Glycobiology and Extracellular Matrices: Proteolytic processing of dentin sialophosphoprotein (DSPP) is essential to dentinogenesis

Qinglin Zhu, Monica Prasad Gibson, Qilin Liu, Ying Liu, Yongbo Lu, Xiaofang Wang, Jian Q. Feng and Chunlin Qin

*J. Biol. Chem.* published online July 13, 2012

Access the most updated version of this article at doi: 10.1074/jbc.M112.388587

Find articles, minireviews, Reflections and Classics on similar topics on the JBC Affinity Sites.

Alerts:
- When this article is cited
- When a correction for this article is posted

Click here to choose from all of JBC’s e-mail alerts

This article cites 0 references, 0 of which can be accessed free at

http://www.jbc.org/content/early/2012/07/13/jbc.M112.388587.full.html#ref-list-1
Proteolytic Processing of Dentin Sialophosphoprotein (DSPP) Is Essential to Dentinogenesis

Qinglin Zhu1,2*, Monica Prasad Gibson1*, Qilin Liu1, Ying Liu1, Yongbo Lu1, Xiaofang Wang1, Jian Q. Feng1, Chunlin Qin1

1Department of Biomedical Sciences, Texas A&M Health Science Center Baylor College of Dentistry, Dallas, Texas 75246, USA
2Department of Operative Dentistry and Endodontics, The Fourth Military Medical University, School of Stomatology, Xi’an, Shaanxi 710032, China

*These authors contributed equally to this work.

Running Title: DSPP Processing in Dentin Formation

To whom correspondence should be addressed: Chunlin Qin, D.D.S., Ph.D. Department of Biomedical Sciences, Baylor College of Dentistry, Texas A&M Health Science Center, 3302 Gaston Ave., Dallas, TX 75246, USA. Tel.: +1-214-828-8292, Fax: +1-214-874-4538, Email: cqin@bcd.tamhsc.edu

Keywords: DSPP, Proteolytic Processing, Posttranslational Modification, Biomineralization, Dentin

Background: In dentin and bone, dentin sialophosphoprotein (DSPP) is processed into the NH2-terminal and COOH-terminal fragments.

Results: The blocking of DSPP processing leads to hypomineralization defects in dentin, similar to those of Dspp-deficient mice.

Conclusion: The proteolytic processing of DSP is an activation step essential to dentinogenesis.

Significance: This study represents major progress in understanding how DSPP functions in dentinogenesis.

SUMMARY

DSPP, which plays a crucial role in dentin formation, is processed into the NH2-terminal and COOH-terminal fragments. We believe that the proteolytic processing of DSPP is an essential activation step for its biological function in biomineralization. We tested this hypothesis by analyzing transgenic mice expressing the mutant D452A-DSPP in the Dspp-knockout (Dspp-KO) background (referred to as “Dspp-KO/D452A-Tg” mice). We employed multipronged approaches to characterize the dentin of the Dspp-KO/D452A-Tg mice, in comparison with Dspp-KO mice and mice expressing the normal DSPP transgene in the Dspp-KO background (named “Dspp-KO/normal-Tg” mice). Our analyses showed that 90% of the D452A-DSPP in the dentin of Dspp-KO/D452A-Tg mice was not cleaved, indicating that D452A substitution effectively blocked the proteolytic processing of DSPP in vivo. While the expression of the normal DSPP fully rescued the dentin defects of the Dspp-KO mice, expressing the D452A-DSPP failed to do so. These results indicate that the proteolytic processing of DSPP is an activation step essential to its biological function in dentinogenesis.

Dentin Sialophosphoprotein (DSP) mRNA was first identified by cDNA cloning using a mouse odontoblast cDNA library in 1997 (1). However, dentin sialoprotein (DSP) and dentin phosphoprotein (DPP), the cleaved products of the DSP protein, were discovered much earlier and were believed to be separate entities until...
the single DSPP transcript was discovered (1, 2,3). Human genetic studies have shown that DSPP mutations are associated with dentinogenesis imperfecta (DGI), an autosomal dominant inherited disease characterized by dentin hypomineralization and significant tooth decay (4-11). Animal studies revealed that Dspp knockout (Dspp-KO) mice manifest hypomineralization defects in dentin. The widened predentin with irregular dentin mineralization in the Dspp-KO mice resembles the dentin defects of human DGI (12). These findings from human subjects and mouse models indicate that DSPP is critical for the formation and mineralization of dentin. However, the exact mechanistic steps by which DSPP functions in dentinogenesis remain largely unknown.

In dentin and bone, DSPP is proteolytically processed into the NH2-terminal and the COOH-terminal fragments (1,13,14). The NH2-terminal fragment of DSPP encoded by the 5’ portion of the DSPP transcript exists in two forms: the core protein form known as “dentin sialoprotein” (DSP) and the proteoglycan form referred to as “DSP-PG” (15-19). The COOH-terminal fragment of DSPP encoded by the 3’ region of the DSPP transcript is found in only one form, referred to as “dentin phosphoprotein” (DPP).

