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Microporous calcium phosphate ceramics driving osteogenesis through surface architecture

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Abstract: The presence of micropores in calcium phosphate (CaP) ceramics has shown its important role in initiating inductive bone formation in ectopic sites. To investigate how microporous CaP ceramics trigger osteoinduction, we optimized two biphasic CaP ceramics (i.e., BCP-R and BCP-S) to have the same chemical composition, equivalent surface area per volume, comparable protein adsorption, similar ion (i.e., calcium and phosphate) exchange and the same surface mineralization potential, but different surface architecture. In particular, BCP-R had a surface roughness (Ra) of 325.4 \pm 58.9 nm while for BCP-S it was 231.6 \pm 35.7 nm. Ceramic blocks with crossing or noncrossing channels of 250, 500, 1000, and 2000 μ m were implanted in paraspinal muscle of dogs for 12

weeks. The percentage of bone volume in the channels was not affected by the type of pores (i.e., crossing vs. closed) or their size, but it was greatly influenced by the ceramic type (i.e., BCP-R vs. BCP-S). Significantly, more bone was formed in the channels of BCP-R than in those of BCP-S. Since the two CaP ceramics differed only in their surface architecture, the results hereby demonstrate that microporous CaP ceramics may induce ectopic osteogenesis through surface architecture. © 2014 Wiley Periodicals, Inc. J Biomed Mater Res Part A: 103A: 1188–1199, 2015.

Key Words: bone grafting, calcium phosphate ceramics, microstructure, surface architecture, osteogenesis

tion without the addition of cells or growth factors through

a process defined as "osteoinduction." Such phenomena has

been reported in several animal models, where hydroxyapa-

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INTRODUCTION

Searching for alternatives to autologs bone graft, which is the current gold standard grafting material to heal bone defects, has become a goal of biomedical engineering.¹⁻³ Calcium phosphate (CaP) ceramics are applied for bone grafting because their chemical composition resembles the bone mineral phase.⁴⁻⁶ Traditionally, CaP ceramics were used as osteoconductive or osteoinductive bone graft substitutes in combination with cells and/or growth factors. For instance, CaP ceramics delivering mesenchymal stromal cells (MSCs) to the bone defect site could induce bone formation.⁷ Alternatively, a subclass of CaP ceramics with specific physicochemical properties can induce ectopic bone forma-

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Correspondence to: H. Yuan; e-mail: h.yuan@utwente.nl Contract grant sponsor: China Scholarship Council tite (HA) ceramic induced heterotopic bone formation in dogs,^{8,9} baboons,^{4,10-12} rabbits,¹³ mouse,¹³ sheep,¹⁴ and goats.^{15,16} Similarly, it was shown occurring also in tricalcium phosphate (TCP) ceramics in dogs,^{4,9,17-20} sheep,²¹ and in biphasic calcium phosphate (BCP) ceramics implanted in dogs,^{5,17,18,22} pigs,⁵ sheep,^{14,23} goat,^{16,24-27} rabbits,^{13,28} and baboons.¹¹ Previous studies have shown the possible clinical importance of such CaP ceramics by comparing them in both nonosseous and osseous sites.^{22,29} Moreover, a resorbable osteoinductive TCP ceramic has

shown equivalent bone forming ability to autologs bone graft and recombinant human bone morphogenetic protein-2 (rhBMP-2) in critical-sized bone defects.²¹ This finding strongly reveals the potential of such specific ceramics to be alternatives to autologs bone grafts.

The available data may have shown the relevance of material properties to its functionality in bone regeneration. Physicochemical modification of CaP ceramics may represent an approach for bone defect repair. It is well known that the chemistry of CaP ceramics influences on their degradation behavior, which in turn affects osteogenesis. For instance, in a previous study of ours, HA and BCP ceramics were implanted intramuscularly in goats: higher bone incidence and amount were found in the BCP ceramic—which contains the more soluble TCP—as compared to HA ceramic.¹⁵ The geometry and macrostructural properties of ceramic scaffolds have been shown to play critical roles in bone formation as well. Ripamonti et al showed that bone formation always started from the concave side of the macropores and never from the convex spaces of HA rods and discs.³⁰

It is generally accepted that the presence of micropores in CaP ceramics plays an important role in inducing heterotopic bone formation. For instance, microporous HA ceramics could induce bone formation after subcutaneous⁸ and intramuscular⁶ implantations in dogs, while no bone formed in those HA ceramics lacking of micropores. The microstructure may improve osteoinduction by enlarging the surface area, thus affecting protein adsorption, ion release and mineral deposition. Protein adsorption, that is the adsorption of growth factors including bone morphogenetic proteins (BMPs),^{31,32} is believed to play a crucial role in bone induction. It has therefore been suggested that micropores in CaP ceramics may concentrate higher amounts of growth factors thanks to their enlarged surface area.^{15,21} It was also proposed that the microporous CaP ceramics can enhance the ion exchange processes because of their enlarged surface area, later affecting osteogenic differentiation of MSCs.³³ The ions (e.g., Ca^{2+} and PO_4^{3-}) in the surroundings might trigger the bioactivity of CaP ceramics, which may later allow bone formation.³⁴ In addition, the Ca^{2+} and PO_4^{3-} in the body fluids could also form a biological apatite layer on the surface of the CaP ceramics.^{16,35} The possible role of the formation of an apatite layer in osteoinduction is supported by the fact that bioactive metals could induce ectopic bone formation. For example, bioactive titanium implants that could get surface mineralization induced bone formation after ectopic implantation, while titanium implants not able to get such mineralization then could not induce any bone formation.35

