Influence of Degrading Calcium Phosphate on the Remodelling and Mineralisation of Avascular Osseous Tissue in a Rat Calvaria Model

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Abstract: Artificial bone grafting is the subject of intensive investigation for craniofacial reconstruction. Among the synthetic candidates, those incorporating Calcium Phosphates (CaP) have gained popularity, due to their osteoinductive properties. Despite many quantitative studies on the degradation rates of CaP biomaterials in vitro, there is lack of quantitative studies in relation to bone remodelling kinetics in vivo. In this investigation, we tested implants of degradable CaP/chitosan-based biomaterial in a rat calvaria model of critical-sized defects. We observed remarkable acceleration of bone growth in the initial stages (~0.8 g/day), which was markedly higher than normal bone growth rates (~0.1 g/day) in implant-free controls. The implanted CaP-based biomaterial significantly enhanced bone formation, with woven bone seen as early as 20 days and lamellar bone at 40 and 60 days post-implantation. However the mineralisation of cortical bone was delayed, due to the overly rapid degradation of the biomaterial: This therapeutically important issue has never been identified before to our best knowledge. A theoretical analysis revealed that during degradation CaP can be rapidly released from the new bone matrix. Hence, while the degradable CaP biomaterial was found to be highly osteoconductive in vivo, future modifications would seek to optimize degradation efficiency for more sustained release of mineral ions, to enhance bone mineralisation at later stages of the healing process after implantation.

Keywords: Dental Graft, Calcium Phosphate, Rat Calvaria, Bone Growth Rate, Mineralisation

Introduction

The loss of teeth and supportive tissues in the jawbone often causes severe bone resorption, which hinders the healing process for dental implants, as well as recovery of physiological function (e.g., chewing, speech) and cosmetic appearance (Vignoletti et al., 2012). Current approaches in bone grafting for dental implants are dominated by autologous grafts and by allografts; however biological bone grafts have several shortcomings. These include donor tissue shortage and morbidity in autologous grafting and immune rejection or transmission of pathogens (e.g., HIV) in allografting (Burg et al., 2000). Over the past decade, artificial bone grafts have been investigated using various synthetic materials, including Hydroxyapatite (HA), Calcium Phosphates (CaP), polyesters, chitosan and their composites (Chen et al., 2012). Among these biomaterials, those containing CaP have been extensively tested (Bohner et al., 2012; Chai et al., 2012) because of their osteoinductive properties (Geffre et al., 2010; Thormann et al., 2013).

Osteoinduction refers to the heterotopic or ectopic formation of new bone at a wound site, due to the biological actions of growth factors (e.g., Bone Morphogenetic Proteins (BMPs) and bone minerals (e.g., CaP, Sr, F) (Habibovic and de Groot, 2007; Barradas et al., 2012). In addition, specific physico-chemical properties of the biomaterial, such as particle size (Ignjatovic et al., 2010), surface area, crystallinity, porosity and composition are also able influence bone formation (Lu et al., 2002; Lange et al., 2011). It has been reported that particle sizes ranging from 80 to 300 µm in diameter
can induce ectopic bone formation, whereas particles greater than 500 µm are not osteoinductive (Balaguer et al., 2010). In vivo studies also indicate that a specific surface area above a threshold level of 1.0 m²/g is critical for CaP to induce osteogenesis, suggesting the importance of optimizing surface areas of bone scaffolds (Chai et al., 2012). It is likely that the influences of particle size and specific surface area on bone formation are mediated via their effects on the degradation rate of CaP materials.

Biphasic materials containing mixtures of HA with CaP demonstrate simultaneously increased matrix solubility and osteoinductivity (Yuan et al., 2006), whereas pure HA is too stable and amorphous Tri-Calcium Phosphate (TCP) is too easily solubilised to be appreciably osteoinductive (Arinzeh et al., 2005). Based on this, we envisage that the osteoinductivity of CaP-based biomaterials is directly controlled by their degradation rates. Other materials properties (e.g., particle size, surface area, crystallinity, porosity and composition) influence osteoinductivity indirectly via degradation rates. Hence, degradation kinetics of biomaterials is a common and important factor that can affect new bone formation.

