FULL REGENERATION OF SEGMENTAL BONE DEFECTS USING POROUS TITANIUM IMPLANTS LOADED WITH BMP-2 CONTAINING FIBRIN GELS

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Abstract

Regeneration of load-bearing segmental bone defects is a major challenge in trauma and orthopaedic surgery. The ideal bone graft substitute is a biomaterial that provides immediate mechanical stability, while stimulating bone regeneration to completely bridge defects over a short period. Therefore, selective laser melted porous titanium, designed and fine-tuned to tolerate full load-bearing, was filled with a physiologically concentrated fibrin gel loaded with bone morphogenetic protein-2 (BMP-2). This biomaterial was used to graft critical-sized femoral bone defects in rats. As a control, porous titanium implants were either left empty or filled with a fibrin gels without BMP-2. We evaluated bone regeneration, bone quality and mechanical strength of grafted femora using in vivo and ex vivo µCT scanning, histology, and torsion testing. This biomaterial completely regenerated and bridged the critical-sized bone defects within eight weeks. After twelve weeks, femora were anatomically re-shaped and revealed open medullary cavities. More importantly, new bone was formed throughout the entire porous titanium implants and grafted femora regained more than their innate mechanical stability; torsional strength exceeded twice their original strength. In conclusion, combining porous titanium implants with a physiologically concentrated fibrin gels loaded with BMP-2 improved bone regeneration in load-bearing segmental defects. This material combination now awaits its evaluation in larger animal models to show its suitability for grafting load-bearing defects in trauma and orthopaedic surgery.

Keywords: BMP, bone graft, bone regeneration, fibrin, metal surface treatment, scaffold, titanium.

Introduction

A major challenge in trauma and orthopaedic surgery is to successfully repair load-bearing segmental bone defects (Einhorn, 1995). This often requires the use of bone grafts or bone graft substitutes to improve bone regeneration by providing an osteoconductive matrix, offering mechanical support, or an osteoinductive and/or osteogenic stimulus (Giammoudis et al., 2011). The golden standard bone graft is still autologous bone (Pape et al., 2010), but the amount of bone that can be harvested is limited and associated with complications in 10-40% (Banwart et al., 1995). These disadvantages motivate the development of biomaterials that can be used as bone graft substitutes (Langer and Vacanti, 1993).

A biomaterial that has the potential to become a bone graft substitute is porous titanium (Alvarez and Nakajima, 2009; Murr et al., 2010; Ryan et al., 2006). Nowadays, porous titanium can be manufactured using additive manufacturing techniques such as selective laser melting (SLM) (Hollander et al., 2006). This enables the design of porous titanium so that its structure and mechanical strength remains suitable to function as a load-bearing osteoconductive matrix in segmental bone defects (Van der Stok et al., 2013a). Osseointegration of titanium is optimised through relatively simple chemical and heat treatments that alter the surface chemistry and (nano-) topography (Amin Yavari et al., 2014a). Thereby, the bioinert titanium surface changes into a bioactive surface that allows spontaneous apatite formation and stimulates proliferation and osteogenic differentiation of osteoprogenitor cells (Amin Yavari et al., 2014b). This surface-treated porous titanium forms a load-bearing osteoconductive matrix, but stimulating bone regeneration and adequate bridging of segmental bone defects may be further improved by addition of effective biological stimuli (i.e. osteoinductive cytokines) (de Wild et al., 2013; Van der Stok et al., 2013a).

Bone morphogenetic proteins (BMPs) such as BMP-2 and BMP-7 play a major role in bone regeneration as osteoinductive cytokines (Urist, 1965). Their osteoinductive effects have been established in a wide range of species,
varying from mice and rats to humans (Murakami et al., 2002). BMP-2 and BMP-7 have received USA Food and Drugs Administration (FDA) approval for use in trauma and orthopaedic surgery (Senta et al., 2009), but their clinical success is limited (Khan and Lane, 2004). This might be because a supra-physiological dosage of BMP needs to be loaded onto an absorbable collagen sponge to reach an effect (Terraat et al., 2005). This high dose has been associated with adverse effects including bone tissue overgrowth, ectopic bone formation, inflammation, and even carcinogenicity (Carragee et al., 2011; Woo, 2013).

To overcome this, numerous slow-release systems have been developed. Interestingly, these slow-release systems, allowing for controlled release of BMP-2 during several weeks, do not resemble the natural bone regeneration process in which BMP-2 is mainly released during the first few days (Cho et al., 2002; Gerstenfeld et al., 2003).