DSP isolated from the extracellular matrix (ECM) of rat dentin migrates at ~95 kDa on 5-15% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) (20). DSP accounts for 5-8% of the non-collagenous proteins (NCPs) in the ECM of rat dentin (21), while DSP-PG appears to be more abundant than DSP (15,19). The two glycosaminoglycan (GAG) chains of DSP-PG isolated from rat dentin are made of chondroitin-4-sulfate, and the two GAG chains of mouse DSP-PG are attached to S242 and S244 in the mouse DSPP sequence (17). In porcine dentin, the DSP-PG GAG chains appear to be made of chondroitin-6-sulfate (19). DSP, which contains few or no phosphates, has no significant effect on the formation and growth of hydroxyapatite (HA) crystals according to in vitro mineralization analyses (22). However, information regarding the effects of DSP-PG on the formation and growth of HA crystals is lacking. In vivo studies involving the transfer of a transgene encoding the NH2-terminal fragment of DSPP into the Dspp-KO background indicate that this fragment might regulate the initiation of dentin mineralization but not the maturation of mineralized dentin (23).

DPP, which accounts for as much as 50% of the NCPs in the ECM of rat dentin (24), contains large amounts of aspartic acid (Asp) and serine (Ser) residues, with the majority of Ser being phosphorylated (25,26). The Asp and phosphorylated Ser (Pse) residues are mostly present in the repeating sequences of (Asp-Pse-Pse)n and (Asp-Pse)n (1,13,14,27-29). The high levels of Asp and Pse give rise to a highly phosphorylated (25) and very acidic protein with the isoelectric point estimated to be 1.1 for rat DPP (30). DPP has a relatively high affinity to calcium (31,32) and is believed to have a direct role in controlling the rate and/or site of dentin mineralization (3,33,34). Several in vitro mineralization studies have indicated that DPP is an important initiator and modulator in the formation and growth of HA crystals (35-37).

The remarkable chemical differences between the NH2-terminal fragment (including DSP and DSP-PG) and the COOH-terminal fragment (DPP) of DSPP suggest that these molecular variants may perform different functions in biomineralization although they are derived from the same mRNA. Studies have shown that significant amounts of DSP, DSP-PG and DPP are present in the ECM of dentin, whereas a very minor quantity of the full-length form of DSPP is found in the dentin (16,38). The abundance of DSPP fragments,
along with the scarcity of full-length DSPP in the dentin, suggests that the processed fragments of DSPP may be the functional forms directly involved in biomineralization.

Previous in vitro studies by our group and others have shown that bone morphogenetic protein 1 (BMP1)/Tolloid-like metalloproteinases cleave mouse DSPP at the NH₂-terminus of Asp⁴⁵², while substitutions of Asp⁴⁵² or two residues that are immediately NH₂-terminal to Asp⁴⁵², block the processing of this protein partially or completely (38, 39, 40). More recently, we generated transgenic mice expressing a mutant DSPP in which Asp⁴⁵² was replaced by Ala⁴⁵²; the transgene expressing this mutant DSPP (referred to as “D452A-DSPP”) was driven by the 3.6-kb rat Col 1a1 promoter, which allows the expression of this transgene in the bone and dentin (40). We observed that the majority of D452A-DSPP was not cleaved in the bone of the transgenic mice in the wild type background, indicating that the D452A substitution effectively blocked the proteolytic processing of DSPP in the mouse bone (40). In the present study, we systematically characterized the dentin of mice expressing D452A-DSPP in the Dspp-KO background (referred as “Dspp-KO/D452A-Tg” mice) in comparison with Dspp-KO mice and mice expressing the normal DSPP transgene in the Dspp-KO background (i.e., without the endogenous Dspp gene). The mice expressing the D452A-DSPP transgene in the Dspp-KO background are referred to as “Dspp-KO/D452A-Tg” mice while those expressing the normal DSPP transgene in the Dspp-KO background are called “Dspp-KO/normal-Tg” mice. The polymerase chain reaction (PCR) primers for detecting the DSPP transgene were: forward, 5′-CCAGTTAGTACCACTGGAAAGAGAC-3′; reverse, 5′-TCATGGTTGGTGCTATTCTTGATGC-3′ (the expected PCR products when using mouse genomic DNA as the template were 521 bp for the transgene and 676 bp for the endogenous Dspp gene). The primers used to identify the endogenous Dspp alleles were: forward, 5′-GTATCTTCTGAGCCTGCATC-3′; reverse, 5′-TGTGTGGTGGCTTGGCTC-3′ (expected PCR product from the endogenous Dspp, 489 bp). The primers specific to the Dspp null allele (containing LacZ gene) in the