In addition to the improved protein adsorption and ion exchange, micropores in CaP ceramics can generate different surface architectures, which have shown to influence the osteogenic differentiation of MSCs through various physical properties such as surface roughness,^{36–40} the size of surface pattern,^{41,42} and surface morphology.^{43–45} For instance, Fu et al. showed that micropost rigidity can affect cell morphology, focal adhesions, cytoskeletal contractility, and differentiation of stem cells.⁴⁶ In addition, Oh et al. showed

the critical role of the nanotube dimensions on osteogenic differentiation of hMSCs in absence of osteogenic inducing media.⁴² Finally, Dalby et al. demonstrated that nanoscale disorder could stimulate osteogenic differentiation of hMSCs *in vitro* without osteogenic supplements.⁴⁵ Therefore the role of surface architecture of CaP ceramics in ectopic osteogenesis cannot be excluded. The surface arthitecture could be varied by controlling the sintering temperature of CaP ceramics: a decrease in grain size and increase in microporosity have shown to positively affect on osteoinductive potential of CaP ceramics.^{16,21} Considering the surface properties, it is known that the presence of a high number of small features on the surface indicates high roughness values.

Therefore, from previous results suggesting that CaP ceramics with small grains trigger heterotopic bone formation, here we expect that CaP ceramics with high roughness will lead to ectopic bone formation. To explore whether the surface architecture affects osteogenesis by microstructured CaP ceramics, we optimized two types of BCP ceramics (i.e., BCP-R and BCP-S) to have the same chemical composition, equivalent surface area per volume, comparable protein adsorption, similar ions (e.g., Ca²⁺ and PO₄³⁻) exchange and same surface mineralization potential, but different surface architecture. We then compared their capacity to induce osteogenesis in a canine osteoinduction model.

MATERIALS AND METHODS

Preparation and optimization of BCP ceramics

Two types of BCP, namely BCP-R and BCP-S, were prepared from the same CaP apatite powder with a Ca/P ratio of 1.64. BCP-R was obtained by mixing CaP apatite powder with 0.1% H₂O₂ solution to get slurry, which was subsequently dried at room temperature and sintered at 1150°C for 8 h. BCP-S was prepared by isostatically pressing the starting apatite powder in a steel cylinder, with inner diameter of 1.25 inch, at a certain pressure for 5 min at room temperature and thereafter sintering it at 1150°C for 8 h. Importantly, the pressure was optimized to get BCP-S having the same protein adsorption as BCP-R (see section "Protein adsorption"): at last the chosen BCP-S was prepared using a pressure of 50 MPa. The ceramic samples were then made into three forms. Discs (\emptyset 10 \times 1 mm) were fabricated using a lathe (Esmeijer, Netherland) and a histological diamond saw (SP-1600, Leica, Germany), while blocks (7 \times 9 \times 12 mm) were made using a diamond saw (Heathway, England). Afterwards, two types of channels (i.e., crossing and noncrossing) with different diameters of 250, 500, 1000, and 2000 μ m were made along the shortest side of the BCP blocks using drills with the required diameter. BCP-R ceramic implants had both crossing and noncrossing channels, while BCP-S ceramic implants only had noncrossing channels. Because of technical challenges, it was not possible to obtain crossing channels with a diameter of 250 μ m in BCP-R. Granules (size 1-2 mm) were obtained by crushing and sieving the ceramics. The crossing channels were made to study the possible role of pore interconnectivity in bone induction. All the samples were ultrasonically cleaned

with acetone, 70% ethanol and demineralized water, then dried at 80° C and sterilized with gamma irradiation (Isotron, Netherland BV, Ede, The Netherlands) before further use.

