The effects of periosteum removal on the osteocytes in mouse calvaria

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Abstract

Objectives: To investigate the early influence of periosteum removal on osteocytes in mouse calvaria.

Material and methods: Fifteen C57BL/6 male mice were used in this study. Under anesthesia, a 2X3mm rectangular shape periosteum, 2mm behind the coronal suture and 1mm beside the sagittal suture, was surgically removed from the left side of calvaria whereas the right side was intact as control. Five mice were sacrificed at 1 day, 3 days and 7 days after the operation and the calvaria together with the surrounding tissues were histologically examined.

Results: At 3 days and 7 days, the number of empty lacunae significantly increased at the periosteum stripped side compared to the intact control side. The empty lacunae were mainly restricted to the outer half of the calvaria under the periosteum stripped area. Sclerostin distribution in lacunae and canaliculi also dramatically decreased at the periosteum stripped side at all the time points.

Conclusion: Periosteum removal would induce early apoptosis of the osteocytes in the outer layer of the bone and decrease sclerostin within this area.

Introduction

Periosteum, the stiff envelope bounds the skeleton, plays an essential role in offering protection and nutrition to the cortical bone [1]. Furthermore, periosteum also serves as a reservoir of progenitor cells and preserves a remarkable bone regeneration capacity, which is crucial for both prenatal bone development and postnatal fracture healing [2]. For immature skeletal system, periosteum-derived ossification makes long bones grow radially [3] and in fracture healing process, periosteum is also inevitable [4]. The formation of cartilage and bone within the callus owes largely to the mesenchymal progenitor cell from the periosteum [5]. When the periosteum is irritated by trauma, infection, neoplasia, or sometimes surgical procedure, new bone would be rapidly formed, which is favorable or unfavorable [6-9]. A previous study showed that the elevation of periosteum can induce new bone formation underneath [10]. In spite of the importance of periosteum, in some surgical circumstances, it has to be completely removed from the bone surface, such as in proper open reduction and in plate applications for treating bone fractures [11], and surgical removal of soft tissue sarcomas (STS) to ensure the adequate surgical margins [12]. In dental implant surgeries, clinicians often apply tension releasing incisions, cutting the periosteum at the base of the muco-periosteal flap to complete a primary closure, especially in ridge expansion procedures [13]. This incision into the periosteal layer leaves a hole, resulting in direct contact of the mucosa to the cortical bone, which can be considered as another kind of the periosteum stripping.

The apoptosis of osteocyte is another highlighted topic, which is already known to be a result of many physiological or pathological changes, such as rapid modeling process [14], age-associated oxidative stress [15], estrogen deficiency [16], overwhelming of glucocorticoid [17]. Also from a mechanical view, unloading of bone which weakens the canalicular fluid shear stress will result in osteocyte apoptosis [18]. Conversely, excessive loading which produces microdamage disrupting the canalicular system integrity also leads to osteocyte apoptosis [19].

Although the studies concerning periosteum stripping or osteocyte apoptosis have been often reported respectively, the reaction of osteocytes to the periosteum stripping has not been clarified yet. In this study, we histologically examined the effects of periosteum removal on the osteocytes in mouse calvaria.

Materials and methods

Animal models

Fifteen male C57BL/6 mice of 12 weeks old, weighing 25-30g, were used in this study. All the experimental procedures had been approved by the Animal Experiment Committee at Tokyo Medical and Dental University. (Approval number: 0140174A).

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Surgical procedure

General anesthesia was performed by an intraperitoneal injection of a combination of ketamine and xylazine (75 mg and 0.5 mg per kg body weight, respectively). The hair of calvarial zone was shaved and the skin over the skull was cleaned with 70% ethanol. Incision of 20mm length was made in the central line of head and partial thickness flap was raised. A 2 X 3mm rectangular (2mm behind the coronal suture and 1mm beside the sagittal suture) area was marked with ink on left side of calvarial periosteum (Figure 1A). Then, the stripping was performed with scalpel and dental excavator to remove the periosteum completely (Figure 1B and 1C). The right side of calvaria was intact as control. Finally the surgical area was closed with suturing.

Five mice were sacrificed for each time point at 1, 3 and 7 days after the operation. Following the anesthesia mentioned above, a perfusion fixation with 4% paraformaldehyde was performed to secure the bone tissue. Then, the whole calvarial bone together with the skin was harvested and fixed in the 4% paraformaldehyde.

Procedure for histological analysis

After fixation in 4% paraformaldehyde at 4°C for 2 days and washing in phosphate buffered saline (PBS) at 4°C for 1 days, samples were decalcified in 20% ethylenediaminetetraacetic acid (EDTA) at 4°C for 14 days and embedded in paraffin. Serial sections of 4 μm thickness were prepared and subjected to hematoxylin and eosin (HE) staining and immunohistochemical (IHC) staining. Images of stained specimens were acquired with an upright microscope (AxioSkop2, Carl Zeiss, Oberkochen, Germany).

Immunohistochemical (IHC) staining

Immunohistochemical analysis of markers for sclerostin was carried out in the paraffin embedded sections. Briefly, deparaffinized and hydrated sections were incubated with 0.1% trypsin solution (Difco™ Trypsin 250, BD Biosciences) for 20 minutes in water bath and hydrated sections were incubated with 0.1% trypsin solution carried out in the paraffin embedded sections. Briefly, deparaffinized Zeiss, Oberkochen, Germany).

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Histomorphometry analysis

For histomorphometric analysis of HE stain slides, high resolution digital images (at 10 times magnification) were acquired on the left side of calvaria within bone area 1mm away from the sagittal suture, and the same area of the control side. For the images of both sides, the number of empty lacunae and total lacunae was counted using the enlarged images. In the same area of HE stain slide, the images of sclerostin immunohistochemical stain slides were also obtained.

