Enamel and dental anomalies in latent-transforming growth factor beta-binding protein 3 mutant mice

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Latent-transforming growth factor beta-binding protein 3 (LTBP-3) is important for craniofacial morphogenesis and hard tissue mineralization, as it is essential for activation of transforming growth factor-beta (TGF-β). To investigate the role of LTBP-3 in tooth formation we performed micro-computed tomography (micro-CT), histology, and scanning electron microscopy analyses of adult Ltbp3-/- mice. The Ltbp3-/- mutants presented with unique craniofacial malformations and reductions in enamel formation that began at the matrix formation stage. Organization of maturation-stage ameloblasts was severely disrupted. The lateral side of the incisor was affected most. Reduced enamel mineralization, modification of the enamel prism pattern, and enamel nodules were observed throughout the incisors, as revealed by scanning electron microscopy. Molar roots had internal irregular bulbous-like formations. The cementum thickness was reduced, and microscopic dentinal tubules showed minor nanostructural changes. Thus, LTBP-3 is required for ameloblast differentiation and for the formation of decussating enamel prisms, to prevent enamel nodule formation, and for proper root morphogenesis. Also, and consistent with the role of TGF-β signaling during mineralization, almost all craniofacial bone components were affected in Ltbp3-/- mice, especially those involving the upper jaw and snout. This mouse model demonstrates phenotypic overlap with Verloes Bourguignon syndrome, also caused by mutation of TLPB3, which is hallmarkmed by craniofacial anomalies and amelogenesis imperfecta phenotypes.

Enamel is the body’s strongest tissue because of its dense mineralized content, which forms in a unique stage-specific process. Enamel formation is a sensitive process susceptible to environmental effects, as well as to the consequences of gene mutations, sometimes encountered in rare diseases (1). Enamel-formation defects observed in rodent genetic models are excellent systems for using to study the basis of analogous human malformations (2). Enamel formation begins when an organic protein-enriched matrix is deposited by ameloblasts during the secretory stage. This protein matrix is then modified through the transition and maturation stages (3). During the secretory stage, polarized columnar-shaped ameloblasts, adjacent to forming enamel, secrete enamel proteins, such as amelogenin, ameloblastin, and enamelin. Tomes’ processes on the distal surface of secretory ameloblasts organize the direction of enamel deposition into rods (4). During the maturation and mineralization stages, enamel matrix proteins are degraded in a stepwise manner by matrix metalloproteinase-20 (MMP-20) and kallikrein-4 (KLK-4) to form unique enamel prisms (5, 6). Murine mutations of enamel processing and degradation proteins produce enamel defects resembling human amelogenesis imperfecta (AI) phenotypes (5–8).

Transforming growth factor beta (TGF-β) is part of a superfamily of growth factors that regulate a broad range of cell growth, differentiation, and extracellular morphogenetic events (9). Inhibited TGF-β signaling (by mutation of TGF-β ligands, TGF-β receptors, or
intracellular SMADs) leads to reduced enamel formation and detachment of ameloblast cells from the dentin surface, causing the secretion of bubble-like masses that form cystic structures (10, 11). For example, mice with a conditional knockout mutation for TGF-β receptor II display enamel attrition with thinner crystals (12). Both Smad3-/- and Smad7-/- mutations reduce enamel mineralization (13, 14). Hence, disrupted TGF-β signaling has stage-specific consequences for amelogenesis in a variety of rodent models.

Transforming growth factor-β family proteins are secreted in the form of high-molecular-mass latency complexes that contain other proteins, including latent-transforming growth factor beta-binding proteins (LTBPs) (15). To date, four members of the LTBPs family (LTBP-1, LTBP-2, LTBP-3, and LTBP-4) are known. Through their interactions with other extracellular proteins, LTBPs are important regulators of the bioavailability and action of TGF-β (16). Immunoprecipitation studies using mouse pre-osteoblast MC3T3-E1 cells revealed that the LTBP polypeptide forms a complex with the TGF-β1 precursor (16). Moreover, LTBP-3 allows latent TGF-β complexes to be targeted to connective tissue matrices and cells (17, 18). Ltbp3-/- mice show severe skull deformities (19, 20) and an osteopetrosis-like phenotype (21), while other LTBPs members have broad functions in many systems (16). Exactly how craniofacial hard-tissue defects occur under Ltbp3 deficiency is still unclear, especially those concerning tooth and enamel malformation.

