Drilling k-wires, what about the osteocytes? An experimental study in rabbits

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Abstract

Introduction The function of osteocytes regarding osteonecrosis has been underestimated for a long time. Recently it has been suggested that apoptosis of osteocytes results in strong osteoclastic bone resorption. Death of osteocytes due to drilling may therefore increase the risk of K-wire loosening. The purposes of our in vivo study were to assess the minimal drill time needed to notice disappearance of osteocytes and to measure the distance of the empty osteocyte lacunae surrounding the drill tract in relation with the insertion time, directly and 4 weeks after drilling Kirschner (K-) wires into the femur and tibia of rabbits.

Materials and methods Trocar tipped K-wires (70 mm length and 0.6 mm thickness) were drilled into the femur and tibia of 14 New Zealand white rabbits [mean body weight 2.81 kg (2.66–3.09 kg)]. Six rabbits were terminated following surgery (t = 0) and eight rabbits were terminated 4 weeks (t = 4) after surgery. Following termination, hematoxylin and eosin stained sections were cut from femur and tibia until the drill hole was visible. The sections were evaluated under a light microscope for the presence or absence of osteocytes in osteocyte lacunae surrounding the drill holes.

Results All osteocyte lacunae were empty around the K-wires in 50 and 87% of the cases, directly and 4 weeks after the surgery, respectively. The osteocytes disappeared especially beyond a drilling time of 37 s (P = 0.011) and 27 s (P = 0.008) at t = 0 and t = 4, respectively. Furthermore, a significantly positive correlation was seen between the distances of the empty osteocyte lacunae surrounding the drill holes in relation with time at t = 0 (P = 0.008) and t = 4 (P = 0.000).

Conclusion Although only drilling without cooling was studied, short drilling times may prevent the disappearance of osteocytes in case cooling is not used in clinical practice as is the case in percutaneous K-wire insertion.

Keywords Animal · Kirschner wire · Drilling · Histology · Insertion time

Introduction

Experiments concerning thermal damage to bone tissue due to drilling Kirschner wires (K-wires) show several variables interfering with heat generation [7, 9–14, 32]. These variables can be categorized into three groups: drilling technique, K-wire characteristics and external factors. The drilling technique is a subtle balance between drilling speed, insertion time and pressure [1, 2, 5, 30, 31, 34]. The characteristics of K-wires differ in diameter and K-wire tip [17, 23, 29]. The main external factor is irrigation with a water spray during drilling [3, 20, 24, 36].

It is believed that critical temperature for bone injury is around 56°C because alkaline phosphatase denaturates at this temperature. Osteocytes, however, may even be more sensitive to heat [23, 24]. Eriksson [9, 10, 12] observed that a temperature of 47°C for only 1 min results in bone resorption.
Bone is capable of remodeling by bone resorbing osteoclasts, bone forming osteoblasts and osteocytes [15, 19, 22, 26]. The latter differentiate out of osteoblasts which have ceased bone production and have become embedded in the bone matrix, in vacuoles cushioned by fluid and large molecules, forming a network [8, 15, 19, 22, 25]. Despite the fact that the osteocytes are the major constituents of bone, their role in bone resorption regulation has remained controversial for a long time [4, 16, 21, 22, 27]. In the past decade, however, understanding of osteocyte physiology has increased dramatically and it has become clear that osteocytes play an important role in bone remodeling [6, 15, 18, 25, 26, 35]. Now we know that micro damage to bone, e.g. by drilling, is associated with an increase in osteocyte death by apoptosis [25]. This process triggers local bone resorption resulting in K-wire loosening, because death of osteocytes turns off the inhibition of osteoclasts [6, 15, 16, 22, 25, 35].

The status of osteocytes after drilling into bone has been investigated before. Thompson [33] describes the absence of osteocytes as an acute cellular reaction, which increases in severity with increase of drilling speed. Pallan et al. [28] continued Thompson’s [33] work and describes the delayed cellular changes in bone after pin insertion which were left in situ for maximum 10 weeks. He also concluded that higher speeds produce relatively higher temperatures and increased tissue response after a long time period [28].

It is well known that in clinical daily practice K-wires are often drilled without proper cooling because most drilling devices do not have an incorporated cooling system and therefore cooling has to be done manually. In case percutaneous drilling is performed, the cooling effect on bone is minimal because of the surrounding soft tissues. An in vivo study was therefore designed which simulated daily practice concerning K-wire drilling to analyze the absence of osteocytes directly \((t = 0)\) and 4 weeks \((t = 4)\) after insertion. Our first aim was to measure the minimal drill time needed to notice disappearance of osteocytes. The second aim was to measure the distance of the disappeared osteocytes around the periphery of the drill holes in relation to drilling time.

