In vitro osteogenesis on a highly bioactive glass-ceramic (Biosilicate®)


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Abstract: One of the strategies to improve the mechanical performance of bioactive glasses for load-bearing implant devices has been the development of glass-ceramic materials. The present study aimed to evaluate the effect of a highly bioactive, fully-crystallized glass-ceramic (Biosilicate®) of the system P₂O₅–Na₂O–CaO–SiO₂ on various key parameters of in vitro osteogenesis. Surface characterization was carried out by scanning electron microscopy and Fourier transform infrared spectroscopy. Osteogenic cells were obtained by enzymatic digestion of newborn rat calvarial bone and by growing on Biosilicate® discs and on control bioactive glass surfaces (Biosilicate® parent glass and Bio-glass® 45S5) for periods of up to 17 days. All materials developed an apatite layer in simulated body fluid for 24 h. Additionally, as early as 12 h under culture conditions and in the absence of cells, all surfaces developed a layer of silica-gel that was gradually covered by amorphous calcium phosphate deposits, which remained amorphous up to 72 h. During the proliferative phase of osteogenic cultures, the majority of cells exhibited disassembly of the actin cytoskeleton, whereas reassembly of actin stress fibers took place only in areas of cell multilayering by day 5. Although no significant differences were detected in terms of total protein content and alkaline phosphatase activity at days 11 and 17, Biosilicate® supported significantly larger areas of calcified matrix at day 17. The results indicate that full crystallization of bioactive glasses in a range of compositions of the system P₂O₅–Na₂O–CaO–SiO₂ may promote enhancement of in vitro bone-like tissue formation in an osteogenic cell culture system. © 2007 Wiley Periodicals, Inc. J Biomed Mater Res 82A: 545–557, 2007

Key words: bioactive glass; crystallization; glass-ceramic; bioactivity; osteogenesis; cell culture

INTRODUCTION

The bioactive glass Bioglass® 45S5 has been known for many years as the bioactive material with the highest bioactivity index, which is defined as the inverse of the time required for 50% of the surface of the material to be intimately bound to the bone. It has been demonstrated that Bioglass® 45S5 affects osteoblast activities that ultimately result in enhanced bone formation both in vitro and in vivo. Indeed, at least seven families of genes are upregulated when primary human osteoblasts are exposed to the ionic dissolution products of bioactive glasses, including genes that encode proteins associated with osteoblast proliferation and differentiation.3–5 Despite its beneficial effects on bone healing, the use of Bioglass® 45S5 and other bioactive glasses for bone engineering applications has been limited due to their relatively poor mechanical properties.6

In this context, the development of novel bioactive glass-ceramics is much needed. Glass-ceramics are materials obtained by controlled crystallization of certain glasses.7 Bioactive glass-ceramics have been developed to improve the mechanical performance of bioactive materials, including Cervital and A/W,8,9 or to introduce other interesting properties, such as the machineable glass-ceramic Bioverit.1 Although...
glass-ceramics may exhibit improved mechanical properties over glasses, the introduction of some crystalline phases may sharply decrease the bioactivity. The result is that the bioactivity indexes of the current commercial glass-ceramics are much lower than those of bioactive glasses.\textsuperscript{10} In spite of decreasing the kinetics of the apatite layer formation in simulated body fluid (SBF) K9, crystallization does not inhibit its development, even in fully-crystallized glass-ceramics.\textsuperscript{11,12} In addition, depending on the characteristics of the crystalline phase that is formed, crystallization provides a temporary good mechanical support.\textsuperscript{13} Regarding this important matter, our research group has developed special nucleation and growth thermal treatments to obtain a novel fully crystallized bioactive glass-ceramic of the quaternary $P_2O_5$–$Na_2O$–$CaO$–$SiO_2$ system (Biosilicate\textsuperscript{1}, patent application WO 2004/074199).\textsuperscript{14} Crystallinity significantly changes the fracture characteristics of the glass. Therefore, full crystallization of the material may lead to enhanced mechanical properties of the bulk material or less sharp and abrasive particles when the material is milled to obtain a powder.

