Partial Bone Formation in Additive Manufactured Porous Implants Reduces Predicted Stress and Danger of Fatigue Failure

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Abstract—New porous implant designs made possible by additive manufacturing allow for increased osseointegration, potentially improving implant performance and longevity for patients that require massive bone implants. The aim of this study was to evaluate how implantation and the strain distribution in the implant affect the pattern of bone ingrowth and how changes in tissue density within the pores alter the stresses in implants. The hypothesis was that porous metal implants are susceptible to fatigue failure, and that this reduces as osseointegration occurs. A phenomenological, finite element analysis (FEA) bone remodelling model was used to predict partial bone formation for two porous (pore sizes of 700 μm and 1500 μm), laser sintered Ti6Al4V implants in an ovine condylar defect model, and was compared and verified against in vivo, histology results. The FEA models predicted partial bone formation within the porous implants, but over-estimated the amount of bone-surface area compared to histology results. The stress and strain in the implant and adjacent tissues were assessed before, during bone remodelling, and at equilibrium. Results showed that partial bone formation improves the stress distribution locally by reducing stress concentrations for both pore sizes, by at least 20%. This improves the long-term fatigue resistance for the larger pore implant, as excessively high stress is reduced to safer levels (86% of fatigue strength) as bone forms. The stress distribution only changed slightly in regions without bone growth. As the extent of bone formation into extensively porous bone implants depends on the level of stress shielding, the design of the implant and stiffness have significant influence on bone integration and need to be considered carefully to ensure the safety of implants with substantial porous regions. To our knowledge this is the first time that the effect of bone formation on stress distribution within a porous implant has been described and characterised.

Keywords—Stress shielding, Rehabilitation, Bone remodelling, Laser sintered titanium alloy implant, Fatigue, Finite element analysis.

INTRODUCTION

The introduction of an implant changes the mechanical environment, resulting in bone adaptation. Bone resorption and aseptic loosening caused by stress shielding can reduce the implant longevity. 11,29,31 Retrospective clinical follow-up for bone cancer survivors demonstrated that promoting extracortical bone formation and osseointegration at the shoulder of distal femoral segmental prostheses using a grooved collar design is effective at reducing failure due to aseptic loosening and improves the survivorship of implants at 10 years from 75% to 98% 9 (Fig. 1). An FEA study to understand the effects of extracortical bone formation on stresses in the intramedullary stem found an 80% decrease in stress concentration due to bone growth, protecting the implant from failure at the stem-collar interface. 12

Recent advancements in additive manufacturing enable implant scaffold designs with extensive interface connectivity to enhance bone growth into porous structures, improving stability and fixation. 18,30,33 An in vivo study compared the outcomes of selective laser sintered (SLS) porous and machined grooved collars in segmental prostheses and found that porous designs
had higher osseointegration. An experimental study found that filling extensively porous implants with epoxy improved their mechanical properties by a factor of 2-7, demonstrating the importance of bone formation in additively manufactured porous implants. However, the long-term performance and potential risk of failure of additively manufactured implants is a concern with radiographic evidence of poor osseointegration in short- to mid-term follow-up, and in vivo evaluation of osseointegration is unable to evaluate safety performance as the test duration is short compared to the implant lifespan in humans. Specified standard mechanical tests are used to determine the fatigue performance of implants, however these tests fail to mimic the different patterns of bone ingrowth and integration. Bone formation in the porous structure reinforces the implant and enhances its performance, but incomplete bone formation could cause failure due to the partially fused, relatively weak porous material.

Computer simulations are useful for modelling adaptive changes caused by implant architecture, to assess the effects on stress and strain within the bone and implant, and to compare with the fatigue strength. FEA assessment of changes in the mechanical environment often assumes full bone formation with homogenous material properties. However, scanning electron microscopy (SEM) and histology results from animal studies have shown that bone formation and tissue regeneration occur only partially and with varying bone density. The assessment of long-term performance in terms of bone ingrowth and its effects on the structural integrity is important for porous metal implants as the structures have thin walls and irregular surfaces that can affect fatigue properties, particularly in notch sensitive materials such as titanium alloy. One method of modelling adaptive bone changes is through mechanotransduction algorithms, using mechanobiological or phenomenological approaches. The former is focussed on the short term, modelling the process by which mesenchymal stem cells (MSCs) differentiate to osteocytes and form bone. The latter is based on the idea of mechanostat, and is concerned with long-term effects at the tissue level. Advantages and limitations of both modelling approaches for orthopaedic implants have been compared. Mechanobiological models have predicted bone ingrowth on coated surfaces and porous implants, but as these scaffolds are non-metallic, fatigue behaviour is usually not evaluated. For metallic implants, bone remodelling algorithms to simulate long-term bone density changes are used to conduct stress analysis. Bone remodelling algorithms have also been coupled with a placeholder and an osteoconnectivity matrix to predict the extent of partial, inhomogeneous bone formation in a grooved titanium segmental prosthesis in a verified adaptive FEA model.

