Nonuniformity in ligaments is a structural strategy for optimizing functionality

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Ligaments serve as compliant connectors between hard tissues. In that role, they function under various load regimes and directions. The 3D structure of ligaments is considered to form as a uniform entity that changes due to function. The periodontal ligament (PDL) connects the tooth to the bone and sustains different types of loads in various directions. Using the PDL as a model, employing a fabricated motorized setup in a microCT, we demonstrate that the fibrous network structure within the PDL is not uniform, even before the tooth becomes functional. Utilizing morphological automated segmentation methods, directionality analysis, as well as second harmonic generation imaging, we find high correlation between blood vessel distribution and fiber density. We also show a structural feature in a form of a dense collar around the neck of the tooth as well as a preferred direction of the fibrous network. Finally, we show that the PDL develops as a nonuniform structure, with an architecture designed to sustain specific types of load in designated areas. Based on these findings, we propose that ligaments in general should be regarded as nonuniform entities, structured already at developmental stages for optimal functioning under variable load regimes.

Periodontal ligament | nonuniformity | fiber directionality | microCT

Significance

Structure and function are intermingled and inseparable. Therefore, the structure–function dependency sequence is mostly unclear. Using the periodontal ligament as a model, employing a technique utilizing a loading system inside a microCT, we were able to visualize in 3D the fresh collagen networks and correlate their distribution and direction with loads exerted on the ligament. We show that the ligament structure is not uniform and is determined before it becomes functional, and therefore we propose that structural nonuniformity is specifically designed to optimize ligament function to the variable forces it sustains.

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phase enhancement which enables 3D visualization of the fibrous network of a fresh and intact PDL due to humidified conditions and mechanical stabilization (13, 14). Using a mouse molar tooth we were able to reach a high resolution with a voxel size of 0.79 μm and generate a detailed structural analysis of the collagen networks inside the PDL.

**A 3D Visualization of the PDL Collagen Networks.** The 3D images of fresh and unsectioned PDL reveal that the fibrous network is organized in longitudinal sheets along the long axis of the roots and not in bundles as described in the literature (Fig. 1). To analyze the different structural features of the collagen networks, we segmented out the tooth and the bone computationally based on morphological features rather than gray values (detailed information about the process can be found in SI Appendix). The segmented PDL images show the longitudinal backbone organization of fibers; however, oblique and horizontal components exist between the longitudinal scaffolds. Together, the longitudinal scaffold and horizontal components are distributed nonuniformly throughout the PDL. There are areas of high fiber concentrations—dense networks (Fig. 1, black arrowheads) while other areas have very few fibers—sparse networks (Fig. 1, gray arrowheads). The sparse networks are seen in the area between the roots—the furcation area (Fig. 1C)—and in the middle of the roots on the buccal side (Fig. 1E).

**Characterizing the Collagen Distribution.** Fiber density maps (Fig. 2) were generated to objectively evaluate the locations of dense and sparse networks (the algorithm is described in SI Appendix). The density maps show a distinct difference between fiber organization at the buccal and lingual sides of the root. In general the lingual side presents a uniform dense network (Fig. 2B), whereas the buccal side contains a dense network strip close to the crown of the tooth (black arrows); however, the middle area of both roots contains only sparse networks (Fig. 2A, white arrows). This difference can be readily seen in coronal views of the roots (Fig. 2C and D). A greater difference in the distribution at the lingual and buccal sides is seen in the mesial root (Fig. 2C). Only sparsely distributed fibrous sheets are seen in the furcation area, between the roots (Fig. 2E, white arrow).