Despite many quantitative studies on the degradation rates of CaP biomaterials in vitro (Bohner et al., 2012; Chai et al., 2012; Barradas et al., 2012), there is lack of similar studies in relation to bone remodelling kinetics in vivo. Therefore, a primary objective of this study was to quantitatively investigate the influence of degrading CaP-based biomaterials on the remodelling of a vascular osseous tissue. To this end, we used the rat calvaria model because of its lack of intra-bone vascular networks (Pannarale et al., 1997). Given that dentin is an avascular tissue, this model particularly suits the evaluation of dental biomaterial in vivo.

Materials and Methods

Biomaterial Preparation

The biomaterial used in this study was a paste made from solid bioceramic powder and an aqueous chitosan solution (2 wt%, pH = 4.5), which was purchased from Polimar Ciencias Nutricio S.A. Chitosan is another biomaterial being intensively investigated for its ability to enhance osteoinductivity of CaP-based materials (Kim et al., 2008). For these reasons, the biomaterial used in this study is a composite of β-Tricalcium Phosphate (β-TCP) and chitosan in its acetylated form.

The bioceramic powder was amixture of β-TCP (Emprove®), calcium oxide and zinc oxide, all purchased from Merck®. The particle size of each powder was measured by laser granulometry using a Mastersizer 2000 (Malvern Instruments). The analysis was conducted using a laser diffraction liquid method on the following suspensions: β-TCP dispersed in propanol and CaO and ZnO dispersed in water. The particle diameters of the β-TCP, CaO and ZnO powders were 15, 3 and 9 µm, respectively. CaO was incorporated mainly for adjustment of pH of the composite and the minor addition of ZnO aimed at bone regeneration at early stages, as indicated by increased markers of osteoblast differentiation, matrix maturation and bone mineralisation in a previous work (Fielding and Bose, 2013).

A series of composites (100 samples) was systematically prepared from bioceramic powders of different β-TCP: CaO: ZnO ratios and mixed with the liquid chitosan solution at various solid/(solid+liquid) percentages. The pH values of these composites were measured. The composites (20 samples) with a pH value between 6.5 and 8.5 were considered provisionally to be safe for biological environments and were evaluated in vitro and the rest (80 samples) were discarded because of anticipated toxicity. The best formula in terms of cytocompatibility (section 3.1) in vitro is listed in Table 1, which had a pH value of ~7.5 and thus was subsequently chosen for the rat wound correction study.

The composite mixture was prepared as follows. The three ceramic powders were gently mechanically mixed for ~5 min and then the mixture was dried in a microwave oven for ~20 min. Each of these dried mixtures was added to the chitosan solution according to the designed percentage (Table 1) to produce a paste. The compressive strength of the paste was ~0.5 MPa. The crystallinity of β-TCP was approximately 30%, as provided by the supplier. The CaO and ZnO powders were amorphous.

Evaluation of Cytocompatibility in Vitro

The biocompatibility of materials in vitro were tested by exposure of proliferating cells to the extract media conditioned by material samples as they degrade, according to the standard cytotoxicity assessment study set by the International Standardization Organization (ISO 10993). In this method, extracts were obtained by soaking the test and control materials in separate cell culture media (DMEM supplemented with 10% Fetal Calf Serum (FCS), 1% L-glutamine and 0.5% penicillin/streptomycin) under conditions of 0.2 g mL⁻¹ of culture medium for 24h at 37°C/5% CO₂ in an incubator. HA was used as a positive control and tissue culture medium (DMEM supplemented with 10% Foetal Calf Serum (FCS), 1% L-glutamine and 0.5% penicillin/streptomycin) alone was used as the negative control. All material samples were sterilized using 70% alcohol/de-ionized water solution. Osteoblast-like MG63 cells from the American Type Culture Collection (ATCC) were seeded in culture media at a density of approximately 2000 cells/well in 96-well tissue culture plates (BD Falcon®). When cells reached 70% confluence (in approximately 2 days), the medium in each well was entirely replaced with 0.2 mL of extract media. All cultures were then further incubated for two days.