**Materials and methods**

**Animals and anesthesia**

A total of 14 healthy, New Zealand white rabbits of female sex weighing a mean of 2.81 kg (2.66–3.09 kg) were used in this investigation. The rabbits were solely housed on a 12 h/12 h (light/dark) cycle and provided with standard diet food and water ad libitum. All animals were housed in the Central Animal Laboratory, Utrecht University, Utrecht, The Netherlands and received care in compliance with the European Convention Guidelines.

The animals were pre-anesthetized with a combination of methadone (10 mg/ml at a dose of 2.5–5.0 mg i.m.), ventraquill (10 mg/ml at a dose of 2.5–5.0 mg i.m.) and etomidaat (2 mg/ml at a dose of 2.0–8.0 mg i.v.).

After introduction of anesthesia the rabbits were cuffed and mechanically ventilated with \(O_2:N_2O\) proportion 1:1, and 2% isoflurane. During the surgical procedure methadone (2.0–5.0 mg i.v.) was given. At the end of the operation nalbuphine (10 mg/ml at a dose of 1.0–2.0 mg/kg i.v.) was administered. After surgery the rabbits were housed at the intensive care for the rest of the day and night. At the end of the study the rabbits were euthanized by an i.v. overdose of pentobarbital.

**Surgical technique**

An operation device was created by the first author (Fig. 1). The operation device consists of a base plate. On this base plate, a dynamic plate was fixed which could be moved up or down. In front of the entire length of this dynamic plate a sideways moving slide was fixed on the base plate. On this sideways moving slide another back-forward slide including the drill was fixed. Forward movement of the drill was initiated using a 1.5 kg weight.
During surgery the rabbit was fixed on the dynamic plate. This plate made it possible that the femur or tibia were on the same height as the K-wire. The sideways moving slide was responsible for the exact position of the K-wire in front of the femur or tibia.

After the rabbits were preanesthetized, X-rays were made to exclude deformities. Thereafter the animal’s hind limb was carefully shaved and prepared with a povidone-iodine solution. After this procedure, the hind limb was fixed on the testing machine. With the animal surgically draped, a straight-line skin incision was made on the lateral aspect of the femur extending from just below the anterior–inferior spine to the distal femur, followed by a straight-line skin incision on the lateral aspect of the tibia extending from just below the joint line proximally to about the joint line distally.

Dissection was carried out down to the periostium. Synthes Trocar tipped K-wires of 70 mm length and 0.6 mm thickness were drilled through the diaphysis. One K-wire was drilled into the femur and one into the tibia. Drilling was performed by a rotary engine fixed at 1,200 rpm. This is the maximum drilling speed used in our daily practice. Cooling was not performed. After insertion, the K-wires were cut short and the K-wire ends were bent to the cortex. After the wounds were closed in layers, X-rays were made to check the position of the K-wires and the condition of the bone. Insertion time could be measured very accurately by analyzing the operations recorded on video camera. All the experiments were performed by the same investigator.

Histological technique

After termination, the femur and tibia were removed from the hind limb and fixed in 4% formaldehyde solution. They were then decalcified, cut transversely next to the K-wire hole, after the K-wire was removed gently by a pair of tweezers, and embedded in paraffin according to standard procedures. Four micrometer thick serial sections were cut until the drill hole was visible, stained with hematoxylin and eosin and evaluated under a light microscope at 400× magnification for the presence or absence of osteocytes in the osteocyte lacunae surrounding the drill holes by a single investigator. The best section was used for evaluation. The distance over which the osteocytes had disappeared perpendicular to the drill holes was measured with an interactive morphometry device (Q-PRODIT, Leica, Cambridge, UK). In each section, four distances from the drill hole to the first osteocyte bearing lacuna were measured and averaged.

Statistics

Pearson’s Chi-square test was used to determine the drilling time needed for osteocytes to disappear. Pearson correlation was used to highlight any significant correlation between the drilling time and the distance of the disappeared osteocytes surrounding the drill holes. A P-value <0.05 was considered statistically significant. The data were analyzed using SPSS 12.0.1 for windows.

