

Matrix metalloproteinase 20 promotes a smooth enamel surface, a strong dentino–enamel junction, and a decussating enamel rod pattern

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Mutations of the matrix metalloproteinase 20 (*MMP20*, enamelysin) gene cause autosomal-recessive amelogenesis imperfecta, and *Mmp20* ablated mice also have malformed dental enamel. Here we showed that *Mmp20* null mouse secretory-stage ameloblasts maintain a columnar shape and are present as a single layer of cells. However, the maturation-stage ameloblasts from null mouse cover extraneous nodules of ectopic calcified material formed at the enamel surface. Remarkably, nodule formation occurs in null mouse enamel when MMP20 is normally no longer expressed. The malformed enamel in *Mmp20* null teeth was loosely attached to the dentin and the entire enamel layer tended to separate from the dentin, indicative of a faulty dentino–enamel junction (DEJ). The enamel rod pattern was also altered in *Mmp20* null mice. Each enamel rod is formed by a single ameloblast and is a mineralized record of the migration path of the ameloblast that formed it. The enamel rods in *Mmp20* null mice were grossly malformed or absent, indicating that the ameloblasts do not migrate properly when backing away from the DEJ. Thus, MMP20 is required for ameloblast cell movement necessary to form the decussating enamel rod patterns, for the prevention of ectopic mineral formation, and to maintain a functional DEJ.

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Enamel development is stage specific. The two predominant stages are the secretory stage and the maturation stage. During the secretory stage, matrix metalloproteinase-20 (MMP20) is expressed while a protein scaffold is formed and while the mineral precipitates as carbonated hydroxyapatite in long thin ribbons that grow out to form the full thickness of the enamel layer. The secretory stage is when the tall columnar ameloblasts of the enamel organ begin moving in rows to form the rod and inter-rod enamel. During the maturation stage, expression of MMP20 ends and the ameloblasts shorten and cease movement. This is when kallikrein-related peptidase 4 (KLK4) assists in removal of the protein scaffold, which allows the enamel ribbons to grow further in width and thickness into large hexagonal crystals (1, 2). The crystals eventually press against one another and interlock as the enamel matures into its final hardened form (3).

Matrix metalloproteinases are a family of proteinases that cleave virtually all extracellular matrix proteins. Matrix metalloproteinases play critical roles in reproduction, development, morphogenesis, wound healing, tissue repair, regeneration, remodeling, and cell migration [reviewed in (4)]. Matrix metalloproteinase-

20 is required for the development of healthy dental enamel. People and mice with homozygous *MMP20* (human)/*Mmp20* (mouse) mutations have soft discolored enamel that may be hypoplastic and abrades easily from the dentin surface (5–11). Although the expression of MMP20 during the secretory stage is necessary to cleave enamel matrix proteins (7, 12–22), MMP20 expression is also required to maintain a normal enamel organ morphology during the maturation stage of development when MMP20 is no longer expressed (6, 7). In the absence of MMP20, ectopic calcified nodules appear on the maturation-stage enamel surface, which disrupts the continuity of the almost linear ameloblast layer.

Here we demonstrate, using the *Mmp20* null mouse, that: (i) calcified nodules form during the maturation stage of enamel development, (ii) the maturation-stage ameloblasts completely cover these nodules, (iii) the enamel is only loosely connected to the underlying dentin, and (iv) the enamel rod pattern is severely malformed or non-existent. These results are consistent with MMP20 playing a key role in cell movement, formation of a strong dentino–enamel junction (DEJ), and prevention of ectopic mineral formation.

Material and methods

All animals used in this study were housed in facilities approved by the Association for Assessment and Accreditation of Laboratory Animal Care, and all operations were performed in accordance with protocols approved by the Forsyth Institute Animal Care and Use Committee.