EXPERIMENTAL PROCEDURES

Generation of Dspp-KO/D452A-Tg and Dspp-KO/normal-Tg mice — The generation of transgenic mice expressing the transgene encoding D452A-DSPP or the transgene encoding normal DSPP in the wild type (WT) background has been described in our previous report (40). In these transgenic mice, the D452A-DSPP or normal DSPP transgene is downstream to the 3.6-kb rat Col 1a1 promoter, which drives the expression of the transgenes in type I collagen-expressing tissues, including bone and dentin. The mouse lines showing the highest expression level of D452A-DSPP [i.e., Line 4 in Zhu et al. (40)] or of normal DSPP [Line 7 in Zhu et al. (40)] in the long bone were crossed with Dspp knockout (Dspp-KO) mice (strain name: B6; 129-Dspptm1Kul/Mmnc; MMRRC, UNC, Chapel Hill, NC). The first crossbreeding generated Dspp-Tg; Dspp+/- mice. Then, the Dspp-Tg; Dspp+/- mice were mated with Dspp-/- mice to generate mice expressing the D452A-DSPP or normal DSPP transgene in the Dspp-KO background (i.e., without the endogenous Dspp gene). The mice expressing the D452A-DSPP transgene in the Dspp-KO background are referred to as “Dspp-KO/D452A-Tg” mice while those expressing the normal DSPP transgene in the Dspp-KO background are called “Dspp-KO/normal-Tg” mice. The polymerase chain reaction (PCR) primers for detecting the DSPP transgene were: forward, 5′-CCAGTTAGTACCACTGGAAAGAGAC-3′; reverse, 5′-TCATGGTTGGTGCTATTCTTGATGC-3′ (the expected PCR products when using mouse genomic DNA as the template were 521 bp for the transgene and 676 bp for the endogenous Dspp gene). The primers used to identify the endogenous Dspp alleles were: forward, 5′-GTATCTTCTGAGCCTGCATC-3′; reverse, 5′-TGTGTGGTGGCTTGGCTC-3′ (expected PCR product from the endogenous Dspp, 489 bp). The primers specific to the Dspp null allele (containing LacZ gene) in the
Dspp-KO mice were: forward, 5′-GTATCTTCATGGCTGTTGCTTC-3′ from the Dspp sequence; reverse, 5′-CCTCTTCGCTATTACGCCAG-3′ from the LacZ sequence (expected size of PCR product, 389 bp). The animal protocols used in this study were approved by the Animal Welfare Committee of Texas A&M Health Science Center Baylor College of Dentistry (Dallas, TX). Multiple approaches were used to characterize the mandibles of the following four types of mice: 1) Dspp-KO/D452A-Tg mice, 2) Dspp-KO/normal-Tg mice, 3) Dspp-KO mice, and 4) WT mice (C57/BL6J mice).

Expression levels of the DSPP transgenes in teeth — Quantitative real-time PCR was performed to evaluate the relative levels of DSPP mRNA in the incisors of the 1-month-old Dspp-KO/D452A-Tg, Dspp-KO/normal-Tg and WT mice. For real-time PCR analyses, total RNA was extracted from the mouse incisors with an RNeasy mini kit (Qiagen; Germantown, MD). The RNA (1 ug/per sample) was reverse-transcripted into cDNA using the QuantiTect Transcription Kit (Qiagen; Germantown, MD). The DSPP primers used for real-time PCR were: forward, 5′-AACTCTGTGGCTGTGCCTCT-3′ (in exon 3) and reverse, 5′-TATTGACTCGGAGCCATTCC-3′ (in exon 4). The real-time PCR reactions were performed as we previously reported (41).

Extraction and isolation of noncollagenous proteins (NCPs) from mouse dentin and detection of DSPP-related proteins — The NCPs, including DSPP-related proteins in the dentin, were extracted from the incisors of the 3-month-old Dspp-KO/D452A-Tg, Dspp-KO/normal-Tg and WT mice. For real-time PCR analyses, total RNA was extracted from the mouse incisors with an RNeasy mini kit (Qiagen; Germantown, MD). The RNA (1 ug/per sample) was reverse-transcripted into cDNA using the QuantiTect Transcription Kit (Qiagen; Germantown, MD). The DSPP primers used for real-time PCR were: forward, 5′-AACTCTGTGGCTGTGCCTCT-3′ (in exon 3) and reverse, 5′-TATTGACTCGGAGCCATTCC-3′ (in exon 4). The real-time PCR reactions were performed as we previously reported (41).

Plain X-ray radiography and micro-computed tomography (µ-CT) — The mandibles from the 3-month-old and 6-month-old mice were dissected from the four groups of mice and analyzed with the Faxitron MX-20 Specimen Radiography System (Faxitron X-ray Corp., Buffalo Grove, IL). For the µ-CT analyses, the mandibles were scanned using µ-CT35 imaging system (Scanco Medical, Basserdorf, Switzerland), as we previously described (42). In the µ-CT program, a scan of the whole mandible in 7.0-µm slice increments was selected for three-dimensional reconstructions to assess the shape and structure of the mouse mandibles.