Mutations in human LTBP3 were first observed in a consanguineous Pakistani family, in which all affected members presented with short stature, vertebral and skull bone alterations, and oligodontia (22). In another family, two sisters with homozygous-recessive truncating mutations in LTBP3 also had oligodontia, short stature, and mitral valve prolapse (23). Our published report identified recessive hypomorphic LTBP3 mutations (including deletion, nonsense, and aberrant splice mutations) in patients with dental anomalies and short stature (MIM: 601216) (24) or Verloes Bourguignon syndrome (25). Using the adult Ltbp3-/- mouse model to characterize all dental hard-tissue components, we previously described very thin or absent enamel in both incisors and molars (24). More complete understanding of Ltbp3-/- mouse dental phenotypes will allow us to address how mutations of this gene in humans produce tooth abnormalities (19, 20), and to clarify the role of LTBP-3 in modulating TGF-β bioavailability (19).

Here we explore dental and cranial morphological differences caused by Ltbp3 deficiency using a three-dimensional (3D) imaging system [X-ray micro-computed tomography (micro-CT), classical histology, and scanning electron microscopy. We observed (i) alterations in enamel formation and deposition of enamel nodules, (ii) maturation-stage ameloblast disruptions, (iii) small bulbous-like formations inside molar roots, and (iv) enamel prism pattern malformations and reduced cementum thickness, collectively providing a framework for investigating the genetic basis of TGF-β signaling defects.

Material and methods

Animals
Ltbp3-/- mice (C57BL/6;Sv129;SW mixed background) were produced and genotyped as previously described (19, 20). All procedures with animals were performed according to the standards approved by New York University School of Medicine Institutional Animal Care and Use Committee.

Micro-CT imaging
The heads of five adult male Ltbp3-/- mutant mice and of five corresponding age/sex-matched wild-type (WT) littermates were examined at 3.5 months of age. To investigate malformations at slightly later stages, two male 5.5-month-old Ltbp3-/- mutants and two WT matching controls were also examined. All samples were scanned using the Quantum FX micro-CT pre-clinical in vivo imaging system (Caliper Life Sciences, Hopkinton, MA, USA), which operates at an energy of 80 kV and current intensity of 160 μA. We found no obvious skull morphological changes with aging. Micro-CT data acquisitions (with resolution at 80, 40, 20, and 10 μm pixel sizes for skulls, mandibles, incisors, and molars, respectively) were reconstructed using Analyze11.0 (Biomedical Imaging Resource, Mayo Clinic, Rochester, MN, USA). Bony dento-cranio-facial anatomy was sorted in coronal, transverse, and sagittal planes at skull, mandible, incisor, and molar levels. The segmented voxels were compiled to represent bony tissue and then were separated according to density to quantitate incisor and molar enamel. Cranial anatomical landmarks were analyzed using Euclidean distance matrix analysis (EDMA) for morphometric analysis (26, 27).

Histology
Samples (heads of 3.5-month-old WT and Ltbp3-/- mice) were fixed in 10% formalin for 30 d or longer and then transferred into 70% ethanol, washed in water, and demineralized in 10% ethylenediaminetaeacetic acid (EDTA) at 37°C for 10 d (the demineralizing solution was changed every day for the first 3 d and then every other day). After thoroughly washing in water, the heads were dehydrated in graded ethanol, cleared in Histo-Clear (National Diagnostics, Atlanta, GA, USA), and embedded in paraffin at 60°C. Ten-micrometer-thick transverse sections were collected, deparaffinized, and stained with hematoxylin and eosin (H&E). A detailed histology protocol can be found at http://www.empress.har.mrc.ac.uk.

Scanning electron microscopy
The upper and lower mure incisors of 3.5-month-old Ltbp3-/- and WT mice were dissected out of the alveolar bone. After rinsing with distilled water, the teeth were dehydrated in a graded series of ethanol, transferred in a solution of propylene oxide/epon resin (1:1, vol/vol), and embedded in Epon 812 (Euromedex, Souffleweyersheim, France). The teeth were sectioned into two halves along their sagittal axes using a water-cooled diamond circular saw (Bronwill Scientific, Rochester, NY, USA), and both surfaces were polished with diamond paste (Escil, Chassieu, France). One-half was etched with a 20% (wt/vol)
citric acid solution for 2 min, rinsed with distilled water, dehydrated in a graded series of ethanol solutions and left to dry at room temperature. The samples were coated with a gold-palladium alloy using a Hummer Jr sputtering device (Technics, Union City, CA, USA). Scanning electron microscopy assessments were performed using a Quanta 250 ESEM (FEI Company, Eindhoven, the Netherlands) operating with an accelerating voltage of electrons of 5 kV.