Physicochemical characterization of the ceramics

The chemical composition of the samples was determined with X-ray diffraction (XRD; Rigaku, Japan). The HA/β-TCP weight ratios in the BCP ceramics were determined with a calibration line obtained from internal standard powders with known HA/B-TCP weight ratios. The surface microstructure of the BCP ceramics was studied with scanning electron microscope (SEM; XL30, ESEM-FEG, Philips, Eindhoven, The Netherlands) in the secondary electron mode. The micro-porosity, micro-pore size distribution, and specific surface area of the ceramic blocks were measured using a mercury intrusion porosimeter (MP; Auto Pore IV 9500, Micromeritics European Analysis Service, Monchengladbach, Germany). The density of the ceramics was measured with the weight/volume method using the ceramic discs. To better evaluate the surface properties, we calculated the surface area per volume as the product between the specific surface area measured with mercury intrusion and the measured density of the ceramic. The surface roughness of the ceramic discs was analyzed with atomic force microscopy (AFM; Bioscope Catalyst AFM, Bruker AXS, Wormer, the Netherlands) using dual scans in tapping mode, at a scan frequency of 1 Hz, on areas of 10 \times 10 μm with a MikroMarsch NCS36 cantilever (total tip height 20-25 μm, tip radius 10 nm and full type cone radius of 40°, spring constant 1.75 N/m). Obtained images were filtered from signal noise and then analyzed with SPIP software (v5.1.11, Image Metrology A/S, Hørsholm, Denmark). The surface roughness was estimated in terms of roughness average (Sa) and root mean square (Sq). Further, measurements on the profile height changes as compared to the zero-plan were also taken and plotted as distribution.

Protein adsorption

Fetal bovine serum (FBS, Lonza, Germany) and bovine serum albumin (BSA, Sigma) were diluted in 25 ppm aqueous NaN₃ solution and sterilized with 0.22 μ m filter to get the solutions of FBS (0.25%) and BSA (100 $\mu g/mL$). After immersing the discs in the solutions (3 mL per disc, n = 5per material) at 37°C for 24 h, the remaining proteins in the solutions were measured with the QuantiProTM BCA assay kit (Pierce, The Netherlands) following the manufacturer's guidelines and absorbance was measured with a spectrophotometer (AnthosZenyth 3100, AnthosLabtec Instruments GmbH, Salzburg, Austria) at 595 nm. The amount of proteins adsorbed from FBS solution onto the discs (microgram of protein per disc) was estimated through a calibration serum protein curve using FBS after determining the total proteins content in pure FBS (i.e., 33.78 ± 0.64 mg/mL). In the meantime, a calibration line for BSA was built with a series of concentrated BSA solutions. All measured values were expressed as mean \pm standard deviation.

rhBMP-2 adsorption

A solution of rhBMP2 (500 ng/mL) was prepared by diluting rhBMP-2 (Reborne, Shanghai) in the basic medium (BM) containing Dulbecco's modified Eagle's medium (DMEM; Gibco, Invitrogen, UK), 10% FBS and 1% penicillin/streptomycin (100 U/mL penicillin, 100 mg/mL streptomycin; Gibco, Invitrogen, UK). The ceramic discs were then soaked in the medium (3 mL per disc, n = 5 per material) at 37°C in humid atmosphere with 5% CO₂ for 24 h. The discs were stored at -80° C for at least 12 h after being washed with PBS three times. Thereafter 0.5 mL of 1% triton was add to each sample and kept at 4°C for at least 12 h and followed by ultrasonically shaking for 5 min. Finally, rhBMP-2 content in the supernatant was measured with BMP-2 Elisa kit (R&D, UK) following the manufacturer's instructions and absorbance was measured with a spectrophotometer at 450 nm. The rhBMP-2 adsorption was estimated through a standard calibration rhBMP-2 curve and expressed as mean \pm standard deviation.

Ion exchange

Ceramic discs were soaked in BM (3 mL per disc, n = 5 per material) at 37°C in humid atmosphere with 5% CO₂ for 24 h. Calcium and phosphate contents in the solutions were measured and compared to the control (i.e., solution with no sample). Calcium and phosphate ion concentration were measured with QuantiChromTM Calcium assay kit (BioAssay, USA) and phosphoWorkTM Colorimetric Phosphate Assay Kit (AATBioquest, USA), respectively following the manufacturer's guidelines. Thereafter, absorbance was measured with spectrophotometer at 620 nm. Both calcium and phosphate contents were expressed as mean \pm standard deviation.

Surface mineralization

Simulated body fluid (SBF) was prepared according to Ref. [47 by dissolving reagent grade chemicals in distilled water strictly in the following order: NaCl, NaHCO₃, KCl, K₂HPO₄·3H₂O, MgCl₂·6H₂O, CaCl₂ (Ca ion standard solution, 0.1M, Metrohm) and Na₂SO₄. The solution was then buffered to pH 7.4 at 36.5°C using Tris ((CH2OH)3CNH3) and 1M HCl. The final solution had an ion concentration (mM) as follows: Na⁺, 142; K⁺, 5; Mg²⁺, 1.5; Ca²⁺, 2.5; Cl⁻, 147.8; $(HCO_3)^-$, 4.2; $(HPO_4)^{2-}$, 1; $(SO_4)^{2-}$, 0.5. Fifty granules (1-2 mm) of each material were soaked in 200 mL of SBF at 37 \pm 1°C for up to 1 week. A minimum of 10 granules were taken out at each time point of 1, 2, 4, and 7 days. They were rinsed five times with distilled water and dried at room temperature. The mineralization of the BCP ceramics was observed with SEM. Four spots were randomly selected from each granule. The mineralization level of the materials was ranked as follows: (a) no mineralization: all the spots were completely bare; (b) few mineralization: one (or more) of the spots had thin and not uniform mineralized layers; and (c) full mineralization: all the spots were covered by thick and uniform mineralized layers. For each classification group, the number of granules was

counted and plotted on a histogram as percentage on the total granules considered.