Histology and immunohistochemistry — Under anesthesia, the Dspp-KO/D452A-Tg, Dspp-KO/normal-Tg, Dspp-KO and WT mice at the ages of postnatal 3 and 6 months were perfused from the ascending aorta with 4% paraformaldehyde in 0.1 M phosphate buffer. The mandibles were dissected and further fixed...
in the same fixative for 24 hours, and then
decalcified in 8% ethylenediaminetetraacetic
acid (EDTA) containing 0.18 M sucrose (pH 7.4)
at 4°C for approximately two weeks. The tissues
were subsequently processed for paraffin
embedding, and serial sections of 5 µm were
prepared. The sections were either stained with
hematoxylin & eosin (H&E) or used for
immunohistochemistry (IHC) analyses. For the
IHC analyses, the anti-DSP-2C12.3 monoclonal
antibody (43) was used at a dilution of 1:800 to
detect DSPP and DSP. The anti-biglycan
antibody (a gift from Dr. Larry Fisher of the
Craniofacial and Skeletal Diseases Branch,
National Institutes of Health, Bethesda, MD,
USA) was used at a 1:1000 dilution to detect
biglycan. Mouse IgG of the same concentration
as that of the primary antibody was the negative
control. All the IHC experiments were carried
out using the M.O.M. kit and DAB kit (Vector
Laboratories; Burlingame, CA) according to the
manufacturer’s instructions.

**Backscattered and resin-casted scanning
electron microscopy (SEM)** — The dissected
mandibles were fixed in 2% paraformaldehyde
and 2.5% glutaraldehyde in 0.1 M cacodylate
buffer solution (pH 7.4) at room temperature.
Four hours later, the fixation buffer was
replaced with 0.1 M cacodylate solution. The
specimens were then dehydrated in ascending
concentrations of ethanol and embedded in
methyl-methacrylate (MMA) resin (Buehler,
Lake Bluff, IL). The surfaces of the dentin
tissues of interest were polished using a micro
cloth with Metadi Supreme Polycrystalline
diamond suspensions of decreasing sizes (0.1
µm, 0.25 µm and 0.05 µm; Buehler, Lake Bluff,
IL). The samples were then washed in the
ultrasonic wash and placed in the vacuum
system overnight. For backscattered SEM, the
surfaces of the teeth embedded in MMA were
polished and coated with carbon. For the resin-
casted SEM, the dentin surface was acid etched
with 37% phosphoric acid for 2–10 seconds and
washed with 5.25% sodium hypochlorite for 5
minutes. The samples were then coated with
gold and palladium as described previously (44).
A FEI/Philips XL30 Field emission
environmental SEM (Philips, Hillsboro, OR)
was used to perform the SEM analyses.

**Double fluorochrome labeling of the dentin**
— Double fluorescence labeling was performed,
as we described previously, to analyze the
mineral deposition rate of the dentin in the
mouse incisors (42, 45). Briefly, calcein (green)
label (Sigma Aldrich, St. Louis, MO, USA) was
injected into the abdominal cavities of the
5-week-old mice at 5 mg/kg. One week later, 20
mg/kg of Alizarin Red label (Sigma Aldrich, St.
Louis, MO, USA) was administered
intraperitoneally. The mice were sacrificed 48
hours after the injection of Alizarin Red label.
The mandibles were dissected and fixed in 70%
ethanol for 48 hours and dehydrated through
ascending concentrations of ethanol (70–100%)
and embedded in MMA. Sections (10 µm thick)
were cut and viewed under epifluorescent
illumination using a Nikon E800 microscope
interfaced with Osteomeasure
histomorphometry software (version 4.1,
Atlanta, GA). The distance between the two
fluorescence labels of the incisor dentin (cross
section of incisors under the mesial root of the
first molar) was determined, averaged and
divided by seven to calculate the mineral
deposition rate, expressed as µm/day.

**RESULTS**

The dentin and bone of the transgenic mice
expressing the normal DSPP transgene or the
D452A-DSPP transgene in the wild type
background were the same as those of normal
mice. In the following section, we describe in
detail the dentin phenotypes of the
Dspp-KO/D452A-Tg mice and
Dspp-KO/normal-Tg mice in comparison with the
Dspp-KO mice and WT mice. We noticed
that the Dspp-KO and Dspp-KO/D452A-Tg
mice also had alveolar bone defects. In this report, we did not include a description of the non-dentin defects in these Dspp-mutant mice.

Expression of the DSPP transgenes in the teeth — Real-time PCR analyses using the mouse incisor RNA as the template revealed that the expression level of the normal DSPP transgene in the Dspp-KO background was approximately 16 folds of the transcription level of the endogenous Dspp in the WT mice, while the expression level of D452A-DSPP transgene was about 13 folds of the endogenous Dspp in the WT mice (Fig. 1).