The present study aimed to evaluate the surface characteristics and the bioactivity of Biosilicate\textsuperscript{1} and to compare to Bioglass\textsuperscript{1} 45S5 (gold standard) and Biosilicate\textsuperscript{1} parent glass. In addition, using calvaria-derived osteogenic cultures, the following key parameters of \textit{in vitro} osteogenesis were assessed: (1) cell morphology and cytoskeleton organization; (2) immunolocalization of the multifunctional proteins bone sialoprotein (BSP), fibronectin, and osteopontin (OPN); (3) growth curve and cell viability; (4) total protein content and alkaline phosphatase (ALP) activity, and (5) mineralized matrix formation. The results showed that Biosilicate\textsuperscript{1} exhibits a hydroxy-carbonateapatite (HCA) layer formation on its surface in SBF-K9 as early as 24 h, which is comparable to the class A bioactive glasses. Furthermore, the enhanced \textit{in vitro} bone-like matrix formation on Biosilicate\textsuperscript{1} suggests that such material is most likely an osteoproductive glass-ceramic. This may indicate that Biosilicate\textsuperscript{1} exhibits a much higher bioactivity level than the current commercial apatite-based glass-ceramics.\textsuperscript{10}

\section*{MATERIALS AND METHODS}

\subsection*{Sample preparation and surface characterization}

High purity silica and reagent grade calcium carbonate, sodium carbonate, and sodium phosphate were used to obtain glass compositions: Bioglass\textsuperscript{1} 45S5 and Biosilicate\textsuperscript{1} parent glass. Raw materials were weighed and mixed for 30 min in a polyethylene bottle. Premixed batches were melted in Pt crucible at a temperature range of 1250–1380$\textdegree$C for 3 h in an electric furnace (Rapid Temp 1710 BL, CM Furnaces, Bloomingfield, NJ) at the Vitreous Materials Laboratory of the Federal University of São Carlos (São Carlos, SP, Brazil). Samples were cast into a 10 mm $\times$ 30 mm cylindrical graphite mold. After annealing at 460$\textdegree$C for 5 h, 3 mm thick glass discs were obtained by cutting the cylinders in diamond-blade. To obtain the fully crystallized Biosilicate\textsuperscript{1} glass-ceramic, Biosilicate\textsuperscript{1} parent glass cylinders underwent cycles of thermal treatment to promote their crystallization. The first thermal cycles were performed at lower temperatures aimed to promote volumetric nucleation of crystals. Afterwards, the nucleated samples were submitted to thermal treatments above the glass transition temperature to lead to a fully crystallized material. The compositions and thermal treatment schedules for obtaining the Biosilicate\textsuperscript{1} glass-ceramic is described in detail in the patent application WO 2004/074199.\textsuperscript{14}

The Biosilicate\textsuperscript{1} cylinders were cut into 3 mm thick discs using a diamond-blade. Finally, 12 mm in diameter and 3 mm thick discs were polished with silicon carbide abrasive powder (1000 grit), immersed in isopropyl alcohol, and cleaned by sonication. The discs were then rinsed with and stored in isopropyl alcohol to avoid surface modification by moisture. For the cell culture experiments, the discs were sterilized in dry heat at 180$\textdegree$C for 2 h.

Surface characterization of the Biosilicate\textsuperscript{1} glass-ceramic and the control bioactive glasses was performed by conventional light microscopy, scanning electron microscopy (SEM), and Fourier transform infrared (FTIR) spectroscopy, as described below.

Light microscopy examinations were carried out to confirm the crystallinity of the fully crystallized Biosilicate\textsuperscript{1} and to characterize its microstructure. The discs were polished with silicon carbide powder up to grit 1000 and with a suspension of finer CeO$_2$. The polished surfaces were treated with 0.5\% HF for 5 s, rinsed with water, acetone, and air-dried. The samples were then examined under reflected light using a Leica DMRX light microscope (Leica, Bensheim, Germany), outfitted with a Sony CCD-IRIS digital camera (Sony, Tokyo, Japan).