Histology

Incisors were from adult C57BL/6 mice. Mice were anesthetized with chloral hydrate and fixed by intravascular perfusion with 1% or 2.5% glutaraldehyde buffered with 0.08 M sodium cacodylate containing 0.05% CaCl₂, pH 7.2. The hemimandibles were removed and most jaws were decalcified at 4°C in 4.13% disodium ethylenediaminetetraacetic acid (EDTA), post-fixed in osmium tetroxide reduced with potassium ferrocyanide, dehydrated in graded alcohols, and embedded in LR White acrylic resin (London Resin, Berkshire, UK), as previously described (23). Jaws fixed with 2.5% glutaraldehyde were either decalcified as described earlier in this paragraph, or left undecalcified. All of these samples were treated with reduced osmium and processed for embedding in Epon 812 (EMS, Hatfield, PA, USA) substitute. Semi-thin sections of incisor segments were cut with glass knives or with a diamond histoknife and stained with toluidine blue for examination by light microscopy.

Scanning electron microscopy

For scanning electron microscopy imaging of surfaces (Figs 1, 2, and 3B), the labial bone encasing the incisors was removed and the enamel organ cells were gently removed from the enamel and discarded. The exposed enamel surfaces were lightly brushed with dry KimWipes (Kimberly Clark, Neenan, WI, USA) and the incisors were examined at ×50 magnification without any further processing in a Hitachi S-3000N (Hitachi High-Tech, Rexdale, ON, Canada) variable pressure scanning electron microscope using the backscatter mode at 25 kV and 20 Pa pressure. In other experiments, whole erupted incisors (Fig. 3A) were air-dried, fastened to stubs, sputter coated with palladium gold, and examined using a JEOL 6400 scanning electron microscope. Scanning electron microscopy evaluations of incisor prism patterns (Fig. 4) were performed on rehydrated freeze-dried mandibular jaws from adult wild-type and *Mmp20* null mice. Samples were washed briefly in diluted sodium hypochlorite solution, rinsed in deionized water, lightly etched with 0.1% nitric acid, re-rinsed, air dried, and examined without coating in backscatter mode using a JEOL-JSM6460LV JEOL (Peabody, MA, USA) scanning electron microscope operated at 20 kV.

Results

Enamel from *Mmp20* null mouse appears histologically as two distinct layers and the enamel surface is marred by calcified nodules

Backscatter scanning electron microscopy imaging of mandibular incisors from adult *Mmp20* null mice revealed that the maturation-stage enamel surface was rough and contained numerous calcified nodules of

irregular sizes and shapes (Fig. 1A,B). These nodules were intermixed with cell debris on the enamel surface. At the end of the secretory stage (Fig. 1C; SEC), ameloblasts appeared to undergo ‘true’ postsecretory transition (Fig. 1C,D; PST) into shorter modulating cells typical of the maturation stage of amelogenesis. Strikingly, both modulating ameloblasts and papillary layer cells were arrayed around nodules which distorted the spatial arrangement of these cells and made them appear highly folded at times (Fig. 1E,G). Unlike normal mice where the enamel surface is flat and smooth, the enamel surface of *Mmp20* null mice was mostly irregular and undulating (Fig. 1D–G). Histological sections (Fig. 1D–G) showed that the abnormally thin enamel was composed of two layers (1 and 2) plus the mineralized nodules at the enamel surface. Interestingly, the inner enamel layer closest to the dentin appeared homogenous and did not vary greatly in thickness. However, the outer layer closest to the ameloblasts displayed large variations in thickness apparently as a result of the ameloblast layer accommodating the mineralized and mineralizing nodules (Fig. 1C–G). The nodules mineralized (Fig. 1F,G; min) during the maturation stage at a somewhat faster pace than the inner two layers of enamel, as observed by backscatter imaging (Fig. 1A,B) and the clear spaces remaining in the tissue following demineralization (Fig. 1G). This demonstrates that absence of MMP20 activity during the secretory stage causes ectopic mineralizations at the enamel surface during the maturation stage of enamel development.