Extraction and separation of NCPs and detection of DSPP-related proteins — Stains-All staining and Western immunoblotting were used to visualize the DSPP-derived proteins in the dentin of the Dspp-KO/D452A-Tg, Dspp-KO/normal-Tg and WT mice. All the chromatographic fractions from the dentin extracts that might contain DSPP-related products were analyzed by SDS-PAGE with Stains-All staining (Fig. 2). In the extracts from the WT and Dspp-KO/normal-Tg mouse incisors, DSP (Fig. 2, hollow arrows) was clearly visualized by Stains-All, along with weak protein bands matching the migration rate of full-length DSPP (Fig. 2, dotted arrows). While large amounts of the full-length DSPP were observed in the Dspp-KO/D452A-Tg mice, protein bands matching the DSP were hardly detectable in the incisors of these mice. In the Dspp-null mice, no DSPP-related signals were observed.

In the Western immunoblotting analyses, DSP and DSPP were clearly detected in the dentin extracts from the WT, Dspp-KO/D452A-Tg and Dspp-KO/normal-Tg mice (Fig. 3). The ratios of DSP (hollow arrow) to DSPP (dotted arrow) varied dramatically between the samples from the Dspp-KO/D452A-Tg mice and the Dspp-KO/normal-Tg mice. The ratio of DSP to DSPP in the Dspp-KO/D452A-Tg mice was 1:10, while that in the Dspp-KO/normal-Tg mice was 15:1, indicating that the full-length form of DSPP in the former mice was approximately 150 folds greater than in the latter. The findings from both the Stains-All and Western immunoblotting analyses indicate that D452A substitution effectively blocked the proteolytic processing of DSPP in the mouse teeth.

Anti-DSP immunostaining — Anti-DSP reactivity was observed in the odontoblasts and the dentin matrix of the WT, Dspp-KO/D452A-Tg and Dspp-KO/normal-Tg mice (Fig. 4). In the matrix, the anti-DSP signals were primarily detected around the dentinal tubules in both the Dspp-KO/normal-Tg mice (Fig. 4C) and Dspp-KO/D452A-Tg mice (Fig. 4D). The presence of anti-DSP activity in the dentin matrix of the Dspp-KO/D452A-Tg mice indicated that the uncleaved DSPP, like its processed fragments, was also secreted into the ECM (Fig. 4D).

Plain X-ray and µ-CT Analyses — Plain X-ray radiography (Figs. 5A-H) and µ-CT (Figs. 5I-P) analyses were performed to reveal the dentin structure in the 3- and 6-month-old mice. These analyses showed enlarged pulp chambers with very thin dentin in the Dspp-KO mice (Figs. 5B, 5F, 5J and 5N). The expression of normal DSPP transgene fully rescued the defects of enlarged pulp and thin dentin in the Dspp KO mice (Figs. 5C, 5G, 5K and 5O), whereas the D452A-DSPP transgene failed to reverse the dentin defects of the Dspp-KO mice (Figs. 5D, 5H, 5L and 5P).

Histological Analysis — At postnatal 3 and 6 months, the pulp chamber in the mandibular molars of the Dspp-KO mice was remarkably larger and the predentin zone was much wider than in the WT mice (Figs. 6A, 6B, 6E and 6F). While the structure of the dentin-pulp complex in the Dspp-KO/normal-Tg mice (Figs. 6C and 6G) was similar to that of the WT mice, the structure of the Dspp-KO/D452A-Tg mice (Figs. 6D, 6H, 6J and 6K) was distinct.
6D and 6H) resembled the one in the Dspp-KO mice. The histology findings confirmed X-ray data showing that the normal DSPP transgene rescued the Dspp-KO dentin defects while the mutant transgene did not.

Anti-biglycan immunostaining — Biglycan immunostaining (Figs. 6I-L) was performed to show the predentin zone since in the teeth, this proteoglycan is primarily localized in the predentin. The biglycan immunostaining analyses clearly showed that the predentin zone in the Dspp-KO (Fig. 6J) and Dspp-KO/D452A-Tg mice (Fig. 6L) was much wider and more irregular than that in the WT (Fig. 6I) or Dspp-KO/normal-Tg mice (Fig. 6K).

Backscattered SEM — Backscattered SEM analyses (Figs. 7A-H) indicated that the dentin in the Dspp-KO mice (Figs. 7B and 7F) and Dspp-KO/D452A-Tg mice (Figs. 7D and 7H) contained more areas that were unmineralized or hypomineralized and resembled interglobular dentin. In the backscattered SEM images, the white areas represent the regions with greater amounts of mineral (higher level of mineralization), while the black areas indicate less mineralization (i.e., unmineralized or hypomineralized). The dentin in the Dspp-KO (Figs. 7B and 7F) and Dspp-KO/D452A-Tg mice (Figs. 7D and 7H) contained more hypomineralized areas compared with the WT (Figs. 7A and 7E) and Dspp-KO/normal-Tg (Figs. 7C and 7G) mice.