The aim of this study was to evaluate the effect of partial bone formation in two porous, laser sintered Ti6Al4V implants on the internal stress and strain, and the potential implications for implant safety and failure. This paper uses the dataset from a FEA model that was verified with histology results for predicting bone formation into porous implants. Its focus was the effect of pore size, coating, and material properties on bone formation, whilst the results reported here investigate the impact of bone formation on the implant. Stress changes due to inhomogeneous tissue densification in the bone and implant were investigated. As histology results were obtained only at one time point, the process of bone ingrowth, where tissue differentiates to bone and becomes rigidly fixed to the implant, was not considered. The changes in tissue density and stiffness with time were modelled using bone remodelling algorithms, considering only the contribution to load carrying capacity from lamellar bone that had formed, but not the smaller effect of immature soft tissue. Stress and strain in the implant and tissues before and after bone remodelling were compared, in relation to the strength and fatigue limit of the materials.

**FIGURE 1.** Distal femoral implant 12 years after insertion, showing grooved HA coated collar (C), extracortical bone formation (EB) and cemented intramedullary stem (IM).
MATERIALS AND METHODS

Animal Model

Porous implants (8 mm diameter, 14.5 mm length) with pore sizes of 700 µm and 1500 µm (Fig. 2A) were used as part of a larger study that evaluated the effect of the combination of pore size and different types of HA coatings on osseointegration. The implant design utilised a strut and plate structure, in a critical sized defect model. The 700 µm implant had struts 300 µm in diameter and 700 µm tall, while the 1500 µm implant had struts 750 µm in diameter and 1500 µm tall. The thickness of the plates corresponded to the diameter of the struts. The 700 µm and 1500 µm implants had porosities of 75% and 70% respectively. The two designs were manufactured as one cylindrical implant using SLS of Ti6Al4V. Bilateral 8 mm × 15 mm defects were created in the medial femoral condyles of 6 mature sheep, and the implants pushed into place to achieve a line-to-line fit in the cancellous bone (Fig. 3). The orientation and location (distal–proximal, left and right leg) of the implant was randomized. 30 cylinders were used (24 coated with calcium phosphate), but only results from the 6 uncoated implants (control group) were considered here as the FEA model cannot accurately predict differences between the different types of HA coating. As the uncoated implant had the least amount of bone formation, it represents the worst-case for failure analysis. Tissue and skin were closed, and all animals recovered well. Due to welfare considerations associated with the reduction of animals in experiments, only one time period was used. All procedures were conducted in accordance with the UK Animals (Scientific Procedures) Act 1986, under personal and project licences from the UK Home Office following review by the local animal welfare and ethical research committee.

After 6 weeks, the implants and surrounding bone were retrieved and fixed in formalin. The specimens were dehydrated and embedded in acrylic resin. Longitudinal thin sections approximately 80 µm thick were prepared by sectioning each specimen through the centre, using grinding and polishing techniques. Toluidine Blue and Paragon staining were used to identify soft tissue and bone within the implant.

FIGURE 2. (a) Implant design used for in vivo sheep study (units: mm); (b) FEA model for 700 µm pore size implant; (c) mesh at 10° cut from loading direction for 700 µm and 1500 µm implants.

FIGURE 3. Radiograph showing implant and defect positions within the femoral condyle.
stained purple and red respectively. Imaging of the stained slides was performed using light microscopy (Axioskop, Carl Zeiss, UK). Thresholding and free-form tools in ImageJ (v1.51, NIH, USA) were used to outline bone regions and quantify bone area ratio. Back scattered scanning electron microscopy (SEM) was used to examine bone structure (JEOL 3500C, Japan). Mann–Whitney U tests were conducted in Origin 2016 (OriginLab Corp., USA) to compare bone ingrowth.