Evaluation of osteoinduction

Animal experiments. Following the permission of the local animal care committee (Animal Center, Sichuan University, Chengdu, China), ceramic blocks (n = 8 per ceramic) were implanted in the para-spinal muscles of 8 adult male dogs (mongrel, 10–15 kg) for 12 weeks. The surgical operation was performed under general anesthesia by abdominal injection of sodium pentobarbital (30 mg/kg body weight) and sterile conditions. Following the surgeries, buprenorphine (0.1 mg/animal) was intramuscular injected to the dogs for 2 days to relieve pain, while penicillin (40 mg/kg) was intramuscular injected for three consecutive days to prevent infection. After operation, the animals were allowed for full weight bearing and received normal diet.

Histological processing. After 12 weeks the animals were sacrificed with a celiac injection of excessive amount of pentobarbital sodium. Implants were then harvested with the surrounding tissues and fixed in 4% buffered formaldehyde solution (pH = 7.4) at 4° C for one week, and embedded in methyl methacrylate (MMA, K-plast, LTI Netherland, Bilthoven, the Netherlands) after dehydration in a series of gradient ethanol solutions. Serial nondecalcified sections with an average thickness of 15 µm were obtained without interruption using a microtome (SP-1600, Leica, Germany) equipped with diamond saw blade having an approximate thickness of 280 µm and a knurled screw to control the thickness of the sections. Being the sectioning done serially, the distance between consecutive sections is ${\sim}280~\mu m$ due to loss of material by the blade. Each slide was stained with 1% methylene blue (Aldrich) and 0.3% basic function (Aldrich). Sections were also prepared from MMA-embedded ceramic blocks (not implanted) to be used as controls.

Histological and histomorphometric analysis. The histological sections were observed with light microscopy (Nikon Eclipse E200, Tokyo, Japan) with respect to tissue formation in the channels. The sections were then scanned using a slide scanner (Dimage Scan Elite 5400II, Konica Minolta Photo Imaging Inc, Tokyo, Japan) to obtain overview images for histomorphometrical analysis, which was performed using Adobe Photoshop Elements 4.0 software. The channel area was selected as a region of interest and the corresponding number of pixels was read as ROI. Then the bone tissue area was pseudo-colored and its respective pixels were counted as B. The percentage of bone area in the channel (B%) was determined as B% = B*100/ROIand the results were presented as mean and standard deviation of all the sections taken at the same depth from all the samples. As shown in the aforementioned equation, we normalized the bone area to the ROI of the corresponding channel to make the comparison of bone formation among the channels with different dimension possible. The ROI of the channels in the control ceramic blocks was measured as well, and the resorption rate (R%) of the ceramics after explant was determined as R% =

(ROI of the control-ROI of the explants)*100/(ROI of the control). The measurements were taken both at the openings and in the central region of the channels.

Statistical analysis

Statistical analysis was performed using a one-way ANOVA followed by a Tukey's multiple comparison test or paired *t*-test. A value of p < 0.05 was considered as significant difference.

RESULTS

Material characteristics

The XRD patterns of the ceramics are given in Figure 1(a), which shows the two BCP ceramics had similar chemistry consisting of 80 \pm 3 wt % HA and 20 \pm 3 wt % $\beta\text{-TCP}.$ The SEM observation demonstrated a rougher surface of BCP-R [Fig. 1(d)] than that of BCP-S [Fig. 1(c)]. This observation was further confirmed by AFM images [Fig. 1(e,f)]. Smaller pores could also be observed on BCP-S than BCP-R as confirmed also by mercury intrusion test [Fig. 1(b)]. The average roughness (Sa) and the root mean square (Sq) resulted significantly higher in BCP-R than BCP-S (p = 0.0014 and p= 0.0019, respectively) indicating that the former material presents a topography having larger and deeper variability in its peaks and hollows heights over the area [Fig. 1(h)]. These results are corroborated by the distribution of surface heights [Fig. 1(g)], which is larger for BCP-R. Interestingly, the distribution of surface heights on BCP-R presents a lager tail on the right as compared to BCP-S, which might be correlated with the preparation method of BCP-S involving a step of isostatic pressure that could have flattened the surface. In addition, the mercury intrusion results showed the two BCP ceramics had different porosity and distribution of micropore size. A micropore range of 0.3 and 1.3 µm with a peak at 0.9 µm for BCP-R, whereas BCP-S had micropore range from 0.3 to 0.7 μ m with a peak at 0.6 μ m [Fig. 1(b)]. The total microporosity was higher for BCP-R (43%) than for BCP-S (28%). Mercury intrusion data revealed also that the surface area per volume of BCP-R and BCP-S were the same (Table I).