Resin-casted SEM — The resin-casted SEM analyses (Figs. 7I-L) revealed that the dentin in the WT (Fig. 7I) and Dspp-KO/normal-Tg mice (Fig. 7K) had well organized and evenly distributed dentinal tubules of similar thickness, whereas in the Dspp-KO (Fig. 7J) and Dspp-KO/D452A-Tg (Fig. 7L) mice, the dentinal tubules were disorganized and collapsed in some areas.

Double fluorochrome labeling — In the WT (Fig. 8A) and Dspp-KO/normal-Tg mice (Fig. 8C), the two labeled zones were regular and evenly distributed. In the Dspp-KO (Fig. 8B) and Dspp-KO/D452A-Tg mice (Fig. 8D), the zones of fluorochrome labeling appeared irregular and diffused; in certain areas, the boundary between the two labels appeared blurry. In the double fluorochrome labeling analyses, the distance between the green (first) labeling and red (second) labeling represented the mineral deposition of the dentin matrix during the period between the two injections (7 days). The quantitative analyses of the distance between the two labels (Fig. 8E) indicated that the mineral deposition rates in the dentin of the Dspp-KO and Dspp-KO/D452A-Tg mice were much lower than in the WT or Dspp-KO/normal-Tg mice.

DISCUSSION

In the ECM of dentin and bone, DSPP is mainly present as the processed NH2-terminal and COOH-terminal fragments (including DSP, DSP-PG and DPP); only a minor amount of full-length DSPP could be detected in the dentin of wild type rat or mouse (38). Based on the abundance of DSPP fragments and scarcity of its full-length form in the dentin, along with the observed roles of DPP in the nucleation and modulation of apatite crystal formation, we hypothesized that the conversion of DSPP to its fragments by proteolytic processing may be an activation event, converting an inactive precursor to active forms, and this activation step may represent one of the controlling mechanisms in dentin formation (16,40,46). In this study, we generated Dspp-KO/D452A-Tg mice lacking the endogenous Dspp gene but expressing the transgenic D452A-DSPP protein, in which Asp452, a key cleavage-site residue, was replaced by Ala452. The dentin of the Dspp-KO/D452A-Tg mice was compared with that of the Dspp-KO mice, WT mice and Dspp-KO/normal-Tg mice that lacked the endogenous Dspp gene but expressed the transgenic expression of normal DSPP protein.
These analyses showed that the D452A substitution effectively blocked the proteolytic processing of this protein in dentin and led to the inactivation of this molecule in dentinogenesis. The findings in the present investigation lend strong support to our hypothesis that the proteolytic processing of DSPP is an activation event, essential to its biological function in biomineralization.

A small portion (10%) of D452A-DSPP was cleaved in the Dspp-KO/D452A-Tg mice. Previously, we showed that substitutions of Asp\textsuperscript{452} and other residues close to this residue could not totally block the cleavage of DSPP by BMP1 in vitro, suggesting the presence of a secondary (cryptic) cleavage site that is currently unidentified (40). Nevertheless, the dentin defects in the Dspp-KO/D452A-Tg mice were very similar to those in the Dspp-KO mice; this observation indicates that the cleavage of DSPP at this cryptic cleavage site has a very limited effect on DSPP activation.

Immunohistochemistry with the monoclonal anti-DSP antibody revealed positive signals for DSP in the dentin matrix of the Dspp-KO/D452A-Tg mice, indicating that the uncleaved full-length DSPP was also secreted into the ECM of dentin. The anti-DSP activity in the dentin of the Dspp-KO/D452A-Tg mice was weaker than in the Dspp-KO/normal-Tg mice or WT mice. The relatively weaker signal for the anti-DSP antibody in the dentin of the Dspp-KO/D452A-Tg mice may be attributed to the difference in the degree of exposure of the epitopes (antigenic determinants); i.e., the epitopes of the processed fragment (DSP) may be more easily exposed and readily recognized by the anti-DSP antibody than the same antigenic determinants wrapped up in the full-length form of DSPP.

In addition to dentin and bone, DSPP has also been found in certain soft tissues such as the salivary glands, cartilage, liver, kidney and brain (41,47). It appears that the DSPP-derived products in the non-mineralized tissues may have posttranslational modifications different from those in the dentin. For example, the majority of DSPP in the condylar cartilage was not cleaved (47), and DSP in the non-mineralized tissues may be devoid of any carbohydrate moieties (41). These variations in the posttranslational modifications of DSPP suggest that the biological role of DSPP in the non-mineralized tissue might differ from that in dentin and bone, in which the cleavage of the full-length protein into its fragment forms is essential to its biological function in the mineralization of these two tissues.

**ACKNOWLEDGEMENTS**

We are grateful to Jeanne Santa Cruz for her assistance with the editing of this article, and to Dr. Paul Dechow for his support with the micro-CT analyses. This work was supported by NIH Grant DE005092 (to CQ).