Results

After surgery (t = 0) six rabbits were terminated. Both hind limbs were used resulting in 24 assessments. From the remaining eight rabbits only one hind limb was operated. Four weeks later at termination this resulted in 16 assessments (t = 4). Two t = 0 and one t = 4 assessments were lost because no sections could be produced showing the drill hole, leaving 22 t = 0 and 15 t = 4 assessments.

Histological response to drilling was seen in most sections next to the drill hole (Fig. 2). At t = 0 and t = 4, all osteocyte lacunae next to the drill holes were empty in 11/22 (50.0%) and 13/15 (86.7%) of bones, respectively. At t = 4, a drilling time longer than 27 s resulted in a significant loss of osteocytes surrounding the drill holes (P = 0.008). At t = 0, a drilling time longer than 37 s resulted in a significant loss of osteocytes surrounding the drill holes (P = 0.011).

A statistically significant positive correlation was seen between the distance of the empty osteocyte lacunae surrounding the drill hole in relation with drilling time, directly (P = 0.008, Fig. 3a) and 4 weeks after K-wire insertion (P < 0.000, Fig. 3b).

![Fig. 2 Status of osteocytes due to K-wire drilling at 1,200 rpm. Disappeared osteocytes due to drilling are marked by a dot. The osteocytes are encircled. The dotted line indicates the border between the present and disappeared osteocytes (hematoxylin and eosin)](image-url)
Discussion

The two most important findings of this experiment are that (1) osteocytes disappear especially beyond a drilling time of 27 s and (2) that there is a positive correlation between the distance of the empty osteocyte lacunae in relation with drilling time.

As far as we know there are no experiments studying the presence/absence of osteocytes while using drilling time as a variable. Therefore it is interesting to notice that our results, representing daily practice by using a drilling speed of 1,200 rpm, clearly indicate that osteocytes disappear after a drilling time above 27 s in this animal model, when no cooling is used. Pallan [28] and Thompson [33] however, noticed absence of osteocytes already after a drilling time varying from 5 to 12 s. The limitation of their experiments is that the insertion force was not standardized. That could be a possible factor explaining the difference in insertion time with matching histological changes.

The effect on osteocytes in our study was more pronounced 4 weeks after insertion compared to the effect on osteocytes directly after surgery. This confirms the results from Thompson [33] where the effect on osteocytes was more pronounced 72 h after the operation than at either 24 or 48 h. Apparently, not all damage to osteocytes is acute and microscopically visible, directly after K-wire insertion. This can be explained by the study of Eriksson et al. [13].

They compared histology and histochemistry for detection of bone viability directly after surgery. The histological sections revealed a 200 μm wide zone of empty osteocyte lacunae around the periphery of the drill hole while histochemistry of the same bone specimens showed a zone of 500 μm lacking diaphorase enzyme activity. This indicates that the empty osteocyte lacunae directly after surgery underestimates the extent of the drilling trauma and becomes clear as time passes. Neovascularisation is probably a reason for the increase of osteocyte disappearance in time. During heating, due to drilling, blood flow will stop in minor vessels preventing the clearance of the osteocyte lacunae [10, 13]. After neovascularisation the osteocyte lacunae can be cleared finally.

Our study also showed an interesting positive correlation between the distance of the empty osteocyte lacunae in relation with time. The zones of disappeared osteocytes varied from 106 to 1,285 μm and the drilling times from 41 to 262 s. A correlation between drill time and distance was not shown before. What we do know is that Eriksson et al. showed a 200 μm wide zone of empty osteocyte lacunae around the periphery of the hole after drilling into femurs of New Zealand white rabbits while conventional irrigation was administered [13]. These results are difficult to compare because Eriksson et al. did not measure time, irrigated with saline, did not standardize the insertion force and drilled with a running speed of 20,000 rpm. Furthermore, Pallan [28] showed that the histological changes, including absence of osteocytes, were never apparent more than 250 to 500 μm from the pins [28]. Like Eriksson they did not standardize insertion force and drilling time varied from 5 to 12 s.

No osteoclastic activity was noticed in the already existing cortex. This was expected as Pallan [28] hardly noticed osteoclastic activity after 6 weeks, and only slight osteoclastic activity after 8 and 10 weeks and our experiments covered a time span of only 4 weeks.

We are convinced that delayed tissue response corresponds with late K-wire loosening seen in daily practice. Although we did not compare drilling, with and without
cooling, this study demonstrates the need for a short drill time, less than 27 s to prevent the disappearance of osteocytes and to limit the bone resorption cascade.

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