Protein adsorption assay

The amount of protein absorbed by the two ceramics is shown in Figures 2(*c*, *d*, *e*), where no significant differences between BCP-R and BCP-S with respect to adsorption of BSA (p = 0.702), of proteins in FBS (p = 0.387), and of rhBMP-2 (p = 0.632) were observed. These results might be explained by the fact that the two ceramics have the same surface area.

Calcium and phosphate ions assay

Changes in calcium and phosphate ion concentrations in BM were evaluated [Fig. 2(a,b)]: instead of releasing calcium and phosphate ions into the medium, both ceramics adsorbed ions from the medium within 24 h. However, BCP-R and BCP-S adsorbed similar amounts of calcium and phosphate ions and no significant differences were observed (calcium: p = 0.153, phosphate: p = 0.088).



FIGURE 1. (a) XRD pattern of the two ceramics, showing that BCP-S and BCP-R have the same chemistry. (b) Micropore distribution of BCP-S and BCP-R as obtained from mercury intrusion measurements. It can be seen that BCP-S has smaller micropore size than BCP-R. The surface microstructure of (c) BCP-S and (d) BCP-R was observed with SEM. AFM allows evaluating the surface roughness of (e) BCP-S and (f) BCP-R. The height distributions of the microstructured surface of the two ceramics is shown in (g). Histograms in (h) allow the comparison of the measured surface roughness average (Sa) and root mean square (Sq) between the two ceramics. Statistically significant differences are marked by * for p < 0.05. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

Surface mineralization

Mineralization on the BCP ceramics was observed as early as after soaking in SBF for one day [Fig. 3(a,b)], and the amount of mineralization increased during time finally leading to thick layers of apatite that uniformly covered the surface of all the samples after seven days [Fig. 3(c,d)]. The trend of surface mineralization over time is illustrated in Figure 3(e), where no significant differences between the two BCP ceramics were observed after seven days.

TABLE I. Density, Specific Surface Area and Surface Area per Volume of the Two BCP Ceramics: The Density and the Specific Surface Area Data Were Experimentally Measured and Used to Calculate the Surface Area per Volume that Resulted to be the Same for the Two Ceramics

| Density (g/mL) | Specific Surface Area (m ² /g) | Surface Aarea per Volume (m²/mL) | | |
|---|---|---|--|--|
| $\begin{array}{r} 1.758\pm0.026\\ 2.242\pm0.026\end{array}$ | 1.057 0.825 | 1.859 1.854 | | |
| | Density (g/mL) 1.758 ± 0.026 2.242 ± 0.026 | Density (g/mL) Specific Surface Area (m²/g) 1.758 ± 0.026 1.057 2.242 ± 0.026 0.825 | | |

Ectopic bone formation

A total of eight samples per ceramic were intramuscularly implanted in eight dogs for 12 weeks, and all were retrieved. Histological observations demonstrated that bone formed in most channels of BCP-R [Fig. 4(b,d)], while in BCP-S little bone amounts were found in few channels and fibrous tissue was dominant [Fig. 4(a,c)]. Bone grew tight to the wall of the channels [Fig. 4(d)]. The distribution of the newly formed bone along the crossing channels with different diameters (i.e., 500 $\mu\text{m},$ 1000 $\mu\text{m},$ and 2000 $\mu\text{m})$ of BCP-R is shown in Figure 4(e). It could be seen that similar trends in the distribution of new bone exist among the different channels: no bone formation at the openings (i.e., at 0 and 7000 μ m on the x-axis); bone formation always started at about 500 µm far from both openings (i.e., at 500 and 6500 μ m on the x-axis); high amount of bone formation was observed in the central part of the channels (i.e., between 750 and 6000 µm on the x-axis). Furthermore, it appeared that higher amount of bone formed in the channels with 1 mm diameter rather than in the others, but no significant differences in the bone amount were seen since



FIGURE 2. (a) Ca^{2+} and (b) PO_4^{3-} concentrations in culture medium after soaking the two ceramics' discs for 24 h: they were significantly lower than the control medium in both cases, but differences in Ca^{2+} and PO_4^{3-} amounts in the medium with BCP-S and BCP-R were not significant. Protein adsorption of the ceramic discs after soaking in (c) BSA, (d) FBS, and (e) rhBMP-2 solutions, indicating that BCP-S and BCP-R have similar protein adsorption potential. Any statistically significant difference is marked with *** for p < 0.001. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]