**Finite Element Model**

Separate FEA models were developed for the 2 pore sizes (Fig. 2), modelling quarter slices using symmetric boundary conditions and repeating pattern to minimise computational costs as the implant orientation and location were not found to affect the results significantly. Both models had dimensions of 5 mm x 5 mm. Soft tissue was assumed to initially fill the implant pores. Homogenised trabecular bone was assumed to surround the exterior of the implant, as limited geometrical information on the trabecular bone was available. The models were axially restrained. The peak load through the medial condyles of the stifle joint during walking was applied. This is equivalent to loads of 89 N and 200 N in the 700 μm (1 mm width) and 1500 μm (2.25 mm width) models respectively, scaled according to the repeating pore and strut pattern width. Static loading with specified peak load has been demonstrated to capture the main changes in bone remodelling, comparable to the daily load history of the Stanford model. Meshes were generated in MSC.Marc 2017.0 (MSC Software Corporation, USA) using linear tetrahedral elements for soft tissue and linear hybrid 4-, 6- and 8-noded elements for bone and implant. This improved geometrical conformance while minimising the total number of elements. Linear elements were used for the soft tissues as mainly compressive load was transmitted through the condyles. The element size was small to capture the geometry around the implants, and the adaptive models were computationally expensive to run. Micromotion was assumed to be minimal and thus tied boundary conditions were used for all contact interfaces. Static stress analysis was conducted to investigate mesh convergence with criteria of 5% error at the fillet and 1% in the soft tissue and bone. The FEA models converged with edge lengths of 0.06 mm and 0.03 mm, yielding 1.33 million and 2.77 million elements for the 1500 μm and 700 μm implants respectively.

Isotropic homogenous material properties were assigned for trabecular bone and implant (Bone: elastic modulus \( E = 1.5 \) GPa, Poisson’s ratio \( \nu = 0.34; \) Ti6Al4V: \( E = 110 \) GPa, \( \nu = 0.34).\) Soft tissue was assigned an initial modulus \( E = 0.5 \) GPa, and Poisson’s ratio \( \nu = 0.3.\) Based on the adaptive elasticity theory, tissue density and thus elastic modulus were permitted to adapt according to the strain energy density per unit bone mass (SED) to model bone formation. The model enforces the sequential laying down of new bone by allowing only elements adjacent to bone, and with SED above the threshold, to adapt their density. Elements that did not fulfil these two conditions had no change in density. The physiological basis is that MSCs diffuse from the bone stock to the stimulus site to form new bone, as bone formation occurs from the edge of the implant towards the centre of the porous structure. In this model no bone formation occurred spontaneously within the centre of the implant. Both bone and soft tissue could exist at the initial modulus of 0.5GPa. The conditions imposed led to initial increases in modulus during ingrowth \( (E_i = 0.5 \text{ GPa}) \), with density decreases due to redistribution occurring only later (number of active elements constant). Change in density \( \rho \) was calculated from SED:

\[
d\rho = B (\text{SED} - k) d\tau \quad 0 < \rho \leq \rho_{cb}
\]

where \( B \) and \( k \) are remodelling rate and reference threshold, set at values of \( 1 \text{ (g cm}^{-3}\text{)}^2/\text{MPa-ctu} \) and 0.0044 \text{ J g}^{-1} \text{ respectively. The upper limit for bone formation } \rho_{cb}, \text{ corresponded to 12 GPa.}^{17,31} \text{ Density of surrounding trabecular bone was assumed to remain unchanged, as resorption of existing bone stock due to implant stress shielding is observed only during long-term follow-up.}\)

The stiffness matrix was updated in each time increment using an established density-modulus relationship from literature \( (E = 3790\rho^\frac{3}{2})^{17} \). Density-modulus relationships calibrated for CT images \(^7\) were not employed, as initially homogenous material properties were assigned. Fixed time step of 0.1 computer time unit (ctu) was used until the number of remodelling elements remained stable. For computational efficiency, adaptive time stepping was used thereafter until equilibrium, at 1.2 \times \text{ of the previous time step.}^7 \text{ Equilibrium was achieved when the change in average tissue density was less than 0.005%. Parametric analysis conducted previously showed that the choice of initial modulus did not affect equilibrium results, but that the initial time step has to be sufficiently small to prevent a numerical overshoot.}^7 \text{ Time units used should be considered arbitrary, as the histology results were only available at 6 weeks. There is insufficient literature for correlating ovine bone formation between simulation (ctu) and actual time, and the time correlation found previously for human subjects may...}
not be applicable for sheep. Quantification of bone area ratio followed the same procedure as histological analysis ("Animal model" section).

Failure Risk Assessment

The failure risks of implant and periprosthetic bone were evaluated using maximum nodal von Mises stress (implant) and largest principal strain (bone), compared with their respective limits. Bone failure strain was assumed to be 0.0305 with strength and fatigue limits of titanium alloys given in Table 1. Custom-written post-processing scripts in Python were used to identify nodes with the highest von Mises stress and principal strain and vertical path plots across the nodes were drawn.