Maturation-stage ameloblast layer of *Mmp20* null mice completely covers the calcified nodules

Rodent incisor maturation-stage ameloblasts normally have a low columnar morphology consisting of single, almost linear, groupings of cells. However, maturation-stage ameloblasts in *Mmp20* null incisors have a markedly altered arrangement and morphology where the null ameloblasts (Am) and associated papillary cells (PL) form a continuous covering over extraneous nodules of calcified material formed at the enamel surface (Fig. 2A–C). This causes distortions in the normal spatial arrangement of the cells so that the cells can appear to swirl or to be layered (Fig. 2A,B). Although it is possible that the nodules are embedded inside the cell layers of the enamel organ (Fig. 2B,D), in our experience the nodules located to the enamel surface (Fig. 2D inset) and the plane of sectioning made them appear within the cell layers. In undemineralized sections, nodules appear layered and to have abnormal structure (Fig. 2D). Nodule-free areas are also sometimes encountered in the maturation stage (Fig. 2E,F). The enamel in these locations often appears thin and the ameloblasts appear somewhat distorted with small pools of protein at their apices (Fig. 2E; arrow). Signs of inflammatory cell infiltration into the enamel organ are also sometimes seen in the regions closest to the gingival margin of the tooth (Fig. 2F; arrows; Fig 3B red arrows). This shows that the ameloblast and papillary layers from *Mmp20* null mice cover and promote the growth of the ectopic calcified nodules.