**Correlating Function with Fiber Distribution.** The loading system in the microCT enables vertical loading of the tooth similar to a physiological mastication process while it is still inside the bone (Fig. 3A) (13). The nonfixed, fresh conditions of the specimen allowed analysis of the tooth trajectory under functional loads. The morphology of molar teeth of many mammalian species, including humans and mice, is such that the long axis of the crown is not continuous with the long axis of the roots. An angle of ~25° toward the buccal side exists between the two axes in the mouse molar (Fig. 3B), similar to the angle in a rat specimen (15). The loading system is designed such that the long axis of the crown will be continuous with the moveable anvil and perpendicular to the occlusal surface of the crown, similar to the direction of functional vertical loads. Here, the 3D scan in the compressed mode showed that the PDL space was getting narrow at the middle portions of the buccal surfaces of the roots as well as at the furcation (Fig. 3C and F), similar to what was shown in a rat model (15). The areas in the PDL that showed prominent narrowing and contact areas between the root and the bone were colocализed with areas in the PDL containing only few collagen sheets—the sparse networks (Fig. 3C–F, black arrows).
Structural Changes at Different Functional Stages. To define whether tooth function is correlated with the distribution of dense and sparse fiber networks, we scanned specimens at different functional stages. We scanned teeth before eruption (2 wk after birth); right after eruption, when the teeth begin to function (3 wk); fully functioning in young adult mice (10 wk); and in adult mice (15 wk). Fig. 4 displays the density maps at the different stages. Interestingly, the same fiber distribution patterns are seen at all ages. The transverse sections (Fig. 4, Right column) show two features: (i) the fiber arrangement, where the roots split, is generated as a sparse network before eruption, and remains as such during the different stages; and (ii) a dense fibrous strip (white arrows) surrounds the furcation area as well as the roots throughout all of the different functional stages, particularly in the preeruptive stage (2 wk). In the sagittal views (Left column), at the buccal side, sparse networks are seen in the middle of the roots (black arrowheads) and a dense network is encircling the upper third. The lingual side is composed of both dense and sparse networks before eruption at 2 wk and gradually transforms into a uniform dense network as function is established. These features can also be seen in the coronal sections (Middle column). The fiber density at the mesial and distal surfaces of the roots is reduced once the tooth is starting to function.

To verify the different fiber densities detected by the microCT method, a second harmonic generation method was used. Fibrilar collagens generate a very bright second harmonic generation (SHG) signal (16). This signal is reflected intensely from noncentrosymmetric structures such as those of triple helical fibrilar collagen molecules (17). We used the SHG signal from a PDL sample after the surrounding bone was reduced in its thickness to a transparency level that enabled the laser to penetrate through it. Fig. 5 is a composite showing the PDL-tooth–bone complex from the buccal side at the level where the roots emerge from the crown. According to the microCT scans this region contains the dense network strip. The SHG signal shows the same dense collagen network feature in this region (white arrows). The bone around the distal root was reduced such that exposure of the PDL along the root, where the sparse network is located, was revealed. Indeed, the SHG signal from the fibers in that area was weaker, corresponding to less organized and fewer collagen networks (black arrow). Verification of the collagen type was done by immunostaining of collagen type I and imaging by SHG with two photon emission; both signals were overlapping (SI Appendix, Fig. S8).

Directionality Analysis. Visualizing the entire PDL fibrous component in 3D revealed that the longitudinal backbone of the fibers displays a certain directionality in the mesial–distal direction. We therefore carried out a 3D directionality analysis comparing the phi and theta angles for fully functioning PDL and preeruptive state PDL. For the analysis, we used the cylinder correlation and trace correlation line modules in Avizo software (XFiber Extension version 9.4, Thermo Scientific; a detailed description can be found in SI Appendix). Briefly, the algorithm assigns a cylinder to each of the detected fibers in the 3D view according to a defined length, diameter, and curvature as specified by the user. Thereafter, analysis of the different cylinder parameters, such as angulation and length, are executed. Our analysis depicts the phi and theta angles as well as the correlation between theta angle values and fiber length. Fig. 6 A–D shows the results for the fully functioning PDL. The phi angles for the majority of the fibers in both the collar and root regions are below 180°; this implies a preferred distal direction of fibers. More specifically, the phi angles of the collar region peak at 30°, whereas the angles in the root region peak at values around 150° (ignoring the 180° peak that is neutral in the sagittal plane). The theta angle analysis shows that the collar region in the fully functioning PDL is mainly containing obliquely to horizontally oriented fibers (80° to the z axis), whereas a large portion of the fibers around the root but below the collar are vertically oriented at angles between 10° and 30° (Fig. 6 B and D). Correlating the fiber length with the theta angle shows that the vertical fibers are longer than the horizontal group. Looking at the PDL fiber direction analysis at preeruptive stage—before function is gained (Fig. 6 E–H), it can be seen that there is a preferred orientation of the collagen network already at that stage. With respect to the collar region, the theta angle distribution is very similar to that of the adult mouse—with preferred distal orientation; however, the phi angles peak at 130° (ignoring the 180° peak). This is explained by the fact that the tooth representing the 2-wk-old mouse is from the contralateral side of the 15-wk-old mouse. Along the root there was no directional preference (Fig. 6 G and H).