Statistical analysis

Statistical analyses were performed with standard Student’s t-tests for comparisons between different sides of within the same time point. The one-way analysis of variance (ANOVA) was employed for longitudinal comparisons between different time points of the same side.

Results

Histological findings

In all 3 time points, the periosteum removal sides showed less staining for sclerostin whereas the control sides showed more sclerostin distribution among osteocytes. Also in control side, a clear network of canaliculi radius, which was positively stained, spread out from the lacunae in the middle or inner layer of the bone could be often observed. However in the periosteum removed side, even in the 1day group, the canaliculi existing sclerostin were seldom observed around the lacunae (Figure 4).

Discussion

The empty lacunae are the most simple and direct indication of osteocyte apoptosis among several histological detection methods in vivo [20]. This phenomenon has been initially reported as the evidence of osteocytes loss in the aged patients about half century ago [21]. Although many factors can promote osteocyte apoptosis, in the present study, the periosteum stripping seems to be the reason that largely increases the number of empty lacunae at day 3 and day 7. This may be considered as the result of cutting off the main blood supply in the area. As reported previously, the stripping of periosteum significantly decreases the perfusion up to 20% of the whole length of tibia [11]. While for an irregular shape bone like mandible, as Saka B...
et al. reported, periosteal blood supply predominated in the body of the mandible [22]. Calvarium also has the similar structure of thick cortex with less marrow as the edentulous mandible. Another evidence to support this speculation is that the empty lacunae are mainly seen in the outer layer of the cavlaria, where can be taken as the periosteum nutritive domain as illustrated by the study of Pazzaglia et al. [23].

From the result of this study, the death signal may be released in the first day after the stripping, but the autophagy of osteocyte seems takes sometime which leads to the similar empty lacunae amount and rates on both sides at day 1. But soon the process ends itself and leaves the empty lacunae under the periosteum stripped side on the 3rd day. Interestingly, this death area did not enlarge to the day 7, indicating that it was determined at the first place when the periosteum was cut off.

This early death of osteocyte may lead to an osteoclastic resorption which could explain the previous finding that the long term bone density and mechanical properties dropped in the extensive circumferential periosteal stripping femur model [12]. Also it can enlighten the reason of high fracture risk after the STS surgical resection from the histological view [24]. For the surgery of oral implant, a tension releasing incision cut into the periosteal layer or placing a membrane and bone substitutes under the muco-periosteal flap may all cause the bone, especially the cortex of mandibular, losing the essential blood supply from periosteum, therefore it may undermine the basal bone where implant will be or already been placed in. Nevertheless, in a pilot study (date not shown), a simple elevation and reposition of periosteum, mimicking an open flap surgery in oral cavity, does not affect the osteocytes.

Sclerostin, the protein product of the SOST gene, acts as a negative regulator of bone formation and recently became a target for the anabolic treatment of osteoporosis [25]. This protein is mainly produced by osteocytes [26] and antagonizes the canonical Wnt signaling pathway to inhibit the osteoblast bone formation [27]. As mentioned previously, periosteum serves as not only a nutrition supply source but also as an important reservoir of osteoblasts. These cells have close interconnections with the osteocytes network via gap junctions [28] and extracellular paracrine signaling pathways [29]. Therefore, the osteoblasts in periosteum could be the most direct targets that osteocyte can aim at with the signals, such as, sclerostin.

Sclerostin is considered as a mechanotransduction key molecule which expression is enhanced by mechanical unloading [30], or suppressed by loading [31]. In the present study, the decreasing of sclerostin around the osteocytes still exist in the inner layer bone of the periosteum stripping side, cannot be explained by the loading theory, since calvaria unlike tibia or femur , is not under a daily body weight loading.

Figure 2. The number of empty lacunae increased significantly on periosteum stripping side than the control side at day 3 and 7 and almost all the empty lacunae were in the outer half of the calvaria. (Above: black arrows show the cutting edges of periosteum, bar as 300μm. Below: high magnificent of rectangular area in picture A, bar as 50μm).

Figure 3. A significant increase in empty lacunae number on periosteum stripping side, comparing to the control side at day 3 and 7. (*=p<0.05).

Figure 4. At all 3 time points, the periosteum stripping sides showed less staining for sclerostin than the control sides. A clear network of canaliculi radius spread out from the lacunae in the middle or inside layer of calvaria could be observed, stained with brown color, while seldom canaliculi could be seen on the periosteum stripping sides in all groups. (bar as 30 μm).
Bone fatigue can induce osteocyte apoptosis which is then followed by the osteoclastic resorption within the region around microcracks [32]. Loss of the periosteum could be a trauma similar to the microcrack inducing osteocyte apoptosis close under it. Therefore, this periosteum stripping would also raise the wave of resorption and remodeling, which requires suppression of sclerostin expression to turn on osteoblast bone formation. This suppression of sclerostin has also been seen in the bone after a dental implant placement [33]. Since lacking of sclerostin may stimulate osteoblast and enhance the bone regeneration process, it would be interesting to observe a long term remodeling of the bone with periosteum stripped in the future study.

Conclusion

Periosteum removal can result in an early apoptosis to the osteocytes in outer layer of skull and a reduction of sclerostin within that area.

Conflict of interest

The authors do not have any conflict of interest.

References

2. Chang H, Knothe Tate ML (2012) Concise review: the periosteum: tapping into a reservoir of clinically useful progenitor cells. Stem Cells Transl Med 1: 480-491. [Crossref]

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