FIGURE 3. Surface mineralization of BCPs after soaking in SBF solution. SEM images of surface mineralization on BCP-S with (a) few and (c) full surface mineralization and BCP-R with (b) few and (d) full surface mineralization. (e) Percentage of BCPs ceramics with no, few and full surface mineralization at days 1, 2, 4, and 7. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

the standard deviation was high (p > 0.05). Table II shows the amount of bone formation in each channel of the BCP ceramics. In BCP-R ceramics: with the same diameter, the type of channel (i.e., crossing and noncrossing) does not affect the amount of bone formation (p > 0.05); within the same type of channel, the diameter of the channel does not affect the amount of bone formation either (p > 0.05). However, the amount of bone formation was significantly affected by the ceramic type (i.e., BCP-R vs. BCP-S), where significantly higher amounts of bone formed in the channels of BCP-R (p < 0.01 or p < 0.001). The histological overviews [Fig. 4(a,b)] show that the channels of both BCP ceramics retained their shape after a 12-week implantation indicating no obvious resorption of both ceramics in vivo. The quantitative data confirmed this observation (Table III) where no significant resorption was seen in both ceramics. In particular, the measurements showed no effect of the

diameter or region (i.e., at the openings and in the central sector) of the channel on the ceramics resorption.

DISCUSSION

In this study, we used optimized BCP ceramics to investigate whether the surface architecture is playing a role in material-directed osteoinduction. We could minimize the number of possible influencing factors and make sure that the two ceramics differed only in their surface architecture. After a 12-week intramuscular implantation, BCP-R gave rise to abundant bone formation while BCP-S could induce the formation of a little amount of bone. The results presented herein thus indicate an important role of surface architecture in the osteogenesis process by CaP ceramics.

Chemistry,^{21,22,42} macrostructure,^{11,48} and microstructure^{16,49,50} have been suggested to play roles in material dependent osteoinduction. The possible influence of chemistry and macrostructure could be excluded in our study. The



FIGURE 4. Histology of the BCP ceramics (a and c: BCP-S; b and d: BCP-R) after 12 weeks in muscle of dogs. The green rectangles in (a) and (b) highlight the areas observed with light microscope in (c) and (d). Figure (e) shows bone formation area along the crossing channels in BCP-R. Legend of symbols: ST stands for soft tissues, M for material, and B for bone. [Color figure can be viewed in the online issue, which is available at wileyonlinelibrary.com.]

two BCP ceramics were made from the same CaP powder and, as shown in the XRD patterns, both BCP ceramics had the same composition with 80 \pm 3% wt HA and 20 \pm 3% wt β -TCP. This indicated that the different manufacture methods did not affect the final chemistry of the BCP ceramics. The macrostructure of the two BCP ceramic

implants was kept equal: both BCP ceramics contained noncrossing channels with different diameters (i.e., 250, 500, 1000, and 2000 μ m), and the channels were created in the same positions in all the blocks.

The histological analysis showed that the newly formed bone was not on the surface of the implants, but it always

| TABLE II. A Summar | of the | Bone Forming | Ability In | Vivo foi | r the Ceramics | Studied |
|--------------------|--------|--------------|------------|----------|----------------|---------|
|--------------------|--------|--------------|------------|----------|----------------|---------|

| Samples | | | Bone Incidence (%) | Bone in Pores (% wt) |
|---------|---|------|--------------------|----------------------------------|
| BCP-R | Noncrossing (Ø: mm) | 2 | 100 | 11.0 ± 3.8 |
| | | 1 | 100 | 17.7 ± 5.7 |
| | | 0.5 | 100 | 7.9 ± 9.4 |
| | | 0.25 | 75 | 15.5 ± 19.5 |
| | Crossing (Ø: mm) | 2 | 100 | $14.3~\pm~5.8$ |
| | C A A A A A A A A A A A A A A A A A A A | 1 | 100 | $\textbf{22.3} \pm \textbf{9.6}$ |
| | | 0.5 | 87.5 | 6.8 ± 9.4 |
| BCP-S | Noncrossing (Ø: mm) | 2 | 50 | 0.6 ± 1.1 |
| | C C | 1 | 25 | 0.9 ± 2.4 |
| | | 0.5 | 62.5 | 1.8 ± 2.3 |
| | | 0.25 | 31.25 | 1.8 ± 3.7 |

It can be observed that the bone incidence as well as the newly formed bone volume were significantly higher in BCP-R than BCP-S when considering the case of closed channels (i.e., with only one opening). The presence of a second opening in the channels (i.e., two sides open case) did not significantly change the *in vivo* performance of BCP-R.

formed inside the channels suggesting the important role of macrostructure in osteoinduction. This result is consistent with a previous study,¹⁹ where it was concluded that a concave structure is necessary for osteoinduction in CaP ceramics. To further investigate the effect of macrostructure dimension on bone formation, we fabricated both BCP ceramic implants with different channels' diameters. When analyzing the ceramics, no significant difference was observed in both the bone incidence and the bone amount among the channels having different diameter. It is worthy of notice that this fact was found for both BCP-R and BCP-S, therefore the diameter of the channels did not seem to be essential for bone ingrowth in the tested range. However, the measured percentage of bone in the 12-week samples could not address the dynamic metabolism of the formed bone regarding its growth towards the center of the channels and its remodeling, therefore the possible influence of the channels' diameter on bone metabolism could not be excluded. It may well be that, at other time points such as later than 12 weeks, the smaller channels would have more bone fill than the larger ones as the latter will take longer to be fully colonized by bone tissue. Furthermore, we observed that there was no significant difference between crossing and noncrossing channels in BCP-R, which is in good agreement with a previous study¹² where it was claimed that interconnected macropores are not necessary for osteoinduction. However, the histological results showed that the incidence and amount of bone induced by BCP-S were lower than BCP-R regardless of the channel diameter (Table II). Therefore, the surface microstructure may have played roles as main determinant of the osteoinductive potential of the BCP ceramics, which was already shown in several studies.^{16,46,49} It is generally suggested that microporous CaP ceramics affects bone formation by virtue of its enlarged surface area since it would result in higher protein adsorption (e.g., BMP-2), higher calcium and phosphate ions release and induced surface mineralization, or their combinations.