Bone regeneration starts with the formation of a fibrin clot, often referred to as the fracture haematoma. This fibrin clot forms the natural binding reservoir for osteoinductive cytokines such as BMP-2 (Gerstenfeld et al., 2003; Martino et al., 2013) and is formed through conversion of fibrinogen by thrombin. Fibrinogen is synthesised in its high molecular weight form, but occurs as a mixture together with partially degraded low molecular weight forms in circulation (Kaijzel et al., 2006). Fibrin gels, made from physiological fibrinogen concentrations (2-4 mg/L), are highly permeable to cells (Kaijzel et al., 2006).

However, at these physiological concentrations, fibrin gels are soft and therefore not suitable for most clinical applications. Consequently, commercially available fibrin sealants contain very high fibrinogen concentrations (50-100 mg/L) (Janmey et al., 2009) at the cost of seriously compromising the favourable cellular permeability of these gels. When incorporated into porous titanium, the use of physiologically concentrated fibrin gels becomes feasible as the metal frame ensures mechanical support, then. The surface-treated porous titanium implants may even improve the network organisation of fibrin fibres (Milleret et al., 2011).

The aim of the current study was to develop a biomaterial capable of improving bone regeneration of segmental bone defects: osteoconductive load-bearing porous titanium filled with physiologically concentrated fibrin gels releasing BMP-2. For this novel combination, the BMP-2 releasing fibrin gel was prepared from purified high molecular weight (HMW) fibrinogen, since HMW fibrinogen increases angiogenesis in vitro and in vivo. (Kaijzel et al., 2006). To determine whether the angiogenic HMW fibrin gel alone is capable of increasing bone regeneration, porous titanium implants were also filled with HMW fibrin gels without BMP-2 and compared to unfractionated (UNF) fibrin gels. Porous titanium implants incorporated with the three above described fibrin gels were compared to empty porous titanium implants in a critical-sized load-bearing segmental femur defect in rats using in vivo (4, 8 and 12 weeks) and ex vivo (after 12 weeks) μCT scans, histology, and biomechanical torsion tests.

Materials and Methods

Porous titanium implants

Porous titanium implants were produced from Ti6Al4V ELI powder (ASTM B348, grade 23) using selective laser melting (SLM, Layerwise N.V., Leuven, Belgium). The implants were a copy of the replaced femoral bone segment and had a height of 6 mm, a maximum outer diameter of 5 mm and a minimal inner diameter of 1.3 mm (leaving an open medullary canal). The porous architecture was based on a dodecahedron unit cell with a strut width of 120 μm and an average pore size of 500 μm, to result in 55 mm³ porous volume. All implants underwent a post-production alkali-acid-heat treatment consisting of (1) immersion in a 5 M aqueous NaOH solution at 60 °C for 24 h; (2) immersion in water at 40 °C for 24 h; (3) immersion in 0.5 M HCl at 40 °C for 24 h; (4) heating to 600 °C at a rate of 5 °C/min in an electric furnace at ambient air pressure, keeping the temperature at 600 °C for 1 h, and subsequent natural cooling (Amin Yavari et al., 2014b).

Reproducibility of porous implant architecture (e.g., pore size, titanium strut thickness and porosity) was verified by μCT (SkyScan 1076; Bruker micro-CT N.V., Kontich, Belgium).

Fibrin gel preparation

Fibrin gel preparation was done as previously described (Holm et al., 1985; Kaijzel et al., 2006). Briefly, plasminogen-rich unfractionated human fibrinogen (Chromogenix, Mölndal, Sweden) was dissolved in Tris buffer (10 mM Tris/HCl, pH 7.4) to a concentration of 5 mg/mL. Saturated (NH₄)₂SO₄ was slowly added to a final concentration of 19 % (v/v) and the solution was mixed for 30 min at room temperature prior to centrifugation for 10 min at 2,000 × g. Repetition of this precipitation step resulted in a HMW fibrinogen pellet (~ 99 % purity), which were dissolved in 5 mL of saline (0.9 % NaCl) and then dialysed against M199 culture medium, as was the UNF fibrinogen. Purity was determined using standard non-reducing sodium dodecylsulphate polyacrylamide gel electrophoresis and concentrations were calculated using the molar extinction coefficient of fibrinogen (EI 1 % 280 nm for fibrinogen is 15.8). The preparations were stored in single-use aliquots at -80 °C until further use.