**Fig. 3.** Correlation between the fiber distribution and compression loads exerted on the PDL. (A) Closeup at the half mandible inside the loading setup with the vertically moving anvil bonded to the first molar. (B) A 2D slice from a microCT scan depicting the difference between the long axis of the crown and the long axis of the root. (C) Coronal 2D view of the mesial root imaged while the tooth was vertically compressed inside the loading setup showing a narrowing of the PDL region in the middle of the root at the buccal side (black arrow). Note the great proximity between the root and the bone. White arrow points at the widened area in the PDL. (D) Coronal 2D view in through the furcation when the tooth is vertically loaded, showing close proximity between the tooth and the bone at the furcation (black arrow). The density maps at an unloaded state (D and E) show that the regions of great proximity between the tooth and bone are occupied with sparse collagen networks (black arrows). B, buccal; L, lingual. (Scale bars: 100 μm.)

**Fig. 4.** Density map at different stages of development and function at 2 wk (before the tooth erupts), 3 wk (right after eruption), 10 wk (functioning tooth in young mouse), 15 wk (adult mouse). Density scale shows hot colors toward yellow (dense), cold colors toward blue (sparse). Black arrowheads point to sparse networks. (Scale bars: 200 μm.)
The longer fibers, however, were oriented vertically already in the prefunctional stage (Fig. 6H).

Nonuniformity of the Noncollagenous ECM in the PDL. The PDL is a highly vascularized tissue (18, 19). We demonstrate here that the sparse areas are correlated with high compression loads which in alveolar bone, unlike long bones, usually lead to bone resorption (20). However, under normal function and healthy conditions, the bone level in the furcation area is preserved despite the high and repetitive compression forces. Therefore, we propose that bone remodeling rates are higher in the sparse than in the dense areas of the PDL. Previous studies showed high correlation between multipotent cells and blood vessels (18, 19, 21). Moreover, blood vessels serve as a conduit for bone marrow multipotent cells to get to the PDL and might play a central role in bone turnover. Fig. 7A shows a coronal section through the mesial root of a wild-type mouse stained with anti-CD31, an endothelial marker (green). Many blood vessels were seen at sparse networks, for example in the middle of the root on the buccal side (B) (white arrows), whereas only a few were seen on the lingual side (L) (black arrows) occupied by a dense network. To image blood vessels in different fiber network areas, we also used endothelial-specific Flk1-Cre; tdTomato transgenic mice (22). A sagittal section of the first molar shows large blood vessels at the furcation area (Fig. 7B, white arrows), but such vessels are absent from other areas in the PDL. To image the 3D distribution of blood vessels, we used multiphoton microscopy to image the PDL of Flk1-cre; tdTomato mice. Few small blood vessels were noted in the dense collar region while large vessels were located in the sparse network close to the furcation area (SI Appendix, Fig. S9). Epithelial rests of Malassez cells are residual cells from Hertwig’s epithelial root sheath which is believed to have a central role in root formation (23). We noticed an extremely high presence of epithelial rests of Malassez cells in the furcation area (SI Appendix, Fig. S10). To further understand the contribution of different matrix components to the nonuniformity of the PDL, we examined the presence of chondroitin sulfate, decorin, and fibronectin in different networks. Using immunohistochemistry we were able to detect noticeable differences in chondroitin sulfate levels between the networks. Higher levels of chondroitin sulfate

Fig. 5. Image of the stacked maximal projection of second harmonic generation signal from a 15-wk-old mouse. White arrows point to dense network, where the fibers are clearly observed, and black arrows point to the area occupied by a sparse network where fewer fibers are observed. Black rectangle in Upper Left Inset depicts the region imaged with the SHG signal. (Scale bar: 100 μm.)