RESULTS

Verification of FEA Model with In Vivo Data

Histology results showed regions of bone formation across the centre of longitudinal sections taken after 6 weeks (Fig. 4). No difference was observed between left/right leg, orientation (inside/outside) and location of the implants tested. A growth gradient was observed with the highest bone formation at the exterior, which reduced towards the centre. The percentage surface integration was 9.8 ± 5.0% and 10.7 ± 4.9% for the 1500 μm and 700 μm implant respectively, not significantly different (Mann–Whitney U test, $p = 0.810$). SEM images of bone formation within the pores showed that the bone is mature and lamellar in structure, indicating that the bone formed is unlikely to be a direct reaction to the surgery (Fig. 5). Regions of bone formation predicted by the FEA models at equilibrium qualitatively agreed with histology results as bone growth was restricted to the outer implant pores irrespective of pore size (Fig. 4). For the 1500 μm implant,

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**TABLE 1.** Mechanical properties of Ti6Al4V, machined and additively manufactured using electron beam melting (EBM). Fatigue strength at $10^7$ cycles $^{21,32,34}$

|               | Machined   | Additively manufactured |
|---------------|------------|-------------------------|
| Yield strength (MPa) | 903        | 882                     |
| Ultimate tensile strength (MPa) | 958        | 979                     |
| Fatigue strength (MPa)       | 510        | 350                     |

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**FIGURE 4.** Correspondence of bone formation in animal model histology (left) and FEA models (right) for both implants obtained at equilibrium: (a) 1500 μm pore size model. (b) 700 μm pore size model; histology: bone stained pink with toluidine blue; FEA: scale shows elastic modulus (MPa).

**FIGURE 5.** Back scattered scanning electron micrograph (SEM) of the bone-implant interface for a 700 μm implant, retrieved at 6 weeks after insertion.
numerical results predicted formation of dense bone in the outer pore, surrounded by the plates. For the 700 μm model, fully dense bone was predicted to form in the outer pore, with the strut blocking bone formation on its underside. There was bone formation originating from the top side of the second strut layer, to about 1/3 the height of the pore. For both implants, histology results showed bone formation only at the first strut and thus FEA models over-predicted bone growth. The bone surface ratio obtained from histomorphometric analysis of 6 implants (one from each sheep) were 0.125 ± 0.087 [range 0.041–0.288] and 0.134 ± 0.067 [range 0.021–0.222] for the 1500 μm and 700 μm implant respectively, not significantly different (Mann–Whitney U test, \( p = 0.575 \)). Predicted percentage bone surface areas of the 1500 μm and 700 μm implants were higher than the histological results at 0.270 ± 0.052 [range 0.196–0.374] and 0.175 ± 0.047 [0.088–0.224] respectively, which were significantly different for the two implants (Mann–Whitney U test, \( p = 0.001 \)).

**Changes to Implant Stress Distribution due to Bone Remodelling**

Stress patterns in both porous implants changed due to bone remodelling. Before remodelling, the highest stress concentrations were at the sharp edges where partial holes intersect the outer ring of material (Fig. 6b). High stresses were also present along the plates, near holes along the loading direction, and at strut fillets. At equilibrium, bone formation extended beyond the top layer of struts in both implants and partially filled the first row of holes on the underside. The predicted depth of bone growth in the 1500 μm model was higher than in the 700 μm implant, which was not observed experimentally (Fig. 6a).

There was an overall stress reduction in both implants with remodelling and bone formation. The highest stress concentration in the 700 μm implant reduced by 23% from 534 to 411 MPa, and by 21% from 383 to 301 MPa for the 1500 μm implant (Table 2). Before remodelling, peak von Mises stresses of both implant models were lower than the tensile strength of machined Ti6Al4V (Table 1), but above the fatigue strength of machined Ti6Al4V at 10^7 cycles for the 700 μm implant. The maximum von Mises stress exceeded the fatigue strength of untreated additively manufactured Ti6Al4V for both pore sizes (Tables 1, 2).

Remodelling changed the location of the highest stress concentration, initially at the edges of irregularly shaped outer holes, to the holes along the plates after remodelling (Fig. 6b/c arrows). This reduction corresponded to regions with bone remodelling, primarily at the geometrical shape change, at the top of the plate and around the fillet above the struts. Stress reduction was observed in the implant adjacent to regions of remodelled bone, but there was a slight increase in stress inside the partially remodelled bone (Fig. 7). There was no observed change in stress levels inside the deeper struts, beyond an arc length of 3 mm for the 700 μm implant.

As adjacent tissues remodelled, the high stresses at the partial holes (Fig. 6b arrows) decreased asymptotically with time. Most of the reduction occurred within 100 ctu (Fig. 8 dashed lines). At equilibrium, von Mises stresses of the 700 μm and 1500 μm implants at the partial holes reduced to 259 MPa and 232 MPa, below the fatigue failure strength of additively manufactured Ti6Al4V. At equilibrium, peak stresses were located at the middle of the first layer of holes along the plates, the thinnest cross-section (Fig. 6c arrows). Stresses at these holes increased slightly from 282 to 301 MPa, and 398 to 411 MPa in the 1500 μm and 700 μm implant respectively (Fig. 8 solid lines). The increase in stress was due to remodelling that occurred above the holes, increasing the stress uptake in the local region that had not remodelled (Fig. 7). The maximum stress in the 700 μm implant at equilibrium was 37% higher than that in 1500 μm model, as the 700 μm implant has a higher porosity (75%) compared to the 1500 μm model (70%) and thus thinner struts and plates.