In a custom-made mould, the porous titanium implants were filled with 55 μL of either HMW fibrinogen (2 mg/mL, HMW-Fb) or UNF fibrinogen (2 mg/mL, UNF-Fb) which were clotted with 0.5 IU/mL of thrombin (Global Siemens Healthcare, Erlangen, Germany) dissolved in a 4.5 mM calcium chloride buffer (Baxter, Utrecht, Netherlands) in a 8.5:1 ratio. HMW fibrin gels with BMP-2 (HWM-BMP-Fb) were made by adding 3 μg BMP-2 (Shanghai Rebone Biomaterials Co., China) in 1 mM saline solution to the HMW fibrinogen solution before clotting. Prior to implantation, after clotting the fibrin-filled implants were wrapped in Parafilm® and incubated for 15-18 h at 6 °C to allow completion of fibrin gelling.
Scanning electron microscopy (SEM)
To determine filling efficacy and to characterise the structure of the fibrin networks polymerised from HMW and UNF fibrinogen, SEM was used as follows: implants were filled with fibrin gels and fixed in 3 % glutaraldehyde for 24 h and rinsed with sodium phosphate buffer (0.1 M, pH 7.2-7.4; Merck). Samples were then consecutively dehydrated in ascending alcohol concentrations (30, 50, 70 and 90 % v/v) with three final incubations in 100 % ethanol for 10 min each. Probes were critical-point-dried in liquid CO₂, and then sputtered with a 30 nm gold layer. Samples were analysed in FEI/Philips XL 30 FEG ESEM (Philips) in a high vacuum environment.

Load-bearing segmental bone defects
Critical-sized segmental bone defects were made in the femora of 40 male 16-weeks-old Wistar rats (446 ± 32 g). Rats were divided into four experimental groups receiving porous titanium implants filled with HMW-BMP-Fb, HMW-Fb and UNF-Fb or were left empty (empty). The Animal Ethics Committee of the Erasmus University approved the study and Dutch guidelines for care and use of laboratory animals were followed. Before surgery, rats received subcutaneous injections of antibiotics (enrofloxacin, 5 mg/kg body weight) and pain medication (buprenorphine, 0.05 mg/kg body weight). Surgery was performed aseptically under general anaesthesia (1-3.5 % isoflurane). The right femur was exposed through a lateral skin incision and separation of underlying fascia. Using three proximal and three distal screws, a 23 × 3 × 2 mm polyether ether ketone (PEEK) plate (RISystem, Davos Platz, Switzerland) was fixed to the femur anterolateral plane. Periosteum was removed over 8 mm of the mid-diaphyseal region before a 6 mm cortical bone segment was removed with a wire saw and a tailor-made saw guide. Subsequently, a porous titanium implant was implanted press-fit into the defect. Finally, fascia and skin were sutured. Subcutaneous injection of pain medication (buprenorphine, 0.05 mg/kg body weight) was given twice a day for the following three days. Rats were sacrificed after twelve weeks with an overdose of pentobarbital (200 mg/kg body weight).

µCT evaluation
Bone regeneration was measured by in vivo µCT scans (SkyScan 1076; Bruker micro-CT N.V., Kontich, Belgium) at four, eight and twelve weeks, and by ex vivo µCT scans on isolated grafted femora at the end of the experiment. Rats were kept under general anaesthesia (1-3.5 % isoflurane) during in vivo µCT scans at 35 μm resolution (95 kV, 105 μA current, 1.0 mm Al/0.25 mm Cu filter, and 0.75° rotation step, 14 min scan). Ex vivo µCT scans were acquired at 18 μm resolution (95 kV, 100 μA current, 1.0 mm Al/0.25 mm Cu filter, and 0.5° rotation step). µCT scan images were reconstructed using volumetric reconstruction software NRecon version 1.6.6 (Bruker micro-CT N.V., Kontich, Belgium).

Bone regeneration was expressed as bone volume (BV), which was measured at four specific regions: 1) total BV: the total volume of bone formed within the 6 mm defect; 2) porous BV: the bone formed inside the porous space of the titanium implants; 3) outer BV: the bone formed outside the porous titanium implants; and 4) inner BV: the bone formed in the medullary canal of the implants. BV values were measured using CTanalyser version 1.13 (Bruker micro-CT N.V., Kontich, Belgium). First the specific region was selected, then the titanium and its border artefacts was excluded from images using a global threshold with a value between titanium and bone and removal of an extra 35 μm border (size of one pixel) surrounding the titanium. Subsequently the bone was extracted by using a second global threshold that differentiated between bone and soft tissue. The global threshold values were chosen on visual inspection and were kept constant for all scans. Bone bridging was assessed on ex vivo scans with DataViewer 1.4 (Bruker micro-CT N.V., Kontich, Belgium). Complete bone bridging was defined as bridging of three or more cortices counted on ex vivo scans in the coronal and sagittal plane. Bone bridging was quantified by measuring the shortest remaining gap size between bone formed at the proximal and distal site of the 6 mm bone defect.