**Statistical analysis**

Non-parametric statistical testing (using the Mann-Whitney U-test) was used to compare the enamel volume of

\[ \text{Fig. 1. External morphology, micro-computed tomography (micro-CT), and histology analysis of lower incisors of a 3.5-month-old wild-type (WT) mouse (left column of panels) or a latent transforming growth factor beta binding protein mutant (Ltbp3-/-) mouse (right column of panels). The same WT or Ltbp3-/- mutant is shown in all respective panels. (A, B) In external views of lower incisors, the incisors of the mutant are shorter, laterally white, and centrally orange. (C, D) Three-dimensional rendered micro-CT images of the distal part of the mandible. The surface of enamel from the Ltbp3-/- mouse appears irregular (yellow arrows), and enamel nodules are seen near the alveolar bone crest area of the incisor (white arrows). (E, F) Two-dimensional micro-CT sagittal sections of the first lower molar and the underlying incisor (obtained from the respective boxed region in panels C and D), showing the enamel density. In the mutant the enamel layer is clearly reduced and has a rough appearance (red arrows). (G, H) Histological sections (hematoxylin and eosin staining) of the lower incisor (respective red boxed area in panels E and F) showing an irregular enamel layer (EN) in the Ltbp3-/- mutant. The odontoblast (OD) and dentin (DE) layers show no obvious changes. AM, ameloblasts; SR, stellate reticulum.} \]

\[ \text{Fig. 2. Histological analysis (hematoxylin and eosin staining) of the lower incisor ameloblasts. Normal ameloblast development starts at the cervical loop region of the incisor and involves differentiation in consecutive secretory, transition, maturation, and postmaturaton stages toward the incisal edge. These regions are illustrated both in central and lateral zones of the incisor in latent transforming growth factor beta binding protein (Ltbp3-/-) mutants (right-side panels) and respective wild-type (WT) controls (left-side panels). The central (A, E, I, M) and lateral (B, F, J, N) zones of WT mice show virtually identical morphology. (A-H) Ameloblasts of Ltbp3-/- mice show no obvious changes at the secretory and transition stages. However, the papillae underlying the ameloblasts have an irregular organization, with an increased space between each papilla. (I–L) At the maturation stage, the ameloblasts of Ltbp3-/- mice differentiate as shorter columnar cells (K). Ameloblasts appear highly abnormal in the lateral zone (L). (M–P) At the postmaturation stage, the ameloblast layer is disorganized in the mutants (O, P), with cell disruptions localizing mainly to the lateral zone, including the detachment of enamel (circled in P).} \]
incisor and molar enamel segmentations. The level of statistical significance was set at $P < 0.05$. For EDMA analysis, a parametric bootstrap approach was performed to validate the statistical significance, as described previously (27). All significance points were then recalculated as length in millimeters, aiding data interpretation.
Results

Alterations in enamel formation and deposition of enamel nodules

Macroscopically, the upper and lower incisors of normal teeth (from WT mice) are naturally yellow in color and the enamel is smooth and present only on the labial side (Fig. 1A). In Ltbp3−/− mutants, reduced coloration and irregular enamel surfaces were observed, with the lateral zones having less enamel than the central zones (Fig. 1B). The 3D rendered micro-CT images showed disrupted enamel patterns at the junction between the alveolar bone and the crown area, where numerous enamel nodules had formed laterally (Fig. 1C,D; white arrowheads). Early enamel formed at the distal part of the lower incisor showed obvious changes, as seen in two-dimensional (2D) sagittal plane reconstructions from micro-CT images (Fig. 1E,F; red arrowheads). This impaired enamel formation was confirmed by histological analysis (Fig. 1G,H). The enamel matrix in Ltbp3−/− mutants was irregular and failed to form a steady pattern (Fig. 1H). Changes in enamel volume were not uniform throughout the tooth crown, as enamel was completely missing from the lateral side of the labial portion of the incisor. However, a band of enamel of similar thickness as in WT mice was present in the central portion of the labial crown of the incisors.