**Strain Within Remodelled Tissue and Bone**

The highest absolute strains (minimum principal strain) before remodelling were at the circumference of the soft tissue in the loading direction, especially adjacent to the outer implant struts (Fig. 9a). The highest compressive strain for the 700 μm implant was 0.08, 10% higher than for the 1500 μm implant. Bone remodelling significantly reduced the compressive strain for both cases to less than 0.02 (Fig. 9b). For the 700 μm implant, there was a redistribution of the minimum principal strain in the tissues within the inner pores, with an increase in magnitude adjacent to the partial holes.

The peak strain magnitude in the bone stock before remodelling was 0.03, close to the fracture limit of ovine bone. Remodelling reduced the peak maximum

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**FIGURE 6.** Effect of bone formation on von Mises stress in the implant. (a) Elastic modulus of tissues showing the extent of bone remodelling; Implant not shown for improved visualisation; (b) stress in implant before remodelling; (c) stress in implant after remodelling, at equilibrium; Arrows indicate regions of maximum von Mises stress.
principal strain to less than 0.008. The highest maximum principal strain remained towards the middle of the arc at 45°/C176, but strain was redistributed across the full thickness of the bone stock, causing an increase in strain in the newly formed bone in the region between 30°/C176 and 60°/C176 from the vertical direction. However, these strains were not conducive to further bone formation deeper into the implant as the SED was below the threshold required for densification.

**DISCUSSION**

Extensively porous implants are being used in orthopaedics with considerable success.3,22,35 However, higher failure rates than expected have been reported5 for the Tritanium acetabular component. Hence, changes in loading on the implant, tissue and bone, as bone grows into the porous structure, have important implications for the design of extensively porous implants. This study used a SED-based algorithm, developed to model extracortical bone formation,7 to evaluate the stress and strain distribution within porous implants of two different pore sizes (700 μm and 1500 μm) due to partial bone formation. Porous Implants made by SLS are being used clinically for

**TABLE 2. Peak von Mises stress experienced by the implant before and after remodelling**

| Implant size (μm) | Peak von Mises stress (MPa) | Before remodelling | At equilibrium |
|-------------------|-----------------------------|-------------------|---------------|
| 1500              | 383                         | 301               |
| 700               | 534                         | 411               |

**FIGURE 7.** Stress plots along loading direction of the implant before and after remodelling (path across location of initial maximum nodal value). Cutting planes show extent of remodelling at equilibrium. (a) 1500 μm implant; (b) 700 μm implant.

**FIGURE 8.** Time evolution of von Mises stress for 1500 μm and 700 μm implants at locations of highest initial stress (Max initial, at partial hole, Fig. 6b arrows) and highest stress at equilibrium (Max eqm, thinnest strut cross-section. Figure 6c arrows).
applications to augment bone growth throughout the structure. The implant modulus depends on pore size and strut thickness, whilst the amount of bone ingrowth depends on implant modulus and loading conditions. The reliance of bone ingrowth on the modulus of a porous material has been demonstrated, and from this study it is apparent that modulus mismatch between the implant and the surrounding bone may lead to regions of the porous structure without bone tissue. The novelty of our study is that we predict bone ingrowth and relate this to the fatigue performance of substantially porous implants. We believe that this is important as short-term failures of these structure have been identified. In our study there was no bone formation in the centre of the implant, and overall stress in the implants reduced up to the depth where bone formation occurred (Fig. 7). There was a corresponding reduction in minimum principal strain in the surrounding bone stock and an increased stress uptake in the newly formed bone, as its elastic modulus increases. This agreed with previous in vitro and FEA studies that demonstrated that bone formation within porous implants improves the implant strength, yield stress and fatigue resistance, or reduces the risk of fracture through stress redistribution. However, the papers that investigated metallic implants considered homogenous bone at discrete stages of growth, whereas histology results
from in vivo studies showed that bone formation is inhomogeneous, as porous structures are incompletely filled.10,20,22

The results showed that the highest initial implant stress concentration was located at irregular geometrical features, and the stress pattern in the plates in Fig. 6 was due to the narrowing of the cross section. The initial peak von Mises stress reduced quickly (within 100 ctu), and by more than 20% in both implant designs, as bone grew into the implant. This reduction is especially important for the 700 μm implant, as the initial stress exceeded the fatigue strength of Ti6Al4V. However, the predicted equilibrium von Mises stresses in the FEA models, 301 MPa and 411 MPa for the 1500 μm and 700 μm implants respectively, are still not within safe limits of untreated additively manufactured Ti6Al4V (fatigue strength: 350 MPa). As the peak stress at equilibrium was located at the holes, the results indicate that geometrical shape changes along the loading direction should be minimised where possible.