Fig. 6. Fiber directionality analysis displaying the phi and theta angles of a fully functioning tooth in a 15-wk-old mouse (A–D) and a nonfunctioning tooth, preeruption stage in a 2-wk-old mouse (E–H). The phi angle is an azimuthal angle in spherical coordinates and is measured in the xy plane from the x axis toward the y axis in a range of 0–360°. Theta angle is measured relative to the z axis in spherical coordinates. Due to the distinctly different characteristics of the collar region relative to the rest of the PDL fibers along the roots, we analyzed the direction of the collar region (A, B, E, and F) and the roots (C, D, G, and H) separately. When comparing the directionality of fibers between the roots, the distal root showed more uniformity in the fiber directionality; therefore, the results for the mesial root are displayed. To reduce processing time and increase the accuracy of the analysis, the root portion of the PDL was divided into three regions: upper, middle, and lower thirds. In general, areas closer to the collar region had a preferred direction compared with the apical thirds. Therefore, we show the results of the middle portion of the root as a representation for all three portions. x axis, red; y axis, green; z axis, blue. Phi angle analysis shows the counts in the y axis vs. measured angle in the x axis. The red line in the theta angle graph depicts the average fiber length in micrometers at each of the displayed angles.
were correlated with dense networks as well as blood vessels (Fig. 7C), and, interestingly, it was very specific to cementocyte lacunae (SI Appendix, Fig. S10). Decorin and fibronectin levels were not associated with collagen density (SI Appendix, Fig. S11).

Discussion

In this study, we show that the periodontal ligament is generated as a nonuniform entity. The distribution pattern of the different networks was shown to be preserved throughout different functional stages, and therefore the structure is preprogrammed to generate specific tooth function.

A very important area for tooth function is the furcation area between the roots of the tooth, acting as a fulcrum and exposed to compressive loads (15). Our results demonstrate that this area contains only a few collagen sheets, but numerous large blood vessels during different functional stages. A similar structure is found at injured sites in other ligament types. Ligaments are designed to work under tensional forces and contain mostly type I collagen (24); nonetheless, injuries to ligaments, typically caused by rupture, induce pathological changes that result in fewer type I collagen fibers with smaller diameters and increased vascularization (25, 26). Such regions are considered scars within ligaments; they function less efficiently in tension, but provide a better mechanism for enduring compressive forces (1, 24). The furcation area in the PDL sustains repeatedly high compression, torsion, and shear loads. Therefore, it can be regarded as a region of constant trauma, necessitating a structure that will reduce tissue damage. The presence of fewer type I collagen fibers and large blood vessels may leave more volume for noncollagenous ECM components. Such components, proteoglycans and GAGs, are found at sites exposed to compressive, torsional, and shear forces in other ligaments where they serve as load dissipaters and lubricators (27–29). However, when we examined the presence of noncollagenous ECM components, we found higher concentrations of chondroitin sulfate in the dense rather than the sparse networks, as well as with association to blood vessels. Noncollagenous ECM components are known to influence the stiffness of blood vessels either as structural components of the blood vessel wall (for example, elastin) or as an external component (28, 30–32). Therefore, due to the ligament preprogramming, blood vessels located in the compression areas might have a different structure and mechanical properties to provide higher stability and resistance to ischemia incidents due to the repetitive compressive forces.

Local ischemia in the PDL is believed to occur due to compression forces and triggers orthodontic tooth movement (20, 33). However, the alveolar bone in the furcation area, constantly under compression forces, is not resorbed under normal conditions (34, 35). Previous studies showed that the contact areas between the tooth and the bone due to compression forces are small and localized (15, 36). As demonstrated here, the size of blood vessels in the compression regions are large and ischemia is therefore unlikely to readily occur there. Moreover, blood vessels serve as a conduit for pluripotent cells with the capacity to differentiate into bone-forming cells. The high presence of the rests of Malassez cells also provides further support for the high-differentiation capacity in the furcation region. We therefore suggest that the furcation area is specifically designed to function under compression loads. This could explain the enigmatic orthodontic observation that teeth with multiple roots move slower than single-rooted teeth at the same force levels. Initially orthodontic loads would trigger recruitment of multipotent cells and other mechanisms to reinforce the resistance to compression loads rather than triggering bone resorption. Loads that exceed the dissipation capacity of the ligament may trigger different sensing units located outside of the PDL, such as osteocytes, and signaling pathways that eventually stimulate bone resorption and generate tooth movement (37).