Maturation-stage ameloblast disruptions

Histological analysis of secretory, transition, maturation, and postmaturation stage ameloblasts of the lower incisors revealed striking alterations in differentiation in Ltbp3−/− mutants (Fig. 2). At the secretory and transition stages, ameloblasts from Ltbp3−/− mice showed no obvious morphological changes compared with ameloblasts from WT mice, although the papillary layer underneath ameloblasts appeared impaired (Fig. 2A–H). Then, during the maturation stage, ameloblasts displayed a clearly disrupted morphology in mutants, especially within the lateral region of the incisor where the most severe ameloblast differentiation defects were observed (Fig. 2L). In addition, the ameloblast columnar epithelium was reduced in thickness within the central region (Fig. 2K). All these changes were more severe at postmaturation stages (Fig. 2M–P; compare WT with Ltbp3−/−).

Small bulbous-like formations inside molar roots

Micro-CT analysis allows us to segment each tissue according to its density, using different density thresholds. Using this method, we performed morphometric analysis of the skull and facial bones of Ltbp3−/− mice (Figs S1, S2). We were also able to segment the enamel to calculate the enamel volume (Fig. 3A,B; Fig. S3). The enamel volume of Ltbp3−/− mice was significantly reduced in all teeth (Fig. S3). In molars, even in the case of complete absence of enamel, the molar cusp pattern was maintained (Fig. 3C,D). As seen in Fig. 3C,D, the overall root pattern and molar surfaces, as revealed by 3D micro-CT, appear well preserved. However, irregular circles (bulbous-like structures) were seen, with an apparent random distribution on the mesial and/or distal molar roots, especially for M1 (Fig. 3H; yellow arrowhead). From histological analysis, small circle-like growths were observed within Ltbp3−/− roots, in contrast to the smooth mineralization pattern in roots from WT mice (Fig. 3I,J; red arrows). These findings indicate that LTBP-3 is not critical for proper root formation, but has important actions in ameloblasts in regulating proper differentiation of enamel.

Enamel prism pattern malformations and reduced cementum thickness – scanning electron microscopy analysis

Figure 4 shows enamel prism patterns on sagittal views of the lower incisor, which were obtained by high-resolution scanning electron microscopy. Mineralization starts by forming an enamel ribbon, with ameloblasts in WT incisors secreting a rod-like structure. This typical pattern was not observed in the Ltbp3−/− incisors (Fig. 4A,B). Figure 4C shows normal decussating enamel rods, with adjacent rows that criss-cross near the dentinoenamel junction. In contrast, enamel from Ltbp3−/− mice was poorly organized, with a clearly impaired prism pattern (Fig. 4D) and almost no rod pattern at the incisal edge (Fig. 4F). At its thickest area, the enamel from Ltbp3−/− teeth appears to be more porous (Fig. 4G,H). The whole outer enamel pattern, oriented in the opposite direction of the inner enamel, exhibited a disturbed structure (Fig. 4L; arrowhead). These observations suggest that ameloblasts from Ltbp3−/− mice fail to complete enamel formation.

Similar alterations in enamel formation were also found in molars, consistent with the observations made for incisors (Fig. 5). Root regions in Ltbp3−/− mutants showed a more condensed, unsteady pattern, according to the presence of bulbous-like formations (Fig. 5B; oval area shown by a broken line), and an irregular distribution of dentinal tubules beneath a reduction of cementum thickness (Fig. 5A,B, white arrowheads; Fig. 5C,D). These findings suggest that LTBP-3 is required for ameloblasts to secrete a proper enamel matrix, to establish a typical enamel rod structure, and for formation of normal dentinal tissue.