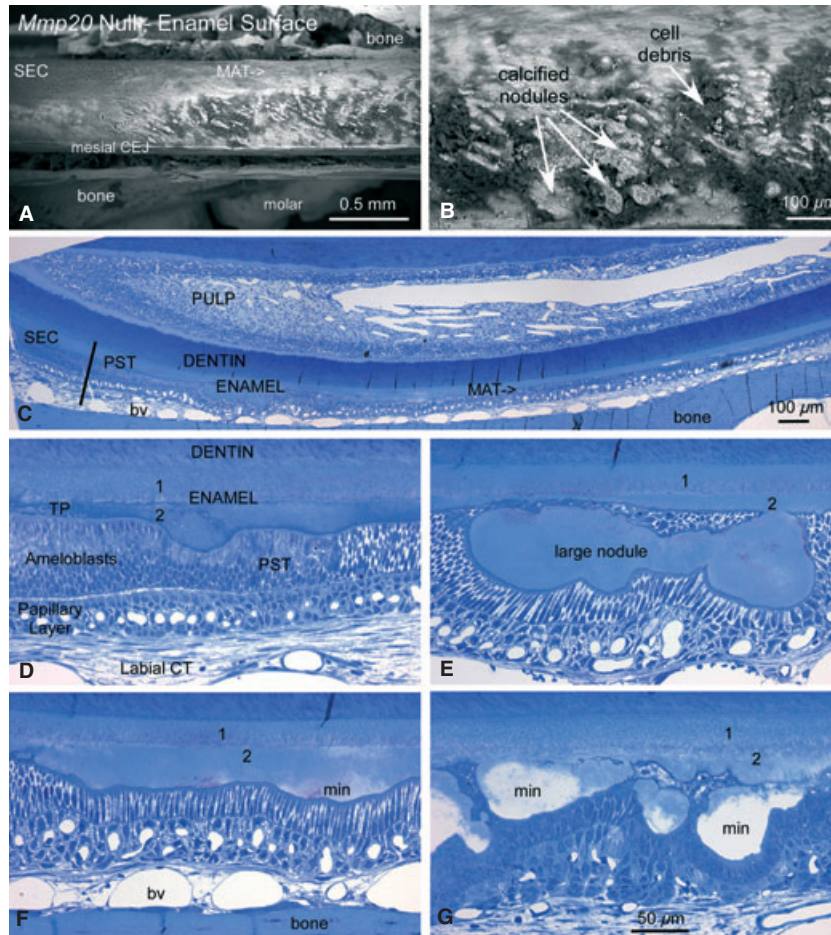


Fig. 1. Enamel from matrix metalloproteinase 20 (*Mmp20*) null mouse was present in two distinct layers and had a rough surface because of the presence of ectopic calcified nodules. Backscatter scanning electron microscopy images of the enamel surface from mandibular incisors of adult *Mmp20* null mice (A,B) revealed malformed enamel. The enamel was especially malformed during the maturation stage (A; MAT) where numerous calcified nodules of irregular sizes and shapes were intermixed with cell debris (B). Semi-thin plastic demineralized histological sections (C–G) show that at the end of the secretory stage (C; SEC) the ameloblasts appear to undergo typical postsecretory transition (C,D; PST) into shorter cells typical of the maturation stage of amelogenesis. However, the enamel was thin and was composed of two layers (1 and 2). Calcified nodules extend from the outermost layer nearest the ameloblasts to form a rough and uneven surface (layer 2). Paradoxically, the innermost layer near the dentin (layer 1) had a relatively smooth and even appearance (D–G). Both ameloblast and papillary layer cells were arrayed around the nodules to cover them, which distorted the spatial arrangement of the cells and, at times, made them appear folded (E,G). The nodules mineralized (F,G; min) during the maturation stage and at a somewhat faster pace than the inner two layers of enamel, as observed by backscatter imaging (A,B) and the clear spaces left in the tissue following demineralization (G). The magnification bar in panel G applies equally to panels D–G. bv, blood vessel; CEJ, cemento–enamel junction; labial CT, labial connective tissue; TP, Tome's processes.

Enamel from *Mmp20* null mouse is loosely connected to the underlying dentin

Enamel in *Mmp20* null mice has almost no resistance to mechanical stress and is so frail that during processing of incisors for scanning electron microscopy analyses, large pieces of enamel often become dislodged from the tooth and fall off (Fig. 3A). The enamel does not crumble but instead dislodges in large sheets from the incisor, suggesting that the interface between enamel and underlying dentin in null mice is particularly weak. Prior to eruption of incisors in *Mmp20* null mouse, a gap can be observed between the dentin and enamel that appears to accumulate debris (Fig. 3B black arrows). This shows that MMP20 is essential for formation of a mechanically strong DEJ typical of normal mice.

Enamel from *Mmp20* null mouse has a disrupted or non-existent rod pattern

Normal rodent incisor enamel has a decussating mineralized enamel rod pattern. Each rod is formed by one ameloblast and each rod preserves a complete record of the migratory path of the ameloblast that formed it. Figure 4A shows normal decussating enamel rods, rows of which can be observed to cross adjacent rows. In contrast, enamel from *Mmp20* null mouse has either a highly disorganized, almost unrecognizable, rod pattern (Fig. 4B,C) or it has virtually no rod pattern at all (Fig. 4D). This suggests that MMP20 is required for ameloblasts to move synchronously in rows that slide by each other to form the normal decussating enamel rod pattern.

