Histological and micro Computed Tomography analysis of a femoral stress fracture associated with prolonged bisphosphonate use

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Summary

Background. The origin of atypical femoral fractures (AFF) associated with bisphosphonate therapy remains to be elucidated. In this study, a biopsy of an AFF site is analyzed to determine whether microdamage and/or morphological changes are present in the area of the AFF.

Material and methods. Cortical bone from an AFF region was obtained during a preventive stabilization in a patient with a symptomatic AFF. This bone was scanned using microCT (resolution=0.01 mm), stained with basic fuchsin and analyzed histologically.

Results. The diameter of the Haversian canals was higher in the vicinity of the AFF compared to the bone further away from the AFF. The bone mineral density within the cortex ranged from 1020 to 1080 mg HA/cm³. We observed penetration of basic fuchsin into the matrix, which is a tell-tale sign of diffuse damage.

Discussion. The higher diameter of haversian canals is likely to result in higher local stresses and consequently increased microdamage. The diffuse microdamage in the biopsy may furthermore be directly related to bisphosphonate use, preventing repair of microdamage, and consequently the development of the AFF.

Conclusion. Increased porosity of the cortex and accumulation of microdamage might have lead to a stress fracture and ultimately a complete AFF.

KEY WORDS: bisphosphonate; spontaneous fracture; histology; atypical femoral fracture.
The integration time was set at 1000 ms, the beam intensity at 55 kV, the current at 145 mA, and the resolution at 0.010 mm. Three-dimensional reconstructions were made with the cone-beam reconstruction algorithm. The contours of the fragment of bone were manually traced, excluding the main fracture and tiny cracks surrounding it as well as the fracture callus. Subsequently, 30 subvolumes were defined in the scan using a rectangular grid. Each subvolume was segmented using the standard procedure (sigma=0.5, support=1, threshold=613 mg HA/cm³) and both the vBMD and the mean diameter of the Haversian channels were calculated using the standard evaluation software (UCT Evaluation v6.5-3, Scanco Medical A.G., Brütisellen, Switzerland). Finally, the contours of the fracture callus present on the endocortical and periosteal surfaces of the cortical bone were manually traced and the vBMD of these newly formed bone layers were compared to that of the cortical bone matrix.

**Histology**

For histology, the biopsy was incubated en block in a series of 1% basic fuchsin (Sigma-Aldrich, St. Louis, MO, USA) solutions in alcohol (supplementary material 2), after which the biopsy was rinsed in 100% alcohol for 1h. Subsequently, the dehydrated and undecalcified biopsy was embedded in methylmethacrylate (MMA). Sections of 5μm thickness were made every 500 μm over a total distance of approximately 4 cm. Sections were analyzed using light microscopy for the presence of microdamage.

**Results**

**MicroCT**

The stress fracture was clearly visible on the microCT scans (Figure 1B). Trabecular bone-like material, likely representing the persisting fracture callus, was clearly visible on the endosteal and periosteal surface of the biopsy (Figure 1A, B). This trabecular material contained woven bone (data not shown), and was calcified, but much less so than the native bone matrix, i.e. the mineral density distribution of the old bone was on average 1070 mg HA/cm³, versus 787 mg HA/cm³ for the bone formed on the endosteal and periosteal surface. The vBMD slightly varied in the endosteal-periosteal direction (x-axis in Figure 3) as well as in the proximal-distal direction within the cortical bone (Vol0 to vol4 in Figure 3), but this variation did not exceed the margin of error for the scan. The vBMD ranged from 1020 to 1080 mg HA/cm³ (Figure 3a).

Haversian channels on the periosteal side of the biopsy were of a normal size (0.050 - 0.080 mm; right hand of the x-axis in Figure 3B) (9, 10). On the endosteal side (Figure 3; left hand of the x-axis in 3b), especially in the neighbourhood of the crack (Figure 3; vol0 to vol2 in 3b), the average diameter was markedly higher than on the periosteal side (Figure 3b; 0.260-0.330 mm).

**Histology**

The stress fracture was clearly visible in the histological sections (Figure 4). In the region around the fracture, in the lamellae, microcracks were visible and basic fuchsin (red) had penetrated the matrix around the fracture. On the periosteal and endosteal side of the cortex a layer of bone was visible with a trabecular appearance, similar to the virtual 2 dimensional microCT images (Figure 2). Osteoclasts were not detectable due to the intense staining with basic fuchsin. There was a lot of erythrocytes visible in the specimen. No blood vessels were observed within the fracture.