Histological evaluation
Histology was performed on two femora per group that represented the mean of the whole group. To select these two femora, all ten grafted femora were sorted according to their total BV after twelve weeks and the two femora closest to the averaged value were chosen. Harvested femora were fixed in 10 % neutral buffered formalin solution for two days, dehydrated in ascending alcohol concentrations (70 to 100 % v/v), and finally embedded in methyl methacrylate (MMA). Sections of ~20 μm were obtained using a diamond saw (Leica SP1600, Rijswijk, The Netherlands) and stained with basic fuchsin 0.3 % (w/v) solution and methylene blue 1 % (w/v) solution to stain bone purple and fibrous tissue blue, respectively. Serial sections were then screened for bone formation, bone-implant contact and bone bridging.

Biomechanical tests
Mechanical strengths of grafted femora were measured by a torsion test conducted on the remaining eight femora of each group. Three contralateral femora, serving as a reference of intact femora, were included as controls. After harvesting the femora, soft tissues and PEEK plates were carefully removed. Specimens were kept in 10 % neutral buffered formalin solution for two days, minimising the effects of formalin conservation on mechanical properties (Unger et al., 2010), and then transferred to phosphate buffered saline. Subsequently, both ends of each femur were embedded in a cold-cured epoxy resin (Technovit 4071, Heraeus Kulzer, Germany). On the upper clamping side, a Cardan joint was used to ensure pure rotation without bending. The lower sides were simply fixed. Tests were performed until failure with a rotation rate of 0.5° s⁻¹ using a static mechanical testing machine that could apply a maximum torque of 450 N.mm (Zwick GmbH, Ulm, Germany). Torsional strength (maximum torque to failure, N.mm) was determined.
Statistics
Statistical analyses were performed using SPSS Statistics 20.0 (SPSS, Inc.). Data are presented as means with standard deviations. One-way analysis of variation (ANOVA) and subsequent post hoc pairwise comparisons with Bonferroni adjustment were used to test for differences between the four groups. A power calculation (β-value > 0.80, SD ~ 25 %) was made to find a true difference in total BV of at least 35 %. Based on this calculation, n = 10 was required. A p < 0.05 was considered statistically significant.

Results
Porous titanium implants incorporated with fibrin gels
Porous titanium implants were produced using SLM in the anatomical shape of the surgically removed cortical bone segment (Fig. 1a). The implants had a porosity of 85 % and a pore size ranging from 460-670 µm (Table 1). The alkali-acid-heat treatment resulted in a titanium oxide layer with an irregular nano-scale features (Table 1). Macroscopic inspection and SEM analyses verified that the pores of the titanium implants were completely filled with fibrin gel (Fig. 1a-b). The protein fibres of both fibrin gels attached intimately to the surface-treated titanium (Fig. 1c). In addition, SEM showed clear differences between HMW-Fb and UNF-Fb gels with respect to their nanofiber structures: as compared to the fibre network formed by unfractionated fibrinogen, resulting in a much denser structures with smaller average pore diameters and

Table 1. Properties of porous titanium implants (Amin Yavari et al., 2014b; Van der Stok et al., 2013a).

<table>
<thead>
<tr>
<th>Titanium implant</th>
<th>165 ± 43 µm</th>
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<tbody>
<tr>
<td>Pore size</td>
<td>577 ± 146 µm (range 460-670 µm)</td>
</tr>
<tr>
<td>Porosity</td>
<td>85 %</td>
</tr>
<tr>
<td>Pore volume</td>
<td>55 mm³</td>
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<tr>
<td>Compression strength</td>
<td>14 MPa</td>
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<tr>
<td>Young’s modulus</td>
<td>0.4 GPa</td>
</tr>
<tr>
<td>Surface area / volume</td>
<td>0.034 µm²</td>
</tr>
<tr>
<td>Surface composition</td>
<td>Oxygen 35 %</td>
</tr>
<tr>
<td></td>
<td>Titanium 60 %</td>
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<tr>
<td></td>
<td>Vanadium 2 %</td>
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<td>Aluminium 3 %</td>
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Surface topography (SEM)

Fig. 1. SEM images of fibrin loaded porous titanium implants. Macroscopic overview, and enlarged details, of a fully fibrin-filled implant (a, b). The fibrin fibres are tightly bound to the implant surface (c). The fibre network resulting from unfractionated fibrinogen is rather dense with thin fibres (d). The fibrin network from HMW fibrinogen reveals a more-open, better-permeable, structure with slightly thicker fibres (e).
thinner fibres (Fig. 1d), polymerisation of high molecular weight fibrinogen appeared to form a more open network with relatively thicker fibres (Fig. 1e).

**Load-bearing segmental bone defects**
All rats were able to tolerate weight-bearing activities immediately after surgery; the implantation sites healed without complications and all animals remained healthy during the follow-up.