Discussion

Transforming growth factor-β proteins are thought to play a major role in the morphogenesis of developing teeth (14). Both TGF-β1 and TGF-β3 are produced by the enamel organ and are activated by components of the basement membrane, including those induced by odontoblast differentiation (28). Inactivation of the Tgfb1 gene (which encodes TGF-β receptor I) increases the proliferation of odontogenic epithelial cells (29). Both Tgfb1 conditional knockout mice and Tgfb1 over-expressing transgenic mice exhibit an abnormal tooth phenotype at early stages of enamel
formation (11, 12). As TGF-β proteins are present in the extracellular matrix as inactive forms, the absence of the targeting LTBP probably results in reduced TGF-β activity, mirroring other Tgfβ mouse mutant models (15). The incisors of Ltbp3-/- mice have a partial white color, with some yellow–orange bands centrally, indicative of loss of enamel, as confirmed by micro-CT, histology, and scanning electron microscopy analyses. In accordance with this result, TGF-β-activating SMAD2 and SMAD3, and TGF-inhibiting SMAD7, are found both in the enamel epithelium and in the dental mesenchyme, with alterations in these SMAD signaling components producing a variety of tooth phenotypes (14, 30–32).

Fig. 3. Alterations of enamel in molars and the molar root phenotype in latent transforming growth factor beta binding protein mutant (Ltbp3-/-) mice. (A, B) High-density renderings of micro-computed tomography (micro-CT) data from wild-type (WT) and Ltbp3-/- molars, highlighting enamel structural deficits. The enamel volume of the Ltbp3-/- molar is significantly reduced, especially at the third molar (also see Fig. S3). (C, D) Three-dimensional rendering of micro-CT images showing less density in molar crowns of Ltbp3-/- mice. The root pattern of Ltbp3-/- mice is similar to that of WT mice. (E–H) Two-dimensional sections confirming enamel reduction in both molars and incisors of Ltbp3-/- mutants. Section planes (F, H) are generated at the level of the yellow line. (H) A bulbous-like structure can be found in some areas of Ltbp3-/- molar roots (yellow arrowhead). (I, J) Histology (hematoxylin and eosin staining; regions highlighted by red boxes in E and G) showing a correctly organized root dentin in the lower molar of WT mouse, whereas irregular dentin staining is seen in the Ltbp3-/- mutant mouse. A bulbous-like structure (red arrowhead) is also observed. The periodontium, cementum, alveolar bone, and dental pulp tissues have no obvious morphological changes at the histological scale. BONE, alveolar bone; PDL, periodontium; PULP, dental pulp tissue.
Both histological and micro-CT analyses confirmed alterations in enamel formation in the mutants, with obvious reductions in incisor and molar mineralized enamel volume. A gradient of reduced enamel mineralization seen from central to lateral zones of incisors, and from the first to the third molars, suggests site-specific or temporally defined roles of LTBP-3. In addition, some incisor enamel areas also showed enamel nodules, especially at the lateral zones near the alveolar bone crest, suggesting that enamel formation and ameloblast differentiation are not maintained throughout tooth development. Normal fully formed murine incisor enamel has a decussating mineralized rod pattern. Each rod is presumably formed from a single ameloblast, preserving a complete record of its migratory path during its formation (3). Our scanning electron microscopy analysis showed enamel defects throughout the lower and upper incisors, as the prism patterns always failed to form. Consistently, SMAD3-signaling mutants showed alterations in enamel mineralization (13). In addition, Klk4 is down-regulated in 7-d-old Tgfbr2 conditional mouse mutants (12). The phenotypic changes in our Ltbp3-/− mice suggest that active TGF-β signaling complexes may regulate enamel matrix protein levels and/or enamel proteases, such as KLK-4, thereby regulating dental hard-tissue mineralization.

Severe ameloblast morphological alterations are observed at the maturation stage, revealed as both shortening and severe morphological alterations of these cells. Transforming growth factor-β family members are known to regulate blood-vessel formation and hematopoiesis (33), and in our histological analysis we observed some vessel enlargements and an impaired papillary layer below the ameloblasts. This suggests that the ameloblast alterations may potentially be caused by a vascularization defect, although this hypothesis will need further analysis to be substantiated. It has been reported that transgenic over-