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At the end of the incubation period, cytotoxicity was quantified using a commercial kit, Tox-7 (Sigma-Aldrich). Spent culture media were collected and the degree of cell death was determined by measurement of Lactate Dehydrogenase (LDH) levels in the culture media (‘DEAD LDH’), as described previously (Liang et al., 2010). Each well containing living cells was then filled with 0.2 mL fresh cell culture medium and cells were treated with Tox-7 lysis solution. These lysates were then used to determine the cellular LDH content, which equates to the number of living cells per well (“LIVE LDH”). LDH level was determined by measuring the absorbance of the supernatant from the centrifuged medium at 490 nm (after subtraction for background absorbance at 690 nm) using a multi well plate format UV-vis spectrophotometer (Thermo Scientific). The absorbance results of LDH can be converted to the number of cells according to a linear standard curve. Hence, cytotoxicity can be expressed as follows Equation 1:

\[ \text{Percentage of dead cells} = \frac{\text{DEAD LDH}}{\text{DEAD LDH} + \text{LIVE LDH}} \times 100 \]  

Animal Model

Various animal models of bone repair with the anatomical capacity to recreate a Critical-Sized Defect (CSD) have been developed for biomaterials research (Kochi et al., 2009; Gosain et al., 2000), but the 5-mm rat calvarial bone defect is one of the most frequently used for in vivo studies (Mills, L.A. and A.H. Simpson, 2012; Develioglu et al., 2009; Jones et al., 2007). Very recently, a systematic review on calvarial CSD (CSD) have been developed for biomaterials research studies (Mills, L.A. and A.H. Simpson, 2000), but the 5-mm rat calvarial bone defect is one of the most frequently used for in vivo studies (Mills, L.A. and A.H. Simpson, 2012; Develioglu et al., 2009; Jones et al., 2007). Very recently, a systematic review on calvarial CSD (CSD) have been developed for biomaterials research studies (Mills, L.A. and A.H. Simpson, 2000), but the 5-mm rat calvarial bone defect is one of the most frequently used for in vivo studies (Mills, L.A. and A.H. Simpson, 2012; Develioglu et al., 2009; Jones et al., 2007). Very recently, a systematic review on calvarial CSD (CSD) have been developed for biomaterials research

Surgical Procedure

Operations were performed on the rats using general anaesthesia, i.e., ketamine (50 mg mL\(^{-1}\); 0.7 mg kg\(^{-1}\)), xilacin (2%, 0.6 mg kg\(^{-1}\)) and acepromazine maleate (10 mg mL\(^{-1}\), 0.6 mg kg\(^{-1}\)). Two circular bone defects were introduced using a trephine bur with a dental implant surgical handpiece (400 rpm). After washing the rat skull with a physiological saline solution, the right side skull bone defect was filled with chitosan/ceramic paste and the left one was left empty as a control (Fig. 1b). The scalp wounds were closed by suture. The rats were sacrificed at 20, 40 and 60 days post-implantation. To minimise experimental variability, the same surgeon performed all procedures.