**µCT evaluation**
Porous titanium implants with HMW-BMP-Fb gels effectively stimulated bone regeneration (Fig. 2). Within four weeks, bone regeneration had occurred throughout the entire length of the porous titanium implants and after eight weeks, bridging of the defect was complete. Only minimal bone regeneration was observed in those defects grafted with HMW-Fb containing porous titanium implants and consequently failed to bridge (Fig. 2). The load-bearing segmental defects grafted with HMW-BMP-Fb containing porous titanium implants fully restored the original bone architecture after twelve weeks (Fig. 2 and 3). Incorporation of HMW-Fb or UNF-Fb did not seem to outperform the empty porous titanium implants (Fig. 3).

Quantitative analysis of regenerated bone, based on in vivo µCT scans, showed that the total BVs of the HMW-BMP-Fb group increased at each time point and reached an average of 65.1 ± 14.9 mm$^3$ after twelve weeks (Fig. 4a). This was significantly higher than all the three control groups. Neither the total BV of the HMW-Fb (37.7 ± 26.4 mm$^3$) group nor that of the UNF-Fb (32.1 ± 13.4 mm$^3$) group was significantly different from

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Fig. 2. Representative longitudinal µCT scans illustrating the bone regeneration process. In vivo scans of defects grafted with porous titanium implants incorporated with high molecular weight fibrin with BMP-2 (HMW-BMP-Fb group) and high molecular weight without BMP-2 (HMW-Fb group) after four (a1-2), eight (b1-2) and twelve weeks (c1-2) as well as ex vivo after twelve weeks (d1-2). In the HMW-BMP-Fb group, rapid bone regeneration throughout the complete length of the defect is already observed after four weeks (a1, arrows). Between eight and twelve weeks, the cortex and medullary canal (indicated by ‘m’) are restored in their original shape (b1, c1, and d1). In the HMW-Fb group, bone regeneration is only observed at the proximal and distal side of the porous implants (a2, arrows); this bone is predominantly situated in the medullary canal and insufficiently bridging the defect (d2). Distally from the titanium implants, bone resorption is observed between eight and twelve weeks (asterisk). Bar indicates 1 mm.
the total BV of the empty group (33.7 ± 16.8 mm³) (Fig. 4a). Also, the porous BV and outer BV of the HMW-BMP-Fb group were significantly higher than that of all three control groups (Fig. 4b-c). After twelve weeks, 51 ± 8 % of the available pore space of the titanium implants with HMW-BMP-Fb gels was filled with regenerated bone, twice as much as in the HMW-Fb (24 ± 18 %) and UNF-Fb (21 ± 5 %) group, respectively. The inner BV of the HMW-BMP-Fb group (3.9 ± 1.6 mm³) significantly increased as compared to the control groups after four weeks. Contrary to the HMW-Fb and UNF-Fb groups, the inner BV of the HMW-BMP-Fb decreased over time (3.1 ± 1.3 mm³ at 8 weeks and 2.6 ± 1.3 mm³ at 12 weeks) and even became significantly less than in the inner BV of the HMW-Fb group (Fig. 4d).

Bone bridging, determined on ex vivo µCT scans at twelve weeks, was only seen in the cortical defects grafted with porous titanium implants filled with HMW-BMP-Fb gels (Fig. 5). Seven defects were completely bridged, and the average remaining gap size in the three defects that were not bridged was 0.8 ± 0.1 mm (Fig. 6a). The remaining gap size in the other three experimental groups was 1.8 ± 1.6 mm (HMW-Fb group), 1.9 ± 0.9 mm (UNF-Fb group) and 1.8 ± 1.4 mm (empty group), respectively.

**Histological evaluation**

Bone quality, assessed using light microscopy, showed that in the HMW-BMP-Fb group bone was formed almost exclusively at the site of the original cortex and an intimate contact between the regenerated bone and the titanium implant was found throughout the entire length of the defect (Fig. 7d vs. a-c). In the HMW-Fb (Fig. 7c), UNF-Fb (Fig. 7b) and empty groups (Fig. 7a), bone formation occurred predominantly at the distal and proximal sites of the titanium implants that were close to the adjacent cortical bone. This bone formation never extended throughout the entire length of the porous implant. The remaining gaps in the defects were rather filled with amorphous fibrous tissue (Fig. 7a, b, and c). Strikingly, in the HMW-BMP-Fb group an open medullary canal was observed after twelve weeks (Fig. 7d), whereas in all control groups bone formation was blocking the medullary canal (Fig. 7a-c).

**Biomechanical evaluation**

Femora of the HMW-BMP-Fb group reached a significantly higher maximum torque than the three control groups (Fig. 6b). The specimens were more than twice as strong as control femora (248 %) and six femora of this HMW-BMP-Fb group were able to resist the maximum torque (450 N.mm) without breaking. Femora of the HMW-Fb group did not differ in maximum torque from the femora of the UNF-Fb or empty group. The average maximum torques were 60 % (HMW-Fb group, 86 ± 29 N.mm), 51 % (UNF-Fb group, 75 ± 20 N.mm), and 53 % (empty group, 77 ± 53 N.mm), respectively, of the average maximum torque measured for control femora (146 ± 19 N.mm) (Fig. 6b).