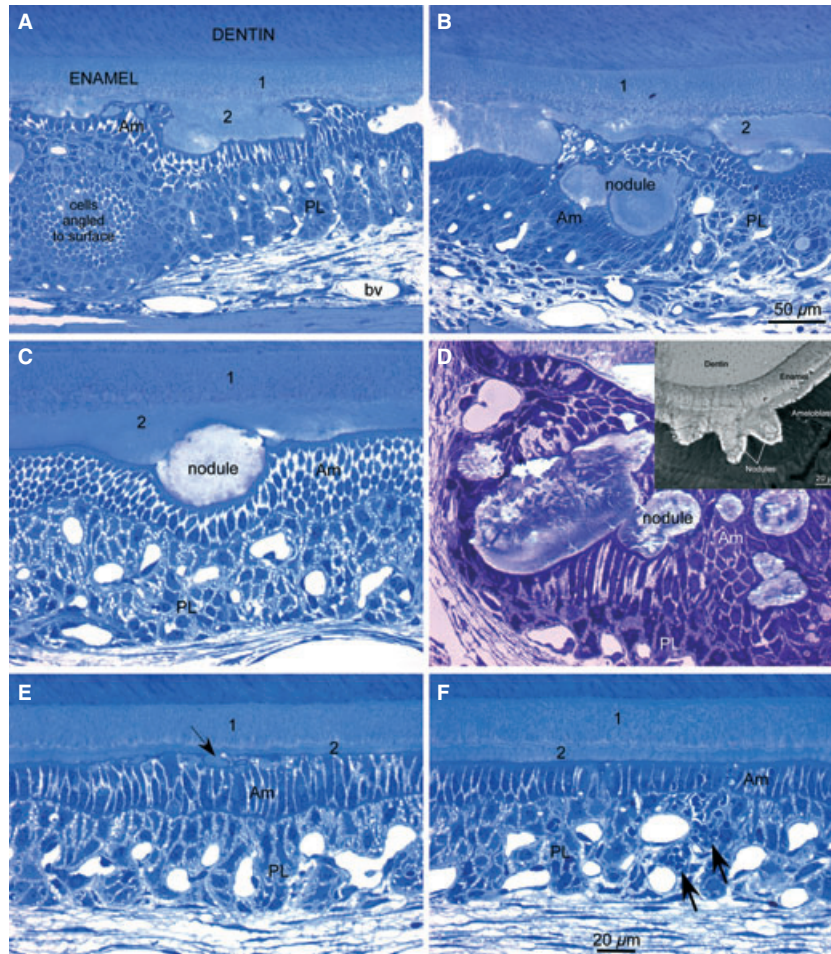


Fig. 2. Maturation-stage ameloblasts from matrix metalloproteinase 20 (*Mmp20*) null mice form a continuous covering over the nodules that project from the enamel surface. Semi-thin plastic demineralized histological sections (A–C, E, F) and a non-demineralized section (D) show maturation-stage enamel organ from mandibular incisors of adult *Mmp20* null mice. The inset in panel D shows backscatter imaging of the nodules at the enamel surface. Modulating ameloblasts (Am) and associated papillary layer cells (PL) formed a continuous covering over all nodules that projected from the enamel surface in *Mmp20* null mice (A–D). This caused distortions in the normal spatial arrangement of the cells, perhaps giving the illusion in some cases that isolated masses of cells were swirling and/or layered (A,B), or that nodules were embedded inside the cell layers of the enamel organ (B,D). In undemineralized sections (D), nodules appeared layered with a malformed structure. Nodule-free areas were sometimes encountered in the maturation stage (E,F). The enamel in these locations often appeared thin and ameloblasts appeared somewhat distorted with small pools of protein at their apices (E, arrow). Inflammatory cell infiltration into the enamel organ was sometimes observed in sections closest to the gingival margin of the tooth (F, arrows). 1 and 2 designate the two different enamel layers. The magnification bar in panel B applies also to panel A; the magnification bar in panel F applies equally to panels C–F.

Discussion

Here we show that enamel organ cells on the incisors of *Mmp20* null mice: (i) form thinner than normal enamel that is arranged histologically in two distinct enamel layers, (ii) form ectopic calcified nodules during the maturation stage that are covered by the ameloblast and associated papillary layers, (iii) form enamel with a very weak DEJ, and (iv) form enamel with a dysplastic or virtually non-existent rod pattern.

In addition to the well-defined roles for MMP20 in hydrolyzing newly secreted enamel matrix proteins and for maintaining normal enamel thickness (7, 12–22), MMP20 is also required to maintain a smooth outer enamel surface during the maturation stage of develop-

ment when MMP20 is no longer expressed (7). If MMP20 is not present during the early stages of development, the enamel surface becomes lumpy and covered with numerous variably sized calcified nodules during later enamel development. Perhaps the presence of MMP20 during early development is crucial for definitive ameloblast progression from the secretory stage to the maturation stage of enamel development. Deregulated developmental progression may promote the observed ectopic calcifications. This is supported by the observation that the *Mmp20* null mouse secretory-stage ameloblasts retract their Tomes' processes as if preparing to enter the maturation stage, but later re-extend their Tomes' processes as if resuming the secretory stage of development (24). Alternatively, it is