The as-designed geometries were used, but there could be geometrical discrepancies and worse fatigue performance due to current resolution limits of additive manufacturing.2,14,18 Titanium alloy is a notch sensitive material and both SLS and electron beam machining (EBM) lead to rougher surface finishing than conventional machining due to partially fused particles 27 that might increase stress concentration and lead to premature failure. It is difficult for surface finishing techniques to access the inner pores, and therefore the fatigue properties of these structures are likely lower than expected. Post-processing of titanium alloys through hot isostatic pressing (HIP) or heat treatment causes titanium alloy phase transformation, resulting in an increase in β-phase titanium that can increase the fatigue strength of Ti6Al4V by up to 40%.15,27,34 SLS and EBM may also lead to structures with lower strength due to increased porosity, hence HIP or other heat treatment is advised for alloy structures made using 3D printing. In this study, the porous structures were modelled as smooth, probably overestimating the fatigue strength. Future work should include conducting micro-CT scans of manufactured implants with metal artefact suppression, and using the reconstructed scans to improve the geometrical fidelity of the FEA models.37 The FE models did not take into account surface microgeometry or changes in the structure of the alloy such as grain size or phase changes. We expect that these changes would only reduce the fatigue strength of the porous material.

Bone remodelling was highest at regions of highest curvature, which agreed with the result obtained using a mechanobiological algorithm to model bone ingrowth.3 One of the goals in porous implant design is to induce bone formation throughout the entire implant, but this remains a challenge in orthopaedics as implant design is often conducted empirically, or to match the average surrounding bone mechanical properties.2,14,18,24,33 Path plots taken across the FEA models indicate that lack of internal bone formation seen in vivo is the result of stress shielding by the implant structure, reducing SED in the tissue to less than 3 MPa (Fig. 7). Thus, the effectiveness of partial bone formation in reducing stress concentration in the implant depends on the location and depth of bone remodelling. Therefore, a trade-off needs to be considered, as sufficiently high stress and strain in the surrounding tissue is required to enhance bone formation, but increases the risk of implant fatigue failure in the absence of bone remodelling.

Bone remodelling improves the mechanical interlock, beneficial for improved load transfer to existing bone stock to prevent long-term resorption. These results highlight the importance of implant design to optimise bone formation. Bone remodelling can reduce stress to be lower than regions unaffected by bone remodelling (Fig. 8), suggesting that implant architecture and stiffness are important since they can affect adaptive changes.2,18,26 Previous work that investigated the effect of reducing the apparent modulus mismatch showed that increasing porosity often increases bone formation.3,26 In particular, bone formation of up to 57% has been reported when Octet truss lattice structures were implanted in canine diaphysis.2 Another approach is to use a different bio-compatible, low modulus, high strength titanium alloy. FEA simulations for a condylar defect model predicted that the use of Titanium-Tantalum alloy (E = 67 GPa) instead of Ti6Al4V (E = 110 GPa) would increase the volume of bone formation from 34 to 65%.8 However, the actual value for in vivo studies may be lower as the model over-predicted the amount of bone formation and these alloys are currently not widely used for SLS.

Large strains were observed in the thin layer of soft tissue between the implant and bone, and at the bone-implant interfaces before remodelling (Fig. 9). This is partially due to the modelled tied contact conditions; the assumption of friction interfaces could be better suited to model the press-fitted implants. Tied contact conditions were used as osseointegration was assumed, as Ti6Al4V is biocompatible and previous histology result showed that bone adjacent to the implant had osseointegrated.9,22 The initial values of −0.08 and 0.04 for minimum and maximum principal strain exceed the 0.03 magnitude limit for ovine bone fracture. After remodelling, these values reduced to safe levels of −0.02 and 0.01 respectively.23 The results suggest that care needs to be taken in the design of implants (e.g.
placement of struts) to avoid thin layers of soft tissue experiencing high strains.