Histology and Electron Microscopy Sample Preparation

After animals were euthanized, the experimental bone defects were retrieved and fixed in 2.5% glutaraldehyde Tris Buffered Saline (TBS) (0.01M, pH 7.4) solution for 48 h at 4°C. After washing with TBS (0.01M, pH 7.4), control samples and two samples of each experimental group were decalcified in 15% Ethylenediaminetetraacetic Acid (EDTA) water solution (pH 7.3) at 4°C for 2 months and the other two samples of each test group remained undecalcified. After rinsing in TBS (0.01M, pH 7.4), the bulk decalcified and undecalcified samples were dehydrated in increasing ethanol concentrations (70-100 %) and embedded in paraffin wax for optical microscope histology or Poly (Methyl Methacrylate) (PMMA) for transmission electron microscopy (Bateman et al., 2012). Then, 5-µm thick slides for histology or 50-70 nm TEM were sectioned along the coronal plane (Fig. 1b) in cooling water with a microtome or an ultrathin microtome, respectively. Histological slides of decalcified bone were stained with Haematoxylin-Eosin (H&E), whereas those of undecalcified bone were stained with Goldner's (Masseon) trichrome.

Histomorphometric Analysis

The above stained samples were imaged with an Aperio Scan Cope® Turbo scanner (Aperio Technologies, Vista, CA, USA/Serial Number AT1681). All scans were conducted at the same resolution and magnification (i.e., 0.497 microns per pixel, 20×). The images of the region of interest were then processed with Adobe Photoshop® version CS2 (9.0) to obtain the masks. In this process, the features of interest were designated in black and the rest of the examined area was designated in white and thus, a black-and-white image (i.e., mask) was created. Image J® 1.46 r software was used to measure the area percentage (Ar. %) of the selected features in the examination field (Egan et al., 2012).

Table 1. Formula of biomaterials used in the work

| Bioceramic powder (wt. %) | Composite (wt. %) |
|--------------------------|------------------|
| β-TCP 96 | CaO 2.5 | ZnO 1.5 | Bioceramic powder/composite 40 |
Fig. 1. (a) Surgical sites indicated on the exposed rat calvaria and (b) diagram of the coronal view of Experimental Site (ES-left) and Control Site (CS-right) which, with surrounding host bone, were retrieved at the end of the experimental period. The histological samples were sectioned across the middle region of each sample along the coronal plane.

**TEM Observation**

The TEM foils were examined with a Tecnai 20 microscope, with soft tissue regions examined at 100kV and bone tissue regions at 200 kV.

**Statistical Analysis**

In vitro experiments were performed with six samples per experimental group and in vivo experiments were performed with 4 animals per experimental group. The statistical outputs are shown in the form of a mean with Standard Error (±SE). A one-way analysis of variance (Mikhailenko et al., 2013) with Tukey’s post hoc test was performed to analyse the significant differences and the significance levels were set at a p-value of less than 0.05.

**Results**

**In vitro Evaluation of Cytocompatibility**

Cellular toxicity was visible in the cultures containing extracts of CaP-chitosan pastes of pH 6.8 and 7.2 (Fig. 2c and d), while the media containing the extracts of the pastes of pH 7.5 and 7.8 were found to support proliferation of MG63 cells (Fig. 2e and f). Quantitative LDH measurements confirmed that the cytocompatibility of the composite paste of pH 7.5 was similar to both culture media and HA controls, with no significant differences detected in cell death percentage between the optimised paste and control groups (Fig. 3). Hence, the subsequent in vivo studied were conducted using the paste of pH 7.5.

**Histological Analysis**

All surgical procedures were performed without complications. Histological examination (Goldner’s Messon trichrome) revealed that the control defects remained empty, with little new bone formed up to 40 days post-implantation and the amount of new bone was considerable only in the samples of the 60-day treatment group (~20 Ar. % of the defects) (Fig. 4). These results are in agreement with previous work on bilateral calvarial defects of 5 mm diameter, which reported that the area percentage of new bone formed in untreated control defects was approximately 20 and 23% at 2 and 3 months, respectively (Vajgel et al., 2013).