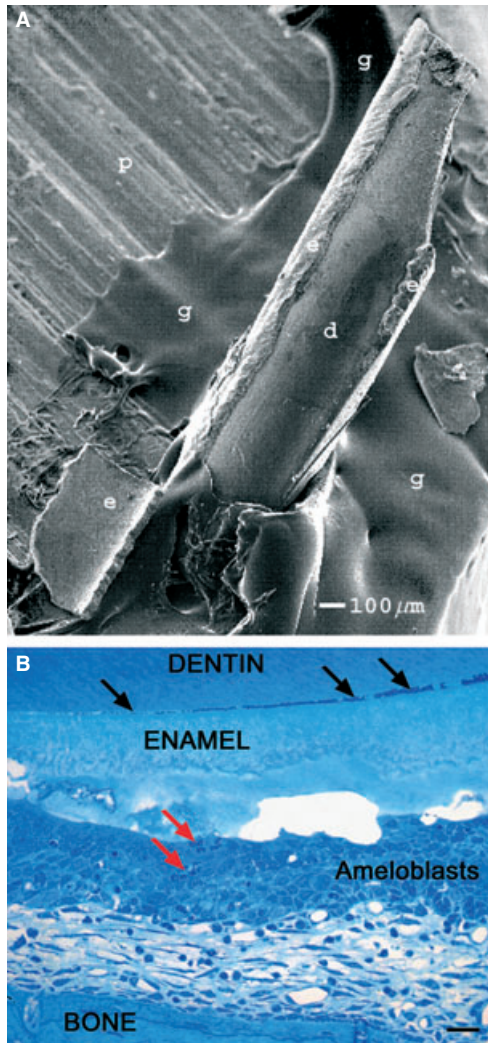


Fig. 3. The dentino–enamel junction (DEJ) is weak in enamel from matrix metalloproteinase 20 (*Mmp20*) null mice. Scanning electron microscopy image of an incisor from an *Mmp20* null mouse glued to a metal pedestal (A). Note that in the process of glueing the incisor to the pedestal a semi-conical piece of enamel fell off the dentin onto the pedestal at the base of the incisor. d, dentin; e, enamel; g, glue; p, metal pedestal. (B) A semi-thin plastic demineralized histological section shows infiltration of inflammatory cells into the ameloblast layer (red arrows). Debris appears to be accumulating at the DEJ where the enamel may already be separating from the underlying dentin (black arrows). Scale bar = 20 μ m.

possible that despite expression of KLK4 in the *Mmp20* null mice, an abundance of unreabsorbed enamel matrix protein at the ameloblast–enamel interface promotes ectopic nodule calcification. In either case, MMP20 is unexpectedly important for enamel maturation after it has normally ceased to be expressed.

The weak DEJ that allows the enamel to fall off in sheets was also interesting because it demonstrates that MMP20 activity is essential for establishing a strong bond between the dentin and forming enamel. Odontoblasts express MMP20 (13, 25) and this suggests that MMP20 may cleave important dentin protein(s), the cleavage products of which are essential for creating a

strong DEJ. Matrix metalloproteinase 20 does not cleave the most abundant dentin protein type I collagen (26), but it does cleave dentine sialophosphoprotein, which is the major non-collagen secretory product of odontoblasts responsible for dentin formation (27). Matrix metalloproteinase 20 also cleaves aggrecan and cartilage oligomeric matrix protein (28), type V collagen (29), type XVIII collagen (30), fibronectin, type IV collagen, tenascin-C, laminin-1, and laminin-5, but not type II collagen (26). Therefore, it is possible that cleavage of one or more of these proteins is essential for formation of the characteristically strong DEJ. Alternatively, because KLK4 is expressed after the enamel has reached its full thickness it may have limited access to the protein present in the deepest enamel layers. Thus, uncleaved protein at the DEJ in *Mmp20* null mice may interfere with the formation of a proper dentin–enamel interface. This could occur through failure of the dentin crystals to properly seed the enamel crystals or could occur from the failure to produce a functional basement membrane at the apical end of the ameloblasts.

The maturation-stage ameloblasts of *Mmp20* null mice maintain continuity by covering nodules projecting from the enamel surface. Perhaps the normal complement of ameloblasts over the abnormally thin enamel causes cell buckling because too many ameloblasts are present in a given area. In this case, cell buckling would precede nodule formation. Conversely, the ameloblasts may simply cover the forming nodules and this may contribute to the observed distorted intercellular spatial relationships. In this case, nodule formation would be concurrent with formation of the irregular ameloblast layer (Fig. 1G and 2A,B). In any case, the distortion becomes progressively worse as enamel matures and may be a contributing factor for infiltration of inflammatory cells as ameloblasts approach the gingival margin where the tooth erupts (Figs. 2F and 3B). Ameloblast cell–cell contacts may also be affected in *Mmp20* null mice. Cadherins are transmembrane proteins with extracellular domains that provide important adhesive contacts between neighboring cells. Previously it was demonstrated that ameloblasts express E-cadherin (31–37) and that MMP20 cleaves the E-cadherin extracellular domain (24). Therefore, in *Mmp20* null mice, the distorted spatial relationships may also result, in part, from cell attachments that would normally not exist because of MMP20 activity. These normally cleaved attachments probably also interfere with the ability of rows of ameloblasts to slide by one another to form the characteristic decussating enamel rod pattern.