To simulate the early changes in surface topography and chemistry that cells will interact with and to allow its proper characterization, discs with the same surface preparation used for cell cultures were immersed in supplemented culture medium (see composition described below) in the absence of cells with a ratio of surface area of material to the volume of solution ($R_{SA/V}$) = 1 cm$^{-1}$ at 37$\textdegree$C in a humidified atmosphere with 5\% CO$_2$. After 12, 24, and 72 h, the discs were briefly washed in distilled water, air-dried, and kept in a desiccator to avoid surface changes by humidity.

The surfaces of randomly-selected Biosilicate\textsuperscript{1} and control discs were examined using a Phillips XL 30 field emission gun scanning electron microscope (Philips, Eindhoven, the Netherlands) operated at 20 kV. Discs stored in isopropyl alcohol (time 0) and exposed to culture conditions in the absence of cells for 12, 24, and 72 h were glued in aluminum sample holders and had their surfaces sputtered with carbon. Micrographs were then taken at different magnifications and processed with the Adobe Photoshop software (version 7.0.1, Adobe Systems).
Bioactivity tests were performed to compare the in vitro bioactivity level of the samples in acellular solution using SBF-K9. This solution has all the ions found in the human blood plasma in a very similar concentration. Such an assay is universally used for testing bioactive materials, in which the samples rest in the solution usually with $R_{SA/VS} = 0.1 \, \text{cm}^{-1}$ and 36.7°C. The discs had their flat surface briefly grounded in water with silicon carbide paper 400 grit and then were immediately rinsed with acetone followed by isopropyl alcohol. After drying, they were immersed in the SBF-K9 solution in a polypropylene flask with impervious sealing.

The surface chemistry was assayed by FTIR spectroscopy in a PerkinElmer Spectrum GX (PerkinElmer Life and Analytical Sciences, Shelton, CT) in reflectance mode. Samples immersed in isopropyl alcohol (time 0) and exposed to the culture conditions in the absence of cells for 12, 24, and 72 h, and to SBF for 24 h were evaluated.

**Cell isolation and primary culture of osteogenic cells**

Osteogenic cells were isolated by sequential trypsin/collagenase digestion of calvarial bone from newborn (2–4 days) Wistar rats, as previously described. All animal procedures were in accordance with guidelines of the Animal Research Ethics Committee of the University of São Paulo. Cells were plated on discs placed in 24-well polystyrene plates at a cell density of 20,000 cells/well. The plated cells were grown for periods up to 17 days using Gibco α-Minimum Essential Medium with l-glutamine (Invitrogen, Carlsbad, CA) supplemented with 10% fetal bovine serum (Invitrogen), 7 mM β-glycerophosphate (Sigma, St. Louis, MO), 5 μg/mL ascorbic acid (Sigma), and 50 μg/mL gentamicin (Invitrogen), at 37°C in a humidified atmosphere with 5% CO₂. The culture medium was changed every 3 days. The progression of cultures was examined by phase contrast microscopy of cells grown on polystyrene.

**Growth curve and cell viability**

Cells grown for periods of 4, 7, and 11 days were enzymatically detached from the culture substrate using 1 mM EDTA, 1.3 mg/mL collagenase, and 0.25% trypsin solution (Gibco, Invitrogen). Total number of cells/well, and percentage of viable and nonviable cells were determined after Trypan blue (Sigma) staining using a hemacytometer (Hausser Scientific, Horsham, PA).