In contrast, the formation of new bone was more extensive in all the experimental sites implanted with the CaP-based biomaterial, with defects being filled with new soft and/or hard tissues (Fig. 5). In the 20-day tissue samples (Fig. 5a), the central region of the defect was occupied by a mixture of biomaterial particles and soft fibrous tissue. This was defined as Region I. Newly formed bone was defined as Region II (Fig. 5a). By 40 days after implantation, the defects were predominately occupied by new bone tissue, with bone marrow scattered in some discrete areas in the intermediate zone of the defects (Fig. 5b). After 40 days, no significant change was observed histologically (Fig. 5c).

The area percentages of Region I, II and bone marrow are given in Fig. 6. Region I was about 50% by area at 20 days but nearly completely replaced by new bone by day 40 (Fig. 6). Assuming the bone was growing at a reasonably constant rate, the average growth rate of the bone is thus estimated to be 2.5 mm/40 days = 62.5 μm day\(^{-1}\) ≈ 1.0×10\(^{-3}\) μm/s in thickness. This rate is remarkably higher than the growth rates of bone remodelling in rats of the same age (Fig. 7). It is estimated by extrapolation of the data in Fig. 7, which was retrieved from literature (Hansson et al., 1972), that the bone growth rate in rats of 4-months old could be as low as several μm/day, which has also been reported by another study (Raman, 1969).

An alternative way to quantify growth rate is weight per day. The density of bone is ~2 g/cm³. The weight of a 5 mm-diameter, 0.8 mm-thickness bone defect is ~0.03 g.
Fig. 2. Images of MG63 cells after one day culture in extract media, which were prepared by soaking test materials in tissue culture medium for 24h (a) Negative control (culture medium only) and (b) positive control of hydroxyapatite samples CaP-chitosan composite of pH (c) CaP-Ch_pH = 6.8, (d) CaP-Ch_pH = 7.2, (e) CaP-Ch_pH = 7.5 and (f) CaP-Ch_pH = 7.8 Magnification is the same for the six images.

Fig. 3. Cytotoxicity of the CaP-chitosan composites determined by measuring the release of LDH after two days of culture. The positive and negative control group was HA and standard culture medium, respectively. The percentage of dead cells in CaP-chitosan composite of pH 6.8 and 7.2 were significantly elevated, but at pH 7.5 was not significantly different from either the negative control or positive control groups ($p>0.05$). The cell death percentage in the CaP-chitosan of pH 7.8 was significantly different from that of the positive control, but not with that of the negative control.
Fig. 4. Low power view of sections of the undecalcified defect tissue from rats that had no implantation of biomaterial particles taken (a) 40 (H&E stained) and (b) 60 (Golders’s Masson trichrome stained) days post-surgery. Almost no new bone formation was observed in the defects retrieved at day 20 and 40 post surgery. After 60 days, new bone was observed, occupying approximately 20 Ar. % of the defect tissue. HB stands for host bone.

Fig. 5. Low power images of H&E stained sections of decalcified CaP biomaterial-implanted tissue (a) 20, (b) 40 and (c) 60 days after implantation. Tissues in Region I consisted mainly of fibrous tissues and CaP particles. Region II consisted of newly formed bone.
Assuming equivalent bone density, the bone growth rate of CaP-grafted defects is thus estimated to be 0.03 g/40 days ≈ 0.8 g/day and for control defects estimated as 0.03 g x 20%/60 days = 0.1 g/day. Again the growth rate of implanted defects was significantly higher than the normal remodelling in rats of the same age.

High power imaging of Region I revealed the disaggregated biomaterial particles embedded in the fibrous tissue at day 20 post-implantation (Fig. 8). Giant cells, as well as blood vessels, were observed surrounding biomaterial particles. Histomorphometric analysis indicated that the area percentages of biomaterial particles in region I were approximately 55±8%.