Implants were modelled as quarter slices, not considering potential end effects of the cylindrical implant and its location within the condyle on the stress distribution, as histological results did not show statistically significant differences in bone formation across the implant or its orientation and location. Peak load in medial condyles was applied in the FEA, as full weight-bearing immediately post-implantation was assumed. A uniform distributed load was assumed based on the weight of the animals used, a common technique. These sources of inaccuracies could explain the over-prediction of bone area ratio in the FEA model. Conducting full gait analysis with instrumented implants would increase the model fidelity. However, the use of a constant peak load represents the worst-case scenario for failure analysis, and it is commonly accepted in literature that bone tissue adapts to the loading could slow rehabilitation. As patient weight differs, the rehabilitation loading scheme should be patient-specific, with a gradual increase in loading to minimise both healing time and risk of implant failure. This study employed histological analysis (which is destructive), an important requirement for progressing towards clinical trials, to quantify the extent of bone ingrowth. Future work should consider larger studies with multiple time points, the inclusion of both non-destructive evaluation and testing of implants to failure, imaged using micro-CT, to further validate the FEA models and evaluate implant performance. A clinical tool could be developed to monitor bone regeneration progress, representing the elastic modulus as a percentage change to the pre-operative value, and the implant stresses as the ratio to material strength.

The effect of partial, inhomogeneous bone formation into the pores of additively manufactured implants was studied. An FEA algorithm previously developed to model bone formation was used to predict the effect of bone remodelling on the stress in the bone-implant structure. Results showed that bone remodelling protects the implant, reducing maximum von Mises stress by more than 20%. The maximum implant stress is still not within safe fatigue limits of additively manufactured Ti₆Al₄V and further improvements to implant design are suggested. Initial rehabilitation should be carefully implemented to load the structure within safe limits, while providing sufficient stimuli for bone formation to occur and to protect against implant failure. The use of extensively porous implants in load bearing situations should proceed with caution as the implant fatigue performance will be determined by the level of bone growth. Regions with only partial bone formation in the porous structure due to inappropriate structural stiffness may be at risk of fatigue failure.
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CONFLICT OF INTEREST

The authors have no conflict of interest to declare.