**Total protein content**

Total protein content was determined using a modification of the Lowry method. Briefly, proteins were extracted from each well with 0.1% sodium lauryl sulfate (Sigma) for 30 min and mixed 1:1 with Lowry solution (Sigma) for 20 min at room temperature (RT). The extract was diluted in Folin and Ciocalteau’s phenol reagent (Sigma) for 30 min and mixed 1:1 with Lowry solution (Sigma) for 20 min at room temperature (RT). The extract was diluted in Folin and Ciocalteau’s phenol reagent (Sigma) for 30 min at RT. Absorbance was measured at 680 nm using a spectrophotometer (Cecil CE3021, Cambridge, UK). The total protein content was calculated from a standard curve, and expressed as μg/mL.

**ALP activity**

ALP activity was assayed in the same lysates used for determining total protein content, as the release of thymolphthalein from thymolphthalein monophosphate using a commercial kit (Labtest Diagnóstica, MG, Brazil). Briefly, 50 μL of thymolphthalein monophosphate was mixed with 0.5 mL of 0.3M diethanolamine buffer (pH 10.1), and left for 2 min at 37°C. The solution was then added to 50 μL of the lysates obtained from each well for 10 min at 37°C. For color development, 2 mL of 0.09M Na₂CO₃ and 0.25M NaOH were added. After 30 min, absorbance was measured at 590 nm, and ALP activity was calculated from a standard curve using thymolphthalein to give a range from 0.012 to 0.4 μmol thymolphthalein/h/mL. Data were expressed as ALP activity normalized for total protein content. Some cultures were also stained with Fast red, as described elsewhere, for histochemical detection of ALP activity during the mineralization phase of the cultures.

**Indirect immunofluorescence for localization of noncollagenous bone matrix proteins**

At days 1, 3, 5, and 14, cells were fixed for 10 min at RT using 4% paraformaldehyde in 0.1M sodium phosphate buffer (PB), pH 7.2. After washing in PB, cultures were processed for immunofluorescence labeling. Briefly, they were permeabilized with 0.5% Triton X-100 in PB for 10 min, followed by blocking with 5% skimmed milk in PB for 30 min. Primary monoclonal antibodies to BSP (anti-BSP 1:200, WVID1-9C5, Developmental Studies Hybridoma Bank, Iowa City, IA), fibronectin (anti-FN 1:100, clone IST-3, Sigma, St. Louis, MO), and OPN (anti-OPN, 1:800, MPIIIIB1-1, Developmental Studies Hybridoma Bank) were used, followed by Alexa Fluor 594 (red fluorescence)-conjugated goat anti-mouse secondary antibody (1:200, Molecular Probes, Invitrogen, Eugene, OR) and Alexa Fluor 488 (green fluorescence)-conjugated phalloidin (1:200, Molecular Probes), as a marker of the actin cytoskeleton. Replacement of the primary monoclonal antibody with PB was used as control. All antibody incubations were performed in a humidified environment for 60 min at RT. Between each incubation step, the samples were washed in PB (3 × 5 min). Before mounting for microscope observation, samples were briefly washed with dH₂O and cell nuclei stained with 300 nM 4',6-diamidino-2-phenylindole, dihydrochloride (DAPI, Molecular Probes) for 5 min. Discs were placed face up on glass slides and covered with 12-mm-round glass coverslips (Fisher Scientific, Suwanee, GA) mounted with Prolong antifade (Molecular Probes). The samples were then examined under epifluorescence using a Leica DMLB light microscope (Leica), with N Plan (×10/0.25, ×20/0.40) and HCX PL Fluotar (×40/0.75, ×100/1.3) objectives, outfitted with a Leica DC 300F digital camera, 1.3 Megapixel CCD. The

acquired digital images were processed with Adobe Photoshop software (version 7.0.1, Adobe Systems).