Both BCP-R and BCP-S used in this study had the same volume and equal surface area as shown in Table I. Therefore, the surface area was not the main factor influencing the bone forming potential. Meanwhile, the two BCP ceramics adsorbed similar amounts of proteins from BSA and FBS solutions, and they also could uptake rhBMP-2 from cell culture medium (Fig. 4). These results confirm the fact that surface area of CaP ceramics influences the protein adsorption. Having equal surface area, BCP-R and BCP-S adsorbed equivalent amounts of proteins regardless of the type of protein. The results also show that the protein adsorption could not be appointed as the main determinant for osteoinduction, since bone formation was different between BCP-R and BCP-S which adsorbed the same amount of proteins.

It has been reported that extracellular calcium and phosphate significantly influenced the growth and osteogenic differentiation of MSCs,⁴⁸ meanwhile the calcium and phosphate supplementation promoted bone cell mineralization.¹⁶ It has been also suggested that have been made the dissolution of calcium and phosphate ions can be improved when biomaterials are precoated with CaP apatite layers. When titanium alloy was covered by an apatite layer, it could dissolve and release ions that can favor osteogenic

TABLE III. A Summary of Ceramic Resorption *In Vivo* (as Measured for the Noncrossing Channels in Both Ceramics), Which Did Not Occur Regardless of the Ceramic Type or Channel Diameter

| | | Peripheral Region of the Channels (Φ : mm) | | | | Central Region of the Channels (Φ : mm) | | | |
|------------------------|----------------|---|------------------------------------|-----------------------------|---|---|------------------------------------|-------------------------------------|----------------------------|
| | | 2000 | 1000 | 500 250 2000 1000 500 | | 500 | 250 | | |
| Resortpion rate (%) | BCP-R BCP-S | $\begin{array}{c} 0.28\pm1.47\\ 2.6\pm2.49 \end{array}$ | -1.42 ± 2.27 3.49 ± 3.5 | -1.21 ± 5.71 1.71 ± 3.39 | $\begin{array}{c} -3.24 \pm 4.42 \\ -4.32 \pm 2.13 \end{array}$ | $\begin{array}{c} 0.23 \pm 2.27 \\ 0.09 \pm 1.16 \end{array}$ | $-0.26 \pm 3.34 \\ -0.11 \pm 2.75$ | -1.99 ± 5.10 1.68 ± 4.65 | 1.28 ± 2.29 1.34 ± 1.49 |

differentiation of progenitor cells⁵¹ indicating a key role in the CaP dissolution. It is therefore possible that the microporous CaP ceramics used in our study initiated inductive bone formation via ion release during material resorption. However, no obvious resorption of both BCP ceramics was observed in this study. The reason could be that the TCP content in BCP ceramics was too low (20%) or the implantation time was too short (12 week) to allow the resorption of the ceramics. Moreover, both BCP-R and BCP-S, instead of releasing, adsorbed similar amount of calcium and phosphate ions from culture medium after 24 h *in vitro*. Therefore, the degradation rate of the materials and the CaP ions release was not the affecting factors in BCP-R ceramicinduced osteogenesis.

The ability to get surface mineralization has also been suggested being important in ectopic osteogenesis because ectopic bone formation was observed quite often in materials that can easily mineralize. Winter and Simpson reported that ectopic bone formation induced by a polymeric material occurred after surface mineralization of the implant.⁵² Fujibayashi et al. studied the in vivo osteoinductivity of titanium implants, which could or not surface mineralize in vitro in SBF. The results demonstrated that those implants that induced an apatite layer formation in vitro had osteoinductive potential in vivo.35 Similarly, Barbieri et al. reported the role of surface mineralization in vitro in materialdirected bone induction. They prepared poly (D,L-lactide) and nanosized apatite composites with various apatite contents that were used to test in vivo osteoinduction. The results showed that only composites with at least 40% wt apatite content could surface mineralize in vitro and induced bone formation in vivo.53 Besides the spontaneous formation of mineralized layers by natural precipitation of ions from biological fluids, precoating biomaterials with CaP layers has shown to improve their biological activity. For example, polycaprolactone/TCP composites with HA-like coating allowed not only better adhesion and proliferation of porcine BMSCs as compared to noncoated counterparts⁵⁴ but also better osteogenic differentiation as the expression levels of various osteogenic indicators were improved. The authors suggested that such apatite-like coating layers would mimic the bone environment thus direct cells towards bone phenotypes.⁵⁴ When a polymer, i.e., polycaprolactone, was coated with a CaP layer and then cultured with osteoblasts,55 it could induce ectopic bone formation subcutaneously. The same study observed significant lower performances of the noncoated polymer indicating a crucial role of apatite-like layers in bone induction. However, our results showed that both BCP-R and BCP-S mineralized in vitro with similar trends, thus the possible influence of surface mineralization on their different osteoinductivity could be excluded.