**REFERENCES**


**FOOTNOTES**

The abbreviations used are: DSPP, Dentin sialophosphoprotein; *Dspp*-KO, *Dspp*-knockout; *Dspp*-KO/D452A-Tg, transgenic mouse expressing the mutant D452A-DSPP in the *Dspp*-knockout background; *Dspp*-KO/normal-Tg, transgenic mouse expressing the normal DSPP in the *Dspp*-KO background; DSP, dentin sialoprotein; DPP, dentin phosphoprotein; DGI, dentinogenesis imperfecta; DSP-PG, proteoglycan form of DSPP; ECM, extracellular matrix; SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis; NCPS, non-collagenous proteins; GAG, glycosaminoglycan; BMP1, bone morphogenetic protein 1; PCR, polymerase chain reaction; β-ME, β-mercaptoethanol; EDTA, ethylenediaminetetraacetic acid; H&E, hematoxylin & eosin; IHC, immunohistochemistry; SEM, scanning electron microscopy; MMA, methyl-methacrylate.

**FIGURE LEGENDS**

*Figure 1. DSPP mRNA levels in the incisor of the *Dspp*-KO/D452A-Tg and *Dspp*-KO/normal-Tg mice.* RNA isolated from the incisor of 1-month-old mice was used for real-time PCR analyses. The
mRNA level in the WT mouse incisor was taken as one, while that of the Dspp-KO/normal-Tg or Dspp-KO/D452A-Tg mice was expressed as folds over the WT mice. The level of the transgenic DSPP mRNA in the Dspp-KO/normal-Tg mice was ~16 folds of that of the endogenous Dspp gene in the WT mice. The level of mRNA from the D452A-DSPP transgene in the Dspp-KO/D452A-Tg mice was ~13 folds of that of the endogenous Dspp gene in the WT mice. The forward primer sequence used for real-time PCR analysis was from exon 3 of the endogenous Dspp gene, while the reverse was from exon 4. The results were from five analyses (n = 5) for each group.

**Figure 2. Stains-All staining of acidic proteins (stained blue or purple) in the NCP extracts from mouse dentin.** The NCPs were extracted from the dentin of 3-month-old WT, Dspp-KO, Dspp-KO/normal-Tg and Dspp-KO/D452A-Tg mice. The extracted NCPs were separated into 118 fractions (0.5 ml/fraction) by a Q-Sepharose ion-exchange column; the digits on the top of each image represent the fraction numbers. Sixty µl of sample from each of the fractions that potentially contained DSPP-derived products was loaded onto 5-15% SDS-PAGE. The dotted arrows denote DSPP, while the hollow arrows indicate DSP; their identities as DSPP and DSP were confirmed by Western immunoblotting (see Figure 3). The major blue bands in fractions 47-65 in the WT and Dspp-KO/normal-Tg mice (solid arrows) was primarily made of DPP, although these bands also contained a small amount of bone sialoprotein (BSP), which was confirmed by anti-BSP Western immunoblotting (data not shown). It should be noted that no anti-DPP antibodies are available to detect DPP in Western immunoblotting analyses. Note the abundance of DSP and DPP in the samples from the WT and Dspp-KO/normal-Tg mouse incisors as well as the large amounts of full-length Dspp in the samples from the Dspp-KO/D452A-Tg mice. M, molecular weight standard; DSP, pure DSP isolated from rat dentin; DPP, pure DPP isolated from rat dentin.

**Figure 3. Western immunoblotting to detect DSPP and DSP in the NCP extracts from mouse dentin.** Western immunoblotting with polyclonal anti-DSP antibodies was used to detect DSPP and DSP in the dentin extracts from the four types of mice. The partially purified rat dentin extract (0.1 µg) containing rat DSPP and DSP was used as a positive control (Ctrl). The Western immunoblotting results from a representative fraction (fraction 29, 60 µl) are shown here. While DSP (hollow arrow) and DSPP (dotted arrow) were detected in the dentin extracts from the Dspp-KO/D452A-Tg, Dspp-KO/normal-Tg and WT mice, the ratios of DSP to DSPP among these three types of samples were remarkably different. Based on our integrated calculation from triplicate analyses (n = 3) using the image J program, we estimated that the ratio of DSP to DSPP in the Dspp-KO/D452A-Tg mice was 1:10, while that in the Dspp-KO/normal-Tg mice was 15:1. In other words, if the quantity of DSP was used for normalization, the amount of DSPP in the Dspp-KO/D452A-Tg mice would be approximately 150 times that of the full-length protein in the Dspp-KO/normal-Tg mice. The findings from both Stains-All and Western immunoblotting analyses showed that the D452A substitution effectively blocked the proteolytic processing of DSPP in the mouse teeth.