**Discussion**

An ideal biomaterial, that can be used as a bone graft substitute, should be able to fully regenerate and bridge load-bearing segmental bone defects within a short period of time (Langer and Vacanti, 1993). Our combination of surface-treated porous titanium with BMP-2 containing...
Fig. 4. Longitudinal quantification of bone regeneration. In vivo µCT scans after four, eight and twelve weeks; total BV (a), defined as all bone formed within the 6 mm defect. Outer BV (b), defined as bone formed outside the titanium implants. Porous BV (c), defined as bone formed inside the porous space of the titanium implants. Inner BV (d), defined as bone formed in the medullary canal of the titanium implants. Values are expressed as mean and SD (n = 10 per group), and a one-way ANOVA test followed by a post-hoc Bonferroni correction was performed to test for statistical significant difference at each time point; p < 0.05 was considered as statistically significant, vertical bars indicate the significant differences found between the groups.

Fig. 5. Illustration of bone bridging. Representative 3D µCT images showing the average extend of bone bridging of the empty (a), UNF-Fb (b), HMW-Fb (c), as well as the HMW-BMP-Fb (d) group. Porous titanium implants appear in transparent grey, whereas bone appears in yellow/dark grey.
Fig. 6. Bone bridging and mechanical strength. The remaining gap size after twelve weeks was used to indicate bridging success (a). Mechanical femoral strength after implantation of porous titanium implants measured by torsion testing (b). As a positive control, three control femora were included to provide a reference of a normal strength of femora during torsion testing (b, control). Values are expressed as mean and SD, and a one-way ANOVA test followed by a post-hoc Bonferroni correction was performed to test for statistical significant differences at each time point; $p < 0.05$ was considered as statistically significant, horizontal bars indicate significant differences between groups.

Fig. 7. Histological evaluation of bone bridging. Representative transversal sections of femur defects twelve weeks after implantation of porous titanium implants; empty (a), or incorporated with UNF-Fb (b), HMW-Fb (c) or HMW-BMP-Fb gels (d). Magnification reveals re-colonisation of the medulla with small round-shaped cells of a typical bone marrow stroma appearance. Sections are stained with basic fuchsin and methylene blue. Basic fuchsin stains bone purple, methylene blue stains fibrous tissue blue. Black bar indicates 1 mm.
physiologically concentrated fibrin gels fully regenerated segmental bone defects in rat femora (Fig. 2 and 5) and fully recovered their mechanical strength (Fig. 6). Moreover, combining osteoconductive titanium with an osteoinductive BMP-2 releasing fibrin gel resulted in “guided” bone regeneration; i.e. new bone was formed throughout the porous titanium implants (a1) without bridging the entire defect (b1). In contrast, HMW-BMP-Fb gels (loaded with the same dose of BMP-2) led to complete bridging with restoration of the medullary canal (a2) and the cortex (b2). Bar indicates 1 mm.

Biomaterials used for load-bearing segmental defects should offer sufficient support to withstand mechanical loading (Giannoudis et al., 2011). A material with such mechanical properties is titanium, and solid titanium implants have been very successfully used in trauma and orthopaedic surgery over the past decades (Learmonth et al., 2007). However, the notable biomechanical mismatch between solid titanium implants and surrounding bone frequently leads to stress-shielding, subsequent bone resorption and implant loosening (Niinomi, 2008). This limitation can be overcome by using mechanically optimised porous titanium implants (Murr et al., 2011; Ryan et al., 2009), the development of which greatly benefited from the introduction of additive manufacturing techniques. Techniques such as selective laser melting (Hollander et al., 2006; Mullen et al., 2009; Stamp et al., 2009), electron beam melting (EBM) (Heinl et al., 2008; Hrabe et al., 2011; Ponader et al., 2010) or similar additive manufacturing techniques (Bandyopadhyay et al., 2010; Bandyopadhyay et al., 2009) allow for a personalised, anatomical implant design and control of its structural and mechanical properties alike. Our SLM-based, femur-shaped implants possessed mechanical properties within the physiological range of the host bone, while its fully interconnected porous structure is considered to be within the range required for osteoconduction (Table 1) (Van der Stok et al., 2013a). Furthermore, the fatigue properties of the porous implants indicate that the biomechanical support is temporary (Amin Yavari et al., 2013). The implants were therefore capable of offering sufficient mechanical support in vivo, while stimulating bone regeneration through osteoconduction in the segmental bone defects (Van der Stok et al., 2013a; Van der Stok et al., 2013b).