The tissue of Region II was avascular cortical bone, characterised by osteocytes and canaliculi (Fig. 9). There were more osteocytes in the cortical bone of 40-day sample (Fig. 9a) than in the bone of 60-days group (Fig. 9b). Almost no biomaterial particles could be observed in Region II under optical microscopy. The newly formed bone was stained red by Goldner’s trichrome after both 40 and 60 days post implantation. Goldner’s trichrome stain is sensitive to the level of mineralisation in bone and thus has been used to discriminate non-mineralised bone (immature bone, red) and mineralised bone (mature bone, green/blue) in undecalcified samples. The red stain by trichrome in 40 and 60 days samples indicated that the mineralisation in the bone remained poor after 60 days implantation, apparently not being enhanced by the degraded CaP-based material.

TEM Examination

TEM examination revealed that many micro- or sub-micron-sized particles presented in what were most likely giant cells in Region I of the 20 days samples (Fig. 10a) and these sub-micro-sized particles were broken down further into nano-sized particles (Fig. 10b). Very fine biomaterials particles were still visible in newly formed woven bone tissue in the 20 days samples (Fig. 10c). However, it was hard to find any biomaterial debris in lamellar bone in 40 and 60 days samples (Fig. 10d).

Summary of Histological Observations

The degradation of CaP particles occurred primarily in Region I. Large agglomerates were first infiltrated and fragmented by fibrous tissue (Fig. 5a) and sub-micron sized particles were phagocytosed by giant cells, further broken down to nano-sized particles, which were fused by lysosomes (Fig. 10b). The CaP material degradation was virtually complete and replaced entirely by new bone.

The critical sized defects were nearly completely filled with new bone tissue after 40 days post-implantation and healed completely with non-vascularised cortical bone.
Fig. 8. Three representative high power views of H&E stained tissue in Region I seen in samples of CaP-biomaterial implanted bone defects retrieved at 20 days post implantation. Biomaterial particles (BP) were surrounded by fibrous tissue with prominent Blood Vessels (BV), where Giant Cells (GC), presumed to be macrophage polykaryons consistent with a foreign body reaction, were noted in proximity to the BP.

Fig. 9. High power of Goldner’s trichrome stained avascular cortical bone after (a) 40 and (b) 60 days implantation.

However, after 40 days implantation, the mineralisation of the non-vascularised cortical bone remained poor, this process apparently not being enhanced by the CaP-based biomaterial.
Fig. 10. Transmission Electron Microscope (TEM) images of (a) biomaterial particles inside a giant cell and (b) the particles near the nucleus of the cells, in Region I in samples retrieved at 20 days post implantation. (c) Woven bone observed in a sample retrieved at 20 days post implantation. (d) Lamellar bone observed in a sample retrieved at day 40 post implantation; similar morphologies were also observed in samples retrieve at day 60 (not shown). BP indicates biomaterials particles. L indicates lysosome and N indicates a cell nucleus.

Discussion

The degradation of the implanted CaP-based particles primarily occurred in Region I where the amount of biomaterial particles, as indicated by significant reduction in average Ar. % of particles from initial 100% at day zero, to 50% at 20-days post implantation. The possible fates of sub-micron-sized particles produced upon degradation of the material include phagocytosis by macrophages (Fig. 10a and b), retention in the newly formed bone (Fig. 8 and 10c) and dissolution (i.e., ionising into Ca$^{2+}$, Zn$^{2+}$ and PO$_4^{3-}$) during remodelling of the bone matrix. Part of the released Ca$^{2+}$, Zn$^{2+}$ and PO$_4^{3-}$ ions could participate in the calcification of the newly formed bone. However excessive amounts of minerals are likely to have been removed via cellular mechanisms, as indicated by the evidence of foreign body reaction (Fig 10b). It is likely that the observed bone marrow and osteocytes (as well as canaliculi) in the newly formed bone (i.e., Region II, Fig. 9) could serve as efficient channels in removing excess mineral ions.