**Mineralized bone-like nodule formation**

At day 17, cultures were fixed with 4% formaldehyde in PB (pH 7.2) for 2 h at RT. The samples were then washed in the same buffer, dehydrated in a graded series of alcohol, and stained with 2% Alizarin red (Sigma) (pH 4.2), for 8 min at RT. They were photographed with a high-resolution digital camera (Canon EOS Digital Rebel Camera, 6.3 Megapixel CMOS sensor, with a Canon EF 100 mm f/2.8 macro lens) and then also imaged by epifluorescence microscopy. The percentage of the disc area occupied by Alizarin red-stained nodules was determined using the software ImageJ, version 1.34 s (NIH, Bethesda, MD). The amount of calcified matrix was also blind scored by six independent observers, using a scale of absent (0), small (1), moderate (2), and large (3).

**Statistical analysis**

Where appropriate, comparisons were carried out using the nonparametric Kruskal–Wallis test for independent samples (level of significance: 5%). If the result of the Kruskal–Wallis test was “significant”, that is occurrence of at least one significant difference, the Fisher’s least significant difference multiple comparisons procedure, computed on ranks rather than data, was performed. The results described below are representative of three sets of primary cultures.

**RESULTS**

**Surface characterization**

Light microscopy revealed that no microstructural features were detected for both bioactive glass controls [Fig. 1(A,B)], whereas Biosilicate® exhibited a fully crystallized structure with average crystal size around 5 μm [Fig. 1(C)]. Prior to the cell culture experiments, SEM images showed that all surfaces stored in isopropyl alcohol were flat, exhibiting randomly distributed features created by the polishing and finishing procedures [Fig. 1(D–F)]. After 12 and 24 h under culture conditions in the absence of cells, no relevant changes in surface microtopography except cracks were detected for all surfaces. Most importantly, higher magnification revealed submicron-scale globular structures on Biosilicate® and on control bioactive glasses as early as 12 h [Fig. 1(G–I)].

Despite their different compositions, original samples of Bioglass® 45S5 glass and Biosilicate® parent glass exhibited very similar infrared spectra [Fig. 2(A)]. The only slight difference in the spectra of the control glasses was the shift of the vibrational bands of the Bioglass® 45S5 glass to a lower wavenumber. It is worthy of note that major changes were observed between the spectra of the Biosilicate® parent glass and the fully crystallized Biosilicate®. Figure 2(A) shows that the vibrational band due to Si–O stretch at 1090 cm⁻¹ becomes sharper, shifts to 1100 cm⁻¹, and splits into two other vibrational bands at 1140 and 1050 cm⁻¹. Similar changes occurred with the vibrational band at 505 cm⁻¹ due to Si–O–Si bend. In the crystalline Biosilicate®, the vibrational band at 505 cm⁻¹ splits to vibrational bands at 530 and 460 cm⁻¹. The sharp vibrational band at 620 cm⁻¹ of Biosilicate® is probably due to Si–O vibrations.

When submitted to the bioactivity test in SBF-K9 at 36.7°C with RS_{SA/VS} = 0.1 cm⁻¹, a well-developed layer of HCA formed on all surfaces after 24 h of exposure, detected by the vibrational bands at 600 and 560 cm⁻¹ [Fig. 2(B)].

Samples exposed to the cell culture conditions in the absence of cells for 12 h at 37°C with RS_{SA/VS} = 1 cm⁻¹ showed significant changes in the infrared spectra compared to the original surfaces [cf. in Fig. 2, (C) with (A)], with Biosilicate® and the control bioactive glasses exhibiting the same vibrational bands [Fig. 2(C)]. For samples exposed for 24 h to the same culture conditions, the main changes compared to the infrared spectra at 12 h are indicated by arrows in Figure 2(D). They indicate that the vibrational bands at 470 and 800 cm⁻¹ decrease and the vibrational band at 590 cm⁻¹ increases. The vibrational bands at 800 and 1175 cm⁻¹ by 12 h shift slightly to 780 and 1230 cm⁻¹ by 24 h, respectively. At 72 h, the composition of the calcium phosphate layer formed is very close to the ones formed at 12 and 24 h. The most significant changes in infrared spectra compared to 24 h are indicated by arrows in Figure 2(