In addition to mechanical support, biomaterials should also offer a surface that facilitates osseointegration, i.e. intimate apposition of bone matrix onto the implant surface (Puleo and Nanci, 1999). The bone-implant interface of most metallic biomaterials including titanium usually consists of an interfacial fibrous-like layer (also called

**Fig. 8.** Direct comparison between BMP-2 release from fibrin and gelatin on bone regeneration. In the same in vivo model, using the same type of porous titanium implants and same batch of BMP-2, results from using HMW-BMP-Fb gels (this work) or gelatin nanosphere gels (Van der Stok et al., 2013b) were compared. Using gelatin nanosphere gels loaded with 3 µg BMP-2 predominantly led to bone regeneration outside or inside the porous titanium implants (a1) without bridging the entire defect (b1). In contrast, HMW-BMP-Fb gels (loaded with the same dose of BMP-2) led to complete bridging with restoration of the medullary canal (a2) and the cortex (b2). Bar indicates 1 mm.
Bone graft substitutes should also be able to induce bone regeneration, which can be induced by a variety of bone morphogenetic proteins (BMPs), including BMP-2 and BMP-7 (Groeneveld and Burger, 2000). BMP-2 is mainly released during the first few days of the natural bone regeneration process, and BMP-7 plays a more important role during the later phase (Gerstenfeld et al., 2003). Both BMP-2 and BMP-7 received FDA-approval (Termaat et al., 2005), but their use in humans is currently heavily debated (Carragee et al., 2011). Although the osteoinductive effect of BMP-2 has been demonstrated in a wide variety of species (including rats, rabbits, dogs, sheep and non-human primates) (An and Friedman, 1999), it is often argued that one must be cautious in assuming that stromal cells from other species may serve as models for inducible osteogenesis in human marrow stromal cells (Diefenderfer et al., 2003). BMP-induced side effects, including cyst-like bone formation and soft tissue swelling, are likely caused by supra-physiological dosages used in humans (Carragee et al., 2011) and these adverse effects were recently reproduced in a similar in vivo model as used in this study with BMP-2 concentrations exceeding 20 µg per defect (Angle et al., 2012; Zara et al., 2011). In contrast, a dose between 2.5 and 10 µg was found to be safe and effective for various other BMP release systems including alginate-based (Boerckel et al., 2011), poly-L-lactic acid (PLLA)-based (Wei et al., 2007) or silk-based (Bessa et al., 2010) scaffolds. Based on these results we used 3 µg BMP-2 per implant. Furthermore, Schmoekel et al. demonstrated that with less soluble nonglycosylated BMP-2, the required cytokine dose could even be further reduced (Schmoekel et al., 2004).

Surface-treated porous titanium was loaded with BMP-2 through incorporation in physiologically concentrated fibrin gels. This is different from fibrin gels that have been used in trauma and orthopaedic surgery as “fibrin glue”, as these sealants are made of highly supra-physiological fibrinogen concentrations (Janney et al., 2009). This supra-physiological concentration (50-100 mg/L) ensures quick and effective clotting, and is therefore primarily used as a haemostatic agent. Supra-physiological fibrinogen concentrations were also used to deliver BMP-2 in several bone defect models (Chung et al., 2007; Kaipel et al., 2012; Kim et al., 2008; Koo et al., 2013; Koo et al., 2012; La et al., 2012; Schmoekel et al., 2004; Schmoekel et al., 2004; Schützenberger et al., 2012; Yang et al., 2010; Yang et al., 2012). Schützenberger et al. showed that fibrin gels outperformed the currently clinically used absorbable collagen sponges as BMP-release properties of fibrin, in contrast to those of collagen, allow to use 85 % less cytokine without compromising the regenerative success (Schützenberger et al., 2012). However, the high fibrinogen concentration of these supra-physiologically concentrated fibrin gels have a limiting effect on cell mobility and ingrowth (Nürnberg et al., 2010; Peterbauer-Scherb et al., 2012). In vivo, fibrin constitutes only 0.25 % of the volume of a blood clot (Weisel, 2004), which was mimicked in the current study by preparing fibrin gels of physiological concentrations (2-4 mg/L). It is tempting to speculate that at these concentrations, fibrin fibres form a more open network that more effectively promoted cell migration and cell ingrowth (Seebach et al., 2014). This open network structure and fibrin fibre adherence was perfectly supported by the surface-treated porous titanium (Fig. 1c). This temporary fibrin network is not expected to remain intact in vivo for more than a few days (Schmokel et al., 2004), but the results obtained in this study suggest that this is sufficient to adequately induce bone regeneration. Our approach mimics physiological fracture healing, during which BMP-2 is entrapped in the spontaneously formed fracture haematoma to induce differentiation of mesenchymal stem cells into osteoblasts (Onishi et al., 1998). These osteoblast subsequently start to produce more BMP-2 and other important osteogenic cytokines to reach a maximum activity after 4-7 days (Kirker-Head, 1995).