We have previously demonstrated that cadherins are essential for dental enamel development. We showed that conditional deletion of the cadherin stabilizing molecule, p120-catenin (p120), from epithelial tissues had a striking effect on mouse enamel development (31). Binding of p120 to the cadherin intracellular domain prevents cadherins from becoming internalized and degraded (38–40). When p120 was deleted from the enamel epithelium, E-cadherin was no longer immunolocalized to the ameloblasts, and the ameloblasts failed to attach properly to neighboring ameloblasts, the

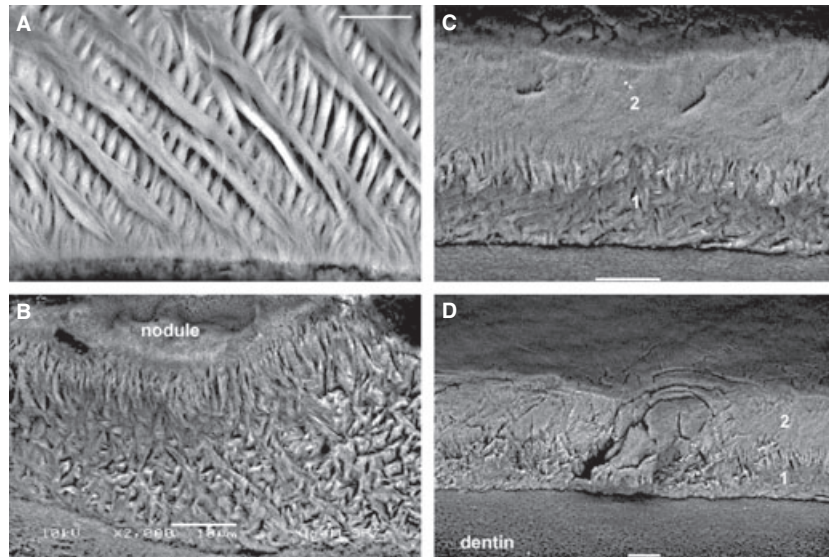


Fig. 4. Enamel rod patterns of mandibular incisors from wild-type and matrix metalloproteinase 20 (*Mmp20*) null mice. The wild-type enamel had criss-crossing (decussating) rows of enamel rods (A). The enamel from *Mmp20* null mice may have a poorly organized rod pattern (B), no rod pattern with a poorly organized rod layer beneath (C), or virtually no rod pattern whatsoever (D). 1 and 2 designate the two different enamel layers. Note that relative to Figs 1 and 2, the enamel from these incisors are displayed upside down. All magnification bars are 10 μm in length.

stratum intermedium, or the enamel surface. Although the general shape of the teeth was normal, the resulting dental enamel had no rod pattern and the malformed enamel was easily abraded from the tooth surface (31). Therefore, stabilization of cadherins on the cell surface is essential for dental enamel formation. This also suggests that a balance exists between proteolytic processing of cadherins for normal ameloblast cell–cell contact vs. a complete disruption of ameloblast function by an almost complete loss of ameloblast adhesive contacts. Thus, too many contacts may inhibit cell movement and too few may cause the ameloblast layer to fall apart.

Absence of cadherin hydrolysis by MMP20 may also be partially responsible for the poor or non-existent rod pattern observed in the *Mmp20* null enamel. Each enamel rod is formed by a single ameloblast (41). Therefore, each enamel rod is a mineralized record of the migratory path of the ameloblast that formed it (42). The enamel rod pattern in *Mmp20* null mice is dysplastic or non-existent, demonstrating that the null ameloblasts fail to migrate properly and implicates a role for MMP20 in modulation of ameloblast cell–cell contacts.

This study demonstrates that MMP20 is an important mediator necessary for maintaining a smooth enamel surface, a strong DEJ, and for establishing the decussating enamel rod pattern.

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Conflicts of interest – The authors declare no conflicts of interest.

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