**Discussion**

An interesting observation was the large osteonal diameter at the endosteal side of the bone (11). The increased osteon diameter near the fracture site is unlikely to be an effect of the fracture, because the bisphosphonates will have stopped the osteoclast activity even before the fracture occurred. Bisphosphonates stop the resorption of bone by osteoclasts but might thereby also be influencing the osteoblasts through disturbed osteoclast-osteoblast coupling (12).

The vBMD within the cortex ranged from 1020 to 1080 mg HA/cm³ and seemed to be slightly lower on the periosteal surface. However, these differences in vBMD are so small, that it cannot be reliably concluded that mineralization is lower on the periosteal surface than the endosteal surface. Arrest of natural remodelling in the bone can lead to an accumulation of unrepaired microdamage in the matrix, leading to fatigue fractures (13, 14). We observed penetration of basic fuchsin into the matrix, which is a tell-tale sign of diffuse damage. It is possible that this microdamage pre-existed prior to the occurrence of the fracture, and has led to the development of the stress fracture (personal communication with Prof. M.B. Schaffler), but it is impossible to exclude the possibility that the microdamage is the effect rather than the cause of the stress fracture. It should be noted that, at the site where the bone was removed, marks of the point of the clamp with which the bone was removed are visible in the histological sections without microcracks, suggesting that damaging the bone does not automatically lead to microdamage in the matrix, and this strengthens the suggestion that the damage preceded the fracture.

The fracture callus was clearly present at 4 weeks after the initial presentation, although the fracture must have been present for a longer period of time. During normal fracture healing, the peak of soft callus is found at 7-9 days and calcified callus appears from 2-6 weeks after fracture and can persist for weeks until bony union (15). The callus in our biopsy had a trabecular-like appearance, and showed no signs of being removed (absence of scalloped surfaces) or remodelled. A limitation of our study is that we do not have the exact date that the stress fracture arose. Unfortunately another limit is that because of our staining method osteoclasts were not visible, but Vigorita et al. suggest they may be present, albeit not active (17). It is not possible to determine precisely whether the callus consisted of bone or calcified cartilage by the methods used in the current study. On microCT the calcification of the callus was lower than that of mature bone. On histology the callus contained woven bone rather than lamellar bone, as would have been expected in mature bone matrix. The findings suggest that the first stages of healing (inflammation, attraction of stem cells) had occurred, but had stopped there. The use of bisphosphonates has possibly prevented progression of the healing by blocking the osteoclast. Blocking of osteoclasts prevents the resorption of damaged bone, but may also impair new bone formation, since resorption of the matrix by osteoclasts releases growth factors such as insulin-like growth factor 1 and...
Figure 1 - A) AP pelvis X-ray with cortical thickening (arrow) on the lateral side of the left femur. B) AP pelvis at 1 year after treatment of the AFF with intramedullary fixation in place.

Figure 2 - A) Virtual 2 dimensional section through the 3 dimensional microCT scan of the entire biopsy. The endosteal side is marked with 'E'. Note the apparently wider Haversian canals on the endosteal side. The fracture is not visible in this virtual section. B) Virtual 2 dimensional section through the 3 dimensional microCT scan of the entire biopsy. The endosteal side is marked with 'E' and periosteal side with 'P'. The fracture is clearly visible in this virtual section.

Figure 3 - A) Quantification of volumetric Bone Mineral Density in the 30 subregions defined within the biopsy. B) Quantification of Haversian canal diameter in the 30 subregions of the biopsy. The fracture runs through vol0, vol1 and vol2, while vol3 and vol4 are situated further away from the fracture. Points on the left hand side of the graph are situated in their respective volumes near the endosteal surface of the biopsy, while points on the right hand of the graph are situated near the periosteal side of the biopsy.
bone morphogenetic proteins among other factors that stimulate osteoblast activity (12).

There was a lot of erythrocytes visible, which most likely were washed into the fracture when the biopsy was taken, as no blood vessels were observed within the fracture. This again suggests that the fracture itself was not progressing beyond a certain stage of the healing process. It has been described that fracture healing is delayed in people after prolonged bisphosphonate use and specifically after AFF (18-20).

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Conflict of interest
Matthijs Paul Somford, Leo J van Ruijven, Peter Kloen, and Astrid D Bakker declare that they have no conflict of interest.

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