentially be replaced or whether these equal the amount lost due to disease. Furthermore, due to the acellular nature of the enamel, regeneration of this tissue poses even more challenges than dentine regeneration (Pandya and Diekwisch, 2019).

Tissue regeneration demands, at minimum, the potential for replacement of cellular and matrix components through proliferation (Krafts, 2010). Dentine regeneration will require replacement of carious dentine matrix by newly formed proliferating odontoblasts and differentiating dental pulp stem cells. To this end, scaffolds along with biological cues pose the most promise allowing for subsequent cellular infiltration, matrix deposition and mineralisation.
Alginates, chitosan, hyaluronic acid, collagen, gelatine, fibrin, silk and hydrogel scaffolds have been effective in supporting dental pulp cell growth and differentiation (Inuyama et al., 2010; Panseri et al., 2016; Prescott et al., 2008; Vagropoulou et al., 2021; Yang et al., 2015). Although these scaffolds promise to promote pulpal regeneration, no studies have shown appreciable levels of mineralisation to replace the amount of dentine lost due to caries. Moreover, the time required to regenerate the bulk tissue destroyed by caries would be extensive and clinically unacceptable. The tooth may need to be out of occlusion to prevent excessive forces from disrupting the regeneration process. Also, it would probably require a full-coverage restoration for better long-term prognosis.

**Dentine MMPs**

MMPs are a family of 28 modular endopeptidases involved in extracellular matrix remodelling and regulation of extracellular signalling networks guiding inflammation, bone development and angiogenesis, among others (Löffek et al., 2011). They are produced by the cellular components of soft and hard tissues, including epithelial cells, endothelial cells, fibroblasts, osteoclasts, osteoblasts, hypertrophic chondrocytes, chondroclasts, inflammatory cells and odontoblasts (Goldberg et al., 2003; Ortega et al., 2003; Sulkala et al., 2002). Most MMPs contain a propeptide domain, responsible for preserving the latent conformation of the enzyme; a zinc-binding catalytic domain, responsible for their proteolytic function; a haemopexin-like domain, responsible for protein-protein interactions. These zymogens are activated by a variety of mechanisms, both proteolytic and nonproteolytic, that make them suitable for their different functions (Van Wart and Birkedal-Hansen, 1990). They can be classified by their substrates, which are largely determined by specificity-determining positions on their catalytic domain (Ratnikov et al., 2014). The MMPs found in dentine include MMP-2, -3, -7, -8, -9, -13, -14, -20, -23 and -25 (Eckhard et al., 2015; Goldberg et al., 2003; Hall et al., 1999; Loreto et al., 2014; Mazzoni et al., 2007, 2018; Sulkala et al., 2002, 2007; Xu et al., 2016) (Table 1).

The dynamics of MMP activation and inhibition remain ill-defined, albeit one of the most important features of tissue remodelling in response to disease. Historically, MMP proteolytic activity has been associated with tissue destruction. MMP-8 has been identified as one of the main collagenases in human dentine involved in the caries process, while increases in MMP-14 activity have also been associated with a response to caries, although its exact role remains unclear (Charadram et al., 2012; Sulkala et al., 2007; Tjäderhane et al., 2015). These endogenous MMPs may be released from the extracellular matrix and/or activated by the caries process and are likely to sustain the disease through their enzymatic activity. Moreover, their loss of function has been associated with decreased caries risk, as in the case of MMP-13 (Loreto et al., 2014; Tannure et al., 2012).

Efforts to slow or prevent disease have focused on inhibiting MMP activity. Failure of resin-based restorations due to hybrid layer degradation has been attributed to the enzymatic degradation of collagen fibrils by MMPs. To date, the use of exogenous MMP inhibitors, such as tetracycline antibiotics and chlorhexidine, have improved the clinical outcomes of resin-bonded restorations by preserving the hybrid layer and bond strength (Breschi et al., 2018; Gendron et al., 1999; Hanemaaijer et al., 2001).
Endogenous TIMP-1 to -4 have been identified in human dentine. TIMP-2 expression increases during the caries process, although this increase is concurrent with enhanced MMP expression (Charadram et al., 2012; Goldberg et al., 2003). The significance of their co-expression has yet to be determined, although MMP/TIMP ratios and substrate-inhibitor specificity may explain and control MMP activity in tissues. TIMP signalling independent of its MMP-inhibitory action has also been proposed. TIMP-1 and -2 expression are important components of scaffolds used in regenerative endodontic procedures (Koh et al., 2007; Suresh et al., 2018). Furthermore, growth factors that contribute to pulp cell proliferation are known to stimulate TIMP expression (Chang et al., 2017). In such cases, MMP co-expression may be explained by MMP counter-regulatory action on TIMP signalling. Ultimately, regeneration requires a balance in matrix turnover. Leveraging MMP activity would require a better understanding of MMP-TIMP protein interactions and biological functions.