Our results also show that the PDL fibrous network is divided into long vertical and short horizontal components. The longitudinal sheets can be regarded as the backbone of the fibrous component of the PDL. They are established before eruption and maintain their longitudinal orientation during the different functional stages. The horizontal element increases as function is gained, establishing the dense networks along the roots particularly on the lingual side. An exception to this pattern is the dense cervical collar region. This area, a unique structure surrounding the tooth just below the crown, contains a dense collagen network composed mostly of horizontal sheets. This structure exists before tooth eruption and is larger at the buccal side. Contrary to other dense regions, its dimensions are reduced after eruption. Since there is no pure horizontal movement of teeth, these dense horizontal fibers are likely to be under tension in all tooth movements and therefore possibly function as a main factor in guiding tooth movement. The reduction in its dimension with age may reflect changes that occur due to function and remodeling capacity. Our data also demonstrate that the fibrous PDL entity has a distal orientation even at preeruption stages and is prominently oriented to drive the tooth in a distal direction. This finding can possibly provide an explanation for the enigmatic physiological drift phenomenon of teeth. Teeth are showing constant drift throughout life in functioning teeth as well as when no functional forces exist. Human teeth have a mesial drift, whereas rodent teeth show a distal movement. Based on our findings, we suggest that the PDL has an internal force for tooth movement directed by the fibrous component, namely the horizontal group. This fact can open avenues for understanding tooth movement as well as bone formation and morphology.

Structural nonuniformities even at developmental stages are not unique to the PDL. The annulus fibrosus of the intervertebral disc, composed of type I collagen networks, was shown to sustain loads in a nonhomogenous pattern (38). Nonuniformities are detected already at the fetal stages; nevertheless, generally they are suggested to represent predisposition to future failure. Based on the data from this study, we suggest that nonuniformities are preprogrammed at developmental stages and are linked to normal ligament function. Support for this notion can be found in the literature, as nonuniformities were also seen in the 3D structure of the Achilles tendon (39) and at the fetal stage of the anterior cruciate ligament (40), suggesting that preprogramming for optimized function does occur.
Conclusions

Here, we have used the periodontal ligament to show that non-uniformities in ligament structure occur before function. The non-uniformity is specifically designed to optimize the tissue function for future load regimes and it is not generated in response to pathological events. Areas that function under compression loads are generated with sparse collagen fibers and large blood vessels. Moreover, the fibers are organized in preferred orientations, providing internal forces that are optimal to particular function. We thereby show that initially non-uniformities in ligaments are not structural modifications due to function, but are generated as a developmental structural strategy to optimize the tissue function.

Materials and Methods

Mice. All animal experiments were performed in compliance with NIH’s Guide for the Care and Use of Laboratory Animals and guidelines from the Harvard University Institutional Animal Care and Use Committee. The study was approved by the Harvard Medical School Institutional Animal Care and Use Committee (Protocol no. 97-074).

Wt-type C57Bl/6 mice were purchased from The Jackson Laboratory and served as controls.

Flk1-Cre mice (22) are a transgenic line generated by fusing the Cre gene to a fragment of the promoter sequence of Flk1. Reporter mice (tdTomato) were purchased from The Jackson Laboratory and crossed with the Flk1Creier-Cre to generate an endothelial cell-specific Cre strain inducing Tomato expression in endothelial cells only.

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MicroCT. The custom-made humidified loading setup is a modification of a previous design published by Naveh et al. (13); a detailed description can be found in SI Appendix.

Histology. Mandibles from 12-wk-old Flk1-Cre; tdTomato and C57Bl/6N mice were used; a detailed description can be found in SI Appendix.

Segmentation Analysis. Detailed description and algorithm are provided in SI Appendix.

Density Analysis. A detailed description can be found in SI Appendix.

Second Harmonic Generation. Hemimandibles were dissected and fixed in 4% formaldehyde for 24 h and washed with PBS; thereafter, the buccal bone was thinned using a dental drill under a Leica stereomicroscope until the tooth roots were visible. SHG signal was collected on a Zeiss 780 confocal microscope equipped with a Mai-Tai DeepSee femtosecond laser. Briefly, the laser was tuned to 860 nm, generating an SHG signal at 430 nm. The back reflected signal was collected through a 20x/1.0 NA water dipping objective onto a nondescanned photomultiplier tube detector behind a 420-480 bandpass filter. Images were analyzed with maximal projection mode in Fiji.

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