Physiologically concentrated fibrin gels prepared from HMW fibrinogen were expected to improve bone regeneration, as compared to gels prepared from UNF fibrinogen, because HMW fibrinogen has been shown to promote angiogenesis in vitro and in vivo in our earlier studies (Kaijzel et al., 2006). UNF fibrin gels contain 30 % low molecular weight fibrinogen and, compared to 100 % HMW fibrinogen, contamination with more than 10 % of LMW fibrinogen gradually decreased the formation of tube-like structure in vitro in a dose-depend manner (Kaijzel et al., 2006). However, implants with HMW-Fb gels alone did not enhance bone regeneration and performed similar to implants with UNF-Fb gels or empty porous titanium implants (Fig. 2 and 5). BMP-2 release from the fibrin gel is apparently providing the only osteoinductive stimulus, while HMW-Fb may still improve cell migration and angiogenesis. Whether using HMW-Fb to release BMP-2 is better than using UNF-Fb cannot be answered here and is a limitation of our study. However, HMW-fibrin gels with BMP-2 clearly outperformed previously used gelatin-based nanoparticle gels with BMP-2 (Van der Stok et al., 2013b), indicating the type of BMP-2 carrier is of crucial importance. Although gelatin gels were capable of a sustained release of BMP-2 (Wang et al., 2013), this resulted in bone regeneration that mainly occurred around and inside the porous titanium implants (Fig. 8, a1). In addition, bone regeneration in that study did not lead to bridging of the grafted defects within twelve weeks (Fig. 8, b1). The mechanical strength also reached up to only 50 % of the original strength in that study (Wang et al., 2013). In contrast, physiological fibrin gels with low doses of BMP-2 boosted bone regeneration in the present study and completely bridged the majority of the grafted defects within four weeks (Fig. 8, b2), filling up more than 50 % of the porous volume of the titanium implants with...
regenerated bone. Twelve weeks following the surgery, the grafted femora were already more than twice as strong as their original strength (i.e. control femora Fig. 6).

**Conclusion**

This study reports the development of a new biomaterial combination, capable of stimulating complete bone regeneration in load-bearing segmental defects in rat femora. This optimal combination enabled quick bone regeneration within four weeks and full restoration of the original bone functionality and anatomical shape in this pre-clinical model. Since all used methods have been used separately in trauma and orthopaedic surgery, this combination should be evaluated in a large animal model or a clinical trial and this might result in an efficient bone graft substitute to graft load-bearing segmental bone defects in trauma and orthopaedic surgery.

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**References**


Bone formation would be impaired if the implants will lose some of their initial strength over time (Amin Yavari et al, 2013b) Treatment of nonunions with nonglycosylated recombinant human bone morphogenetic protein-2 delivered from a fibrin matrix. Vet Surg 33: 112-118.


Discussion with Reviewers

Reviewer I: How can such an implant be fixed? In addition, can the fixation be integral part of the construct?

Authors: This is an intriguing question and obviously an important aspect of implant development. Because porous titanium implants are developed using selective laser melting (an additive manufacturing technique) options for screw or pin fixation can be completely integrated into the implant design. Examples of porous implants designs are more thoroughly described by Murr (Murr et al, 2010).

Reviewer II: When you look more than 12 weeks, may it be possible that the titanium mesh is impairing further bone formation as it is not degrading?

Authors: Bone formation would be impaired if implantation of the porous titanium implants result in a significant mechanical mismatch with the surrounding bone. This could cause stress shielding and subsequent bone resorption. However, extensive mechanical testing of the produced porous titanium indicated that due to fatigue the implants will lose some of their initial strength over time (Amin Yavari et al., 2013). This gradual weakening of the implants will avoid stress-shielding and is more likely...
to stimulate bone formation rather than impairing bone formation. The main disadvantage of the fact that titanium implants do not degrade is the long-term risk of infection.

Reviewer III: Is this laboriously produced fibrin much better than commercially available material?

Authors: That may depend on the application. We have already shown that HMW-Fb stimulates angiogenesis in vitro (Kaijzel et al., 2006). As our commercial UNF-Fb does not behave much differently from HMW-Fb, with respect to bone bridging, we cannot clearly answer this for the present study. However, our novel combination of a fibrin gel within a highly porous titanium implant results in very good bone regeneration – at least when using HMW-Fb with low doses of BMP-2. The limitations of this study are now thoroughly addressed.