The migration kinetics of excessive ions in bone can be predicted using one-dimensional Fick’s second law:

$$\frac{\partial C(x,t)}{\partial t} = \frac{\partial}{\partial x} \left( D \frac{\partial C(x,t)}{\partial x} \right)$$

(2)

where, $C$ is the concentration of the diffusing ion, which is the function of time $t$ and distance $x$. $D$ is diffusion coefficient of the ion in the bone matrix, which is $10^{-7}$ cm$^2$/s in bone (Fernandez-Seara et al., 2002). If the diffusion coefficient is independent of concentration, Equation 2 simplifies to:

$$\frac{\partial C(x,t)}{\partial t} = D \frac{\partial^2 C(x,t)}{\partial x^2}$$

(3)

The concentration $C$ of mineral ions is approximately considered as 1 at the biomaterial site and zero at the sink site (Fig. 11). When the ions migrate from degrading particles to blood vessel to reduce the concentration from $C=1$ to the physiological level (the weight percentage of bone minerals in bone is 0.7). The boundary conditions are summarised as follows Equation 4:

$$t > 0, \begin{cases} C = C_0 = 0 \text{ at } x = 0 \\ C = C_a = 1 \text{ at } x = \infty \end{cases}$$

(4)

Application of these boundary conditions to Equation 3 yields the solution:

$$\frac{C(x,t) - C_a}{C_0 - C_a} = 1 - e^{\left( -\frac{x}{2\sqrt{Dt}} \right)}$$

(5)
For $C(x, t) = 0.7$, Equation 5-7 becomes:

$$\text{erf} \left( \frac{x}{2\sqrt{Dt}} \right) = 0.7$$

(6)

i.e.,

$$\frac{x}{2\sqrt{Dt}} \approx 0.75$$

(7)

A typical distance between lacunae and biomaterial particles was ~50 µm (Fig. 8). If $x = 50$ µm, then $t \approx 120$ s. Hence, once a biomaterial particle is ionised, it only takes several minutes for ions to migrate to a sink (lacuna or bone marrow) to reduce their concentration to the physiological level (70-75wt%). The slow mineralisation process after 40 days might be due to the rapid release of the $\text{Ca}^{2+}$ and $\text{PO}_4^{3-}$ ions from Region II at 40 days post implantation. Considering both ions are readily soluble, an ideal CaP material should have a relatively slow degradation rate to provide a more sustained source of mineral ions for new bone tissue, rather than an initial pulse. This slow-release may enhance the mineralisation of new bone at later stages. Future experiments will explore the use of CaP of high crystallinity and other forms of CaP, such as OCP, which has been shown to be more stable and able to stimulate osteoblasts in a mouse calvaria model (Sasano et al., 1999).

**Conclusion**

This study describes a histological investigation of the remodelling process of critical-sized defects in rat calvarial bone at both the optical and electron microscopic levels, following repair with degradable biomaterials containing CaP, with a focus on its influence on mineralisation. We found that implanted CaP/chitosan particles remarkably accelerated the bone growth rate of the defects at the initial stage, with an estimated growth rate of 0.8 mg day$^{-1}$, which is almost ten-fold higher than the bone growth rate (0.1 mg day$^{-1}$) in the non-implanted control group. However, we also noticed that implanted CaP did not enhance the mineralisation of regenerated cortical bone tissue at the later stage of bone remodelling. A theoretical analysis using Fick’s law indicated that the excessive amounts of mineral ions due to the fast degradation of the biomaterial may be released rapidly from the new bone tissue. Hence, a CaP material with a reduced degradation rate could offer a sustained resource of mineral ions and thus enhance bone mineralisation at later stages.

**Author’s Contributions**

TF, QZC and CHV designed the study. Samples were prepared by MAQ. Animal study experiments were carried out by TF. The cell culture experiments were carried out by YL and GAT. The data were analysed by TF, YL, GAT, MAO and QZC. The manuscript was written by QZC with contributions from TF, CHV, YL, GAT, DFN and MAO.

**Ethics**

There are no ethical issues.

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