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**Fig. 4.** Comparative scanning electron microscopy images of midsagittal sections of lower incisor enamel at various distances from the incisor extremity (the approximate location of the areas shown is given in the upper, low-magnification views), comparing teeth from wild-type (WT) (A, C, E, G, I, K) and latent transforming growth factor beta binding protein mutant (Ltbp3-/−) (B, D, F, H, J, L) mice. (A, B) Early maturation stage of enamel crystal deposition showing enamel ribbon patterns in WT mouse, whereas no clear pattern is seen in Ltbp3-/− mouse. (C, D) Junction area between the alveolar bone crest and crown analog of incisor, showing a criss-cross pattern in WT mouse, whereas a poor organization of enamel rods is observed in Ltbp3-/− mouse. (E, F) Incisal (or tip) area of incisor showing a typical enamel prism pattern in WT mouse, whereas no such pattern is observed in mutant mouse. (G, H) Fully mineralized enamel showing a rough external surface in Ltbp3-/− mouse. The enamel thickness in this zone is similar in WT and Ltbp3-/− mice. Boxed areas indicate areas shown under higher magnification in the next panels. (I, J) The enamel layer appears porous and non-decussated in Ltbp3-/− mutant mouse compared with WT mouse. (K, L) Under higher magnification, the outer enamel shows poorly organized, non-structured prism patterns (arrowhead), while an increased roughness at the outer enamel surface is also observed in the Ltbp3-/− mutant. DE, dentin; EN, enamel.

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expression of TGF-β1 in early secretory stage ameloblasts triggers dentin adhesion and the formation of cyst-like structures and mineralized globules (11). The loss of enamel ribbon formation observed in Ltbp3-/− mice confirms that enamel defects must start at the beginning of enamel deposition, suggesting that LTBP-3 signaling regulates enamel mineralization at early stages, with the effects continuing through amelogenesis.

Dental-targeted over-expression of TGF-β1 (using a Dspp–Tgfβ1 transgene) results in a phenotype similar to dentin dysplasia (34), and over-expression of TGF-β2 in the dentin matrix results in the reduction of dentin hardness and elasticity in male mice (35). Our histological analysis shows no obvious changes in dentin or odontoblasts in Ltbp3-/− mutants. Osteodentin, which was found in teeth from Dspp–Tgfβ1 transgenic mice (11), was not found in Ltbp3-/− mice. Even so, scanning electron microscopy analysis reveals that Ltbp3-/− mice have an impairment of dentinal tubular patterns, suggesting an indirect effect through alterations of enamel differentiation.

Some regions of the Ltbp3-/− molar roots have irregular bulbous-like formations under the cementum. Despite this, the overall root pattern is consistently similar to that of WT mice. In vivo assays of TGF-β signaling effects in ameloblasts using Ki67, a cellular marker for proliferation, showed marked proliferation in the Hertwig’s epithelial root sheath (HERS) at postnatal day 7 (12). Likewise, the presence of LTBP-3 in outer dental epithelium could regulate molar proliferation (24), explaining the reduction of cementum thickness. Ltbp3-/− mice also had striking craniofacial defects, such as reductions of viscerocranium and dome-shape neurocranium, which suggests premature ossification of some cranial sutures (19). Our results are consistent with previous work on Ltbp3-/− mice, which described craniofacial malformations (20) and a high bone-mass phenotype (21). Our measurements show that the reduction of the upper jaw is greater than that of the mandible (see molar relationship classification in Fig. S4).

Our previous study reported that four unrelated families had mutations in LTBP3, which probably produced AI and brachyolmia phenotypes. This highlights the role of LTBP-3 and TGF-β signaling in amelogenesis (24). Here, our findings of morphological changes within all tooth components, including enamel and ameloblasts, of Ltbp3-knockout mice supports multiple roles of LTBP-3 in odontogenesis. Understanding how LTBP-3 regulates hard-tissue mineralization and tooth development might provide treatment strategies for patients with a variety of craniofacial anomalies.

The Ltbp3-/− mouse phenotype described herein is characterized by severe enamel defects and overlaps with those of human LTBP3 mutations and mouse mutants for other genes of the TGF-β pathway. This study demonstrates that LTBP-3 is necessary for initiating and maintaining an enamel matrix, for regulating ameloblast differentiation, for complete enamel prism formation, and for establishing root mineralization.

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Conflicts of interest – The authors have no conflict of interest to declare.

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**Supporting Information**

Additional Supporting Information may be found in the online version of this article:

**Fig. S1.** Anatomical landmarks used for morphometric analysis.

**Fig. S2.** Form difference matrix analysis of skulls and mandibles.

**Fig. S3.** Bar graph of mean enamel volume of WT vs. *Ltbp3*-/− mutants.

**Fig. S4.** Molar relationship classification: the wild-type mice have molar class I occlusions (a normal occlusion), but *Ltbp3*-/− mutants have a class III molar bite (otherwise known as an anterior bite).