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REFERENCES

1. Aldieri, A., M. Terzini, G. Osella, A. M. Priola, A. Angeli, A. Veltri, A. L. Audenino, and C. Bignardi. Osteoporotic hip fracture prediction: is t-score-based criterion enough? A hip structural analysis-based model. J. Biomech. Eng. 140:111004, 2018.
2. Arabnejad, S., B. R. Johnston, J. A. Pura, B. Singh, M. Tanzer, and D. Pasini. High-strength porous biomaterials for bone replacement: a strategy to assess the interplay between cell morphology, mechanical properties, bone ingrowth and manufacturing constraints. Acta Biomater. 30:345–356, 2016.
3. Bandyopadhyay, A., A. Shivaram, S. Tarafder, H. Sahasrabudhe, D. Banerjee, and S. Bose. In vivo response of laser processed porous titanium implants for load-bearing implants. Ann. Biomed. Eng. 45:249–260, 2017.
4. Boccaccio, A., A. E. Uva, M. Fiorentino, L. Lamberti, and G. Monno. A mechanobiology-based algorithm to optimize the microstructure geometry of bone tissue scaffolds. Int. J. Biol. Sci. 12:1–17, 2016.
5. Byrne, D. P., D. Lacroix, J. A. Planell, D. J. Kelly, and P. J. Prendergast. Simulation of tissue differentiation in a scaffold as a function of porosity, Young’s modulus and dissolution rate: application of mechanobiological models in tissue engineering. Biomaterials 28:5544–5554, 2008.
6. Carlh, A. V., L. C. Warth, K. L. de Mesy Bentley, and B. J. Nestor. Short to midterm follow-up of the tritanium primary acetabular component: a cause for concern. J. Arthroplast. 32:463–469, 2017.
7. Cheong, V. S., G. W. Blunn, M. J. Coathup, and P. Fromme. Adaptive 3D finite element analysis model to simulate extracortical bone growth. Comput. Methods Biomech. Biomed. Eng. 21:129–138, 2018.
8. Cheong, V. S., P. Fromme, A. Mumith, M. J. Coathup, and G. W. Blunn. Novel adaptive finite element algorithms to predict bone ingrowth in additive manufactured porous implants. J. Mech. Behav. Biomed. Mater. 87:230–239, 2018.
9. Coathup, M. J., V. Batta, R. C. Pollock, W. J. Aston, S. R. Cannon, J. A. Skinner, T. Briggs, P. S. Unwin, and G. W. Blunn. Long-term survival of cemented distal femoral endoprostheses with a hydroxyapatite-coated collar: a histological study and a radiographic follow-up. J. Bone Jt. Surg. Am. 95:1569–1575, 2013.
10. de Wild, M., S. Zimmermann, J. Rüegg, R. Schumacher, T. Fleischmann, C. Ghayor, and F. E. Weber. Influence of microarchitecture on osteoconduction and mechanics of porous titanium scaffolds generated by selective laser melting. 3D Print. Addit. Manuf. 3:142–151, 2016.
11. Dickinson, A., A. Taylor, and M. Browne. Implant–bone interface healing and adaptation in resurfacing hip replacement. Comput. Methods Biomech. Biomed. Eng. 15:935–947, 2012.
12. Fromme, P., G. W. Blunn, W. J. Aston, T. Abdoola, J. Koris, and M. J. Coathup. The effect of bone growth onto massive prostheses collars in protecting the implant from fracture. Med. Eng. Phys. 41:19–25, 2017.
13. Goodship, A., and J. Kenwright. The influence of induced micromovement upon the healing of experimental tibial fractures. J Bone Joint Surg Br. 67:650–655, 1985.
14. Hedayati, R., S. Janbaz, M. Sadighi, M. Mohammad-Aghdam, and A. A. Zadpoor. How does tissue regeneration influence the mechanical behavior of additively manufactured porous biomaterials? J. Mech. Behav. Biomed. Mater. 65:831–841, 2017.
15. Hrabe, N., T. Gnäupel-Herold, and T. Quinn. Fatigue properties of a titanium alloy (Ti–6Al–4V) fabricated via electron beam melting (EBM): effects of internal defects and residual stress. Int. J. Fatigue 94:202–210, 2017.
16. Huiskes, R. If bone is the answer, then what is the question? J. Anat. 197:145–156, 2000.
17. Huiskes, R., H. Weina, H. Grootenboer, M. Dalstra, B. Fudala, and T. Slooff. Adaptive bone-remodeling theory applied to prosthetic-design analysis. J. Biomech. 20:1135–1150, 1987.
18. Kang, H., J. P. Long, G. D. U. Goldner, S. A. Goldstein, and S. J. Hollister. A paradigm for the development and evaluation of novel implant topologies for bone fixation: implant design and fabrication. J. Biomech. 45:2241–2247, 2012.
19. Liu, X., and G. L. Niebur. Bone ingrowth into a porous coated implant predicted by a mechano-regulatory tissue differentiation algorithm. Biomach. Model Mechanobiol. 7:335–344, 2008.
20. Long, J. P., S. J. Hollister, and S. A. Goldstein. A paradigm for the development and evaluation of novel implant topologies for bone fixation: in vivo evaluation. J. Biomech. 45:2651–2657, 2012.
21. Mohammadhosseini, A., D. Fraser, S. H. Masood, and M. Jahedi. Microstructure and mechanical properties of Ti–6Al–4V manufactured by electron beam melting process. Mater. Res. Innov. 17:s106–s112, 2013.
22. Mumith, A., M. J. Coathup, M. Chimutengwende-Gordon, W. Aston, T. Briggs, and G. W. Blunn. Augmenting the osseointegration of endoprostheses using laser-sintered porous collars: an in vivo study. Bone Joint J. 99-B:276–282, 2017.
23. Nafie, A., C. C. Danielsen, F. Linde, and I. Hvid. Properties of growing trabecular ovine bone: Part I: mechanical and physical properties. Bone Joint J. 82:910–920, 2000.
Reznikov, N., O. R. Boughton, S. Ghouse, A. E. Weston, L. Collinson, G. W. Blunn, J. R. T. Jeffers, J. P. Cobb, and M. M. Stevens. Individual response variations in scaffold-guided bone regeneration are determined by independent strain- and injury-induced mechanisms. Biomaterials 194:183–194, 2019.

Sanz-Herrera, J. A., J. M. García-Aznar, and M. Doblare. Micro–macro numerical modelling of bone regeneration in tissue engineering. Comput. Methods Appl. Mech. Eng. 197:3092–3107, 2008.

Schouman, T., M. Schmitt, C. Adam, G. Dubois, and P. Rouch. Influence of the overall stiffness of a load-bearing porous titanium implant on bone ingrowth in critical-size mandibular bone defects in sheep. J. Mech. Behav. Biomed. Mater. 59:484–496, 2016.

Sing, S. L., W. Y. Yeong, and F. E. Wiria. Selective laser melting of titanium alloy with 50 wt% tantalum: Microstructure and mechanical properties. J. Alloys Compd. 660:461–470, 2016.

Taylor, W. R., B. M. Poepplau, C. König, R. M. Ehrig, S. Zachow, G. N. Duda, and M. O. Heller. The medial-lateral force distribution in the ovine stifle joint during walking. J. Orthop. Res. 29:567–571, 2011.

Taylor, M., and P. J. Prendergast. Four decades of finite element analysis of orthopaedic devices: where are we now and what are the opportunities? J. Biomech. 48:767–778, 2015.

Thesleff, A., R. Branemark, B. Hakansson, and M. Ortiz-Catalan. Biomechanical characterisation of bone-anchored implant systems for amputation limb prostheses: a systematic review. Ann. Biomed. Eng. 46:377–391, 2018.