Taken together, the role of surface architecture (as characterized by surface roughness) in heterotopic bone formation induced by CaP ceramics could be clearly seen in this study with the well-designed materials. In fact, previous studies already showed that osteoblast-like cells favor rougher surface to smooth surface *in vitro*.³⁷⁻⁴⁰ It also has been demonstrated that microscale roughness with a high density of nanoscale features were superior to microscale roughness also in promoting osteogenic induction of the human osteosarcoma cell line MG63, determined by increased osteocalcin, osteoprotegerin, and vascular endothelial growth factor. Moreover, Hu et al. examined the changes in hMSC proliferation and differentiation after growth on silk/tropoelastin materials with different content of human tropoelastin, together with different surface roughness and micro/nanoscale surface patterns. They found that higher surface roughness with micro/nanoscale surface patterns, and by increasing the content of human tropoelastin in the materials, enhanced the proliferation and osteogenic differentiation of hMSCs.³⁶ In our study, the different methods used to prepare the materials resulted in different micro-pore size [Fig. 1(b)] and thus in different surface roughness [Fig. 1(e-h)]. The result herein showed a relationship between surface properties of materials and bone induction in vivo. Many biochemists and biologists induce osteogenic differentiation of cells by topography factors in vitro, but there is no well-understood way of controlling such factors in vivo. We demonstrated in this report a clear mechanism for osteoinduction based on the surface architecture providing an explanation for the control of bone induction by specific surface roughness. It is most probable that surface architecture could physically affect osteogenic differentiation of stem cells or undifferentiated inducible osteoprogenitor cells in ectopic sites and lead to the new heterotopic bone formation.

How the surface architecture of CaP ceramics drives osteogenesis in ectopic sites is not clear as yet. It should be mentioned that, upon injury due to the surgery, an inflammatory reaction is immediately triggered to limit the damage while blood vessels restrict to stop bleeding. During this stage, macrophage migration to the injured site and quick formation of fibrin-based platelet-rich clot may have happened.^{56,57} Macrophages and platelets, together with endothelial cells in the damaged vessels, might have released various biomolecules (e.g., cytokines and several growth factors) prompting angiogenesis and migration of (stem) cells to the site.^{56,57} In this study, the presence of the microstructured surface of BCP implants may have driven such (stem) cells towards osteogenic differentiation favoring the observed ectopic bone formation. To confirm this suggestive mechanism, a histological analysis of short-term implants, that is within few days from implantation, is recommended to analyze the blood clot formation and presence of inflammatory cells in future studies.

Recently it has been shown that scaffolds seeded with human mesenchymal progenitor cells and enriched with rhBMP-7 could ectopically form bone tissue that resembled the typical characteristics of a physiological bone, including a trabecular structure surrounded by a cortex-like layer.⁵⁸ We could not unequivocally identify the fat-like tissue present in the large channels of BCP-R [Fig. 4(d)] as fat marrow and therefore it is not possible to claim that the observed bone can be considered as a complete organ bone in the current study. However, the use of histochemical stains is

suggested in future studies to identify the possible presence of bone marrow and better understand if material-driven ectopic bone formation can lead to a complete functional bone tissue.

The optimized CaP ceramics allowed us to study whether the surface architecture of CaP ceramics could act as a cue for osteoinduction of CaP ceramics *in vivo*. Although we have not pinpointed a mechanism where surface architecture triggers ectopic bone formation of CaP ceramics, to the best of our knowledge it is the first work describing the significant role of a surface architecture in bone formation *in vivo*, instead of *in vitro* osteogenic differentiation of stem cells driven by surface characteristics as often documented in literature.

CONCLUSION

Given the same chemical composition, surface area per volume, proteins adsorption, ion release rate, and surface mineralization ability, the surface architecture (i.e., surface roughness) affects bone forming ability of biphasic CaP ceramics, with a surface roughness of Ra 325.4 ± 58.9 nm as a favorite scale for bone formation in this study. The results also showed that the percentage of bone volume in the channels was significantly affected by the surface architecture of the ceramics instead of the type (i.e., crossing vs. noncrossing) and size of the channels. Although the biological mechanism of ectopic osteogenesis of microporous CaP ceramics is far beyond understood, the current results demonstrate the surface roughness a cue for material-induced bone formation.

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