**Figure 4. Anti-DSP immunohistochemistry.** The specimens were from the first mandibular molars of four types of mice at postnatal 3 months. The samples from the Dspp-KO mice (B) were used as negative controls. Anti-DSP activity was observed in the dentin matrix of the WT (A), Dspp-KO/normal-Tg (C) and Dspp-KO/D452A-Tg (D) mice. The signal for the anti-DSP antibody in
the dentin of the Dspp-KO/D452A-Tg mice was weaker than in the WT or Dspp-KO/normal-Tg mice. Bar: 50 µm

Figure 5. Plain X-ray analyses (A-H) and the µ-CT analyses (I-P) of mandibles from 3- and 6-month-old mice. At postnatal 3 months, the WT mice had evenly distributed and well mineralized dentin (A). The mandibular molars in the Dspp-KO mouse (B) had an enlarged pulp chamber and thinner dentin compared with the WT mice. The tooth defects in the Dspp-KO/D452A-Tg mouse (D) were similar to those of the Dspp-KO mouse, whereas the teeth of the Dspp-KO/normal-Tg mouse (C) resembled those of the WT mice. At postnatal 6 months, the teeth in the Dspp-KO/normal-Tg mouse (G) also appeared the same as those in the WT mice (E), while the teeth of the Dspp-KO/D452A-Tg mouse (H) resembled the Dspp-KO mouse teeth (F). In the µ-CT analyses of mandibles, the teeth of the Dspp-KO (J, N) and Dspp-KO/D452A-Tg (L, P) mice had similar dental defects, which included enlarged pulp chambers and thinner dentin. The WT (I, M) and Dspp-KO/normal-Tg (K, O) mice had normal dental structures. Bar: 200 µm

Figure 6. H&E staining of the dentin-pulp complex in 3- and 6-month-old mice (A-H) and biglycan immunostaining of predentin/dentin in 3-month-old mice (I-L). At postnatal 3 months, the Dspp-KO mouse (B) had wider predentin (solid arrow), uncoalescent calcospherites (hollow arrow) and an irregular dentin-predentin border compared to the WT mouse (A). The Dspp-KO/D452A-Tg mouse (D) showed dentin abnormalities similar to those of the Dspp-KO mouse (B). The dentin of the Dspp-KO/normal-Tg (C) mouse resembled that of the WT mouse (A). At postnatal 6 months, the dentin-pulp structures in the Dspp-KO/normal-Tg (G) mouse resembled those of the WT mouse (E), while the Dspp-KO/D452A-Tg mouse teeth (H) resembled those of the Dspp-KO mouse (F). Bar: A-H = 50 µm. In the biglycan immunostaining of predentin/dentin, the predentin stained by the anti-biglycan antibody (brown color) in the WT mice (I) and Dspp-KO/normal-Tg mice (K) was thin, smooth and evenly distributed, while the predentin in the Dspp-KO mice (J) and Dspp-KO/D452A-Tg mice (L) was wider and unevenly distributed. Bar: I – L = 100 µm

Figure 7. Backscattered SEM analyses (A-H) and resin infiltration and acid-etched SEM analyses (I-L) of the mandibular first molar from 3-month-old mice. In the backscattered SEM images, the white areas represent the regions with greater amounts of mineral (higher level of mineralization), while the black areas indicate those with less mineral (lower level of mineralization). The dentin in the Dspp-KO (B, F) and Dspp-KO/D452A-Tg mice (D, H) had more black areas (dotted arrows) than in the WT (A, E) or Dspp-KO/normal-Tg (C, G) mice, indicating that the former two had more hypomineralized areas than in the latter two. The images in E-H are enlarged views of the boxed areas in figures A-D. Bar: A-D = 100 µm, E-H = 40 µm. In the resin infiltration and acid-etched SEM images of the mandibular first molar, the dentinal tubules in the WT mice (I) and Dspp-KO/normal-Tg mice (K) had uniform diameters and were evenly distributed, running parallel to each other and perpendicular to the dental enamel junction (DEJ, hollow arrows). In contrast, the dentinal tubules in the Dspp-KO (J) and Dspp-KO/D452A-Tg (L) mice were tangled, had uneven diameters and appeared collapsed. Bar: I – L = 50 µm

Figure 8. Double fluorochrome labeling. The specimens were from the dentin of 5-week-old WT (A), Dspp-KO (B), Dspp-KO/normal-Tg mouse (C) and Dspp-KO/D452A-Tg mouse (D). In these
analyses, the first injection (calcein) produced a green label, while the second injection (Alizarin Red) made a red label. The distance between the green zone and the red zone reflected the width of the dentin matrix that was mineralized in seven days. Compared to the normal dentin in the WT and Dspp-KO/normal-Tg mice, the labeling zones in the Dspp-KO mice and Dspp-KO/D452A-Tg mice were diffused, indicating an irregular deposition of mineral. Bars = 10 µm. E, quantitative analyses (N = 3) showed that the dentin of the Dspp-KO mice and Dspp-KO/D452A-Tg mice had a remarkably lower mineral deposition rate compared to the WT and Dspp-KO/normal-Tg mice.
Fig. 4
Fig. 5
Fig. 8