MMPs play a crucial role during tooth development. MMP-2 and -9 are the earliest MMPs to be expressed and may be involved in the degradation of the basement membrane, marking the onset of the terminal differentiation of ameloblasts and odontoblasts (Heikinheimo and Salo, 1995). While important at early stages, MMP-2 (and MMP-20) loss of function results in higher levels and broader distribution of non-collagenous proteins known to promote dentine mineralisation (Bourd-Boittin et al., 2005). Still at later stages of development, the loss of function of other MMPs, such as MMP-14, results in root dentine defects (Xu et al., 2016). Evidence indicates that MMPs are not redundant proteins with interchangeable functions. Their specificities of action can mediate tissue development and their dysregulation carries significant implications for tissue integrity.

Dentine regeneration and tertiary dentine formation depend on the appropriate signalling molecules’ potential for cell differentiation, migration and proliferation – and appropriate remodelling of newly deposited dentine matrix. MMPs are crucial for these processes. Thus, it might not be feasible to completely abolish their proteolytic activity to achieve regeneration. In fact, certain MMPs, although present during disease, may play a protective role by facilitating the repair process. MMP-3 exhibits optimal proteolytic activity and calcium affinity in a pH-dependent manner (range pH 5.3-5.5). Thus, its optimal proteolytic activity lies at the critical pH of demineralisation of enamel and dentine, such as those found in carious environments (Abou Neel et al., 2016; Wilhelm et al., 1993). MMP-3, however, has been associated with angiogenesis and reparative dentine deposition (Zheng et al., 2009).

**Leveraging MMPs for dentine regeneration**

MMPs have been implicated not only in the regulation of developmental processes, but also in the release of biologically active molecules and modulation of repair during tertiary dentine formation (Chaussain-Miller et al., 2006; Sternlicht and Werb, 2001). MMPs clear the way for incoming progenitor cells as well as activate growth factors responsible for angiogenesis, immune regulation and cellular differentiation. MMP activity in response to disease may arise from exogenous (bacterial products) and endogenous (immune cells) sources as well as from dentine matrix reservoirs. Bacterial products from carious lesions may also lead to signalling cascades that activate MMP secretion by odontoblasts.

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Odontoblasts adjacent to reactionary dentine express high levels of MMP-2, which has been associated with increased proteolytic activity in this zone (Charadram et al., 2012). This activity could be responsible for the maturation of collagen fibres and the initiation of mineral formation in the newly formed dentine (Jontell and Linde, 1983).

MMPs contribute to debris clearance and new tissue formation. Animal models of tissue regeneration support the importance of these enzymes in the regenerative process. MMP upregulation is one of the earliest steps in newt limb regeneration after amputation. MMP activity is essential and its inhibition stunts limb regeneration (Vinarsky et al., 2005). MMP-9, -3 and -13 were the most highly expressed in this particular model (Miyazaki et al., 1996). Similarly, pulp injury results in an inflammatory response characterised by the invasion of polymorphonuclear cells and the release of proteases, such as MMP-9 (Gusman et al., 2002; Mente et al., 2016). These MMPs contribute to the degradation of exposed carious dentine, angiogenesis and cell migration, thus activating pathways that lead to tertiary dentine deposition (Charadram et al., 2012). MMP-9 is currently being used in endodontics as a diagnostic and prognostic measure of pulpal inflammation to help guide clinical decisions (Mente et al., 2016; Sharma et al., 2020; Zehnder et al., 2011).

In vivo, MMP-3 treatment results in reversal of mild irreversible pulpitis partly due to its anti-inflammatory properties. These properties include the inhibition of IL-6 expression and a decrease in macrophage and antigen-presenting cell infiltration (Eba et al., 2012). Furthermore, MMP-3 is able to stimulate CTGF production, independent of its proteolytic activity, thus enhancing the migration of dental pulp cells (Muromachi et al., 2012; VanHook, 2008). MMP-3 has also been localised to endothelial cells and stimulates angiogenesis and reparative dentine deposition in pulp injury models, in vivo (Zheng et al., 2009). Increased MMP-3 activity is not found in irreversibly injured pulps further supporting a regenerative role for the protein (Gusman et al., 2002). Future models of dentine regeneration should especially consider MMP-3 as a mediator of regeneration.

Lining the periphery of the pulp chamber is a palisade layer of odontoblasts responsible for the formation of dentine. During dentine deposition, these cells secrete bioactive molecules that orchestrate the mineralisation of the tissue. Similarly, in response to dentine destruction resulting from attrition, carious exposures and chemical insults, these cells are stimulated to secrete new dentine. However, during regeneration and repair, the chemical signals and cellular dynamics responsible for primary dentine formation are absent. Thus, the tissue relies on bioactive molecules to stimulate the proliferation, migration and differentiation of cells responsible for dentine regeneration and repair (Abbass et al., 2020).

Mature dentine is a reservoir of bioactive molecules once involved in its physiological deposition. As such, the tissue contains within it a defence mechanism for environmental insults. TGF-β1, PDGF-AB, VEGF, PlGF and FGF2 are found sequestered in dentine (Bègue-Kirn et al., 2004; Roberts-Clark and Smith, 2000; Zhao et al., 2000) (Fig. 1). Solubilisation of these growth factors promote angiogenesis, odontoblast differentiation and tertiary dentine deposition; ultimately resulting in dentineal bridges that are thicker, denser and structurally similar to physiological dentine (Galler et al., 2011; Liu et al., 2021). Furthermore, isolation of these factors from plasma can sustain tooth-bud cell viability and
has resulted in the regeneration of teeth in porcine animal models (Yang et al., 2012). Dentine conditioning with EDTA is a promising strategy to growth factor release and stimulation of gene expression that potentiates odontoblast differentiation (Sadaghiani et al., 2016). MMPs provide another method for the release of these factors from dental tissues. The use of phosphoric acid etch-and-rinse adhesive systems has the potential to expose these proteases while maintaining their function (Tezvergil-Mutluay et al., 2013). Promotion of pulp healing has been associated with many of these proteases. Direct pulp capping agents consisting of MMP-digested dentine matrix components have shown similar regenerative properties in vivo (Okamoto et al., 2018). These studies identified MMP-1, -9, -13 and -20 as promoters of pulpal healing. Endogenous MMP activity has been leveraged in the delivery of growth factors from scaffolds in models of tissue regeneration. In a recent study, Huang et al. (2020) have engineered a scaffold containing an MMP-2 cleavage site and heparinbinding sites to bind growth factors. Such hydrogel scaffolds would be useful for the release of growth factors when activated by MMP2 in vivo.

Concentrated growth factors from venous blood, consisting of PDGF-BB, TGF-β1, VEGF and others, have been shown to inhibit proinflammatory cytokine release by dental pulp cells and result in the regeneration of dentine-pulp complex, in vivo (Xu et al., 2019). Immunomodulation of the pulpal inflammatory response to injury has also been achieved using melatonin-induced dental pulp TGF-β secretion (García-Bernal et al., 2020). Overexpression of PDGF-BB promotes regeneration through recruitment of dental pulp stem cells, enhancing their proliferation and odontogenic differentiation as well as enhancing pulp angiogenesis (Zhang et al., 2017). Similarly, bioactive pulp-capping agents containing BMPs, such as BMP-2 and -4, as well as FGF2 induce cell differentiation and tertiary dentine deposition (Ishimatsu et al., 2009; Nakashima, 1994).

Cells, signalling molecules and scaffolds form the tissue engineering triad. Successful scaffolds must persist in tissues for long enough to allow for cellular colonisation, after which they must undergo enzyme-mediated degradation (Murphy et al., 2013). MMPs have been leveraged in the timely removal of hydrogel scaffold systems, thereby sustaining cell colonisation and long-term proliferation, qualities essential for neovascularisation and angiogenesis (Lutolf et al., 2003; Turturro et al., 2013). Native cells within the regenerating tissues are crucial for maintaining the scaffolds. Secretion of anti-inflammatory cytokines, such as IL-10, induces TIMP expression and prevents scaffold degradation by MMPs (Ye et al., 2011). Conversely, inflammatory cells within the regenerating tissues are also known to secrete MMPs responsible for remodelling these scaffolds and newly deposited extracellular matrices (Hong et al., 2017).

As stated above, MMPs are instrumental in the formation of tertiary dentine. These enzymes function in the maturation of the dentine collagen matrix, enhancing the bioavailability of signalling molecules, and the cellular events that result in dentine repair. Activation of signalling molecules is particularly important in the mineralisation process. In addition to maturation of the collagenous components of the dentine matrix, MMPs are involved in the maturation and function of a group of non-collagenous proteins called SIBLINGs. Proteins in this family include OPN, DMP1, DSPP, MEPE and BSP2.
SIBLINGs bind specifically to MMPs, activating both latent MMPs and TIMP-inhibited MMPs (Fedarko et al., 2004). The known partners include OPN/MMP-3, DMP1/MMP-9 and BSP/MMP-2. MMP-2 can also cleave DMP1 to release biologically active peptides and MMP-9 can cleave DSPP into DSP and DPP (Chaussain et al., 2009; Yuan et al., 2017). DSPP and MEPE have no MMP partner known to date (Ogubreke and Fisher, 2004). SIBLINGs represent most of the phosphorylated extracellular matrix proteins in dentine and have been implicated in mineralisation and odontoblast differentiation (Almushayt et al., 2006; Eapen and George, 2015). The relationship between these two protein families may represent an opportunity for dentine repair.

Conclusion

MMPs are a family of proteinases responsible for matrix maturation and remodelling as well as modulation of non-collagenous proteins and signalling molecules that result in dentine repair. These proteinases play a developmental role during disease and repair, having functions as diverse as the protein family itself. The existing literature shows that MMPs do not all have similar properties. Some can have a more destructive while others a more regenerative function. To date, dentine regeneration has been approached in either of two ways: (1) by stimulating dentine deposition by stem cells; (2) by generating scaffolds to facilitate the deposition of mineralised tissue at the site of the dentine defect. The spatiotemporal regulation of MMP expression, their multifunctional properties and their ability to autoregulate make these multifaceted proteins ideal candidates for stimulation of dentine regeneration. Future research should focus on leveraging the properties of these enzymes in the promotion of dentine regeneration.

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List of Abbreviations

| Abbreviation | Description                        |
|--------------|------------------------------------|
| BSP2         | bone sialoprotein                  |
| CTGF         | connective-tissue growth factor    |
| DEJ          | dentineoenamel junction            |
| DMP1         | dentine-matrix protein 1           |
| DPP          | dentine phosphoprotein             |
| DSP          | dentinesialoprotein                |
| DSPP         | dentine sialophosphoprotein        |
| EDTA         | ethylenediaminetetraacetic acid    |
| FGF2         | fibroblast growth factor 2         |
MEPE  matrix extracellular phosphoglycoprotein
MMP  matrix metalloproteinase
OPN  osteopontin
PDGF-AB  platelet-derived growth factor AB
PIGF  placenta growth factor
SIBLING  small integrin-binding ligand, N-linked glycoprotein
TGF-β1  transforming growth factor-β1
TIMP  tissue inhibitors of metalloproteinases
VEGF  vascular endothelial growth factor

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Fig. 1. Dentine is a reservoir of bioactive molecules.
MMPs are a family of proteinases responsible for dentine repair during environmental insults to the tooth, such as dental caries. MMPs orchestrate the activation of non-collagenous proteins and signalling molecules that, in turn, stimulate angiogenesis as well as proliferation, differentiation and migration of dental pulp cells during dentine repair.
Endogenous MMPs have been associated with caries susceptibility and dentine repair potential. Evidence indicates that MMPs are not redundant proteins with interchangeable functions.

| Dentine MMP | Functions                                                                 | References                                                                 |
|-------------|---------------------------------------------------------------------------|---------------------------------------------------------------------------|
| MMP-2       | Associated with developmental processes and caries; known to bind bone sialoprotein; cleaves dentine matrix protein 1 | Charadram et al., 2012, Goldberg et al., 2003, Mazzoni et al., 2007       |
| MMP-3       | Associated with angiogenesis and reparative dentine deposition; known to bind osteopontin | Eba et al., 2012, Hall et al., 1999, Muromachi et al., 2012, VanHook, 2008, Zheng et al., 2009 |
| MMP-7       | Role remains unclear                                                      | Mazzoni et al., 2018                                                     |
| MMP-8       | Associated with caries                                                    | Sulkala et al., 2007                                                     |
| MMP-9       | Associated with developmental processes and caries; known to bind dentine maxtrix protein 1; cleaves dentine sialophosphoprotein | Charadram et al., 2012, Goldberg et al., 2003, Mazzoni et al., 2007, Okamoto et al., 2018 |
| MMP-13      | Associated with increased caries risk                                     | Loreto et al., 2014, Okamoto et al., 2018, Tannure et al., 2012           |
| MMP-14      | Role remains unclear                                                      | Xu et al., 2016                                                          |
| MMP-20      | Role remains unclear                                                      | Okamoto et al., 2018, Sulkala et al., 2002                               |
| MMP-23      | Role remains unclear                                                      | Eckhard et al., 2015                                                     |
| MMP-25      | Role remains unclear                                                      | Eckhard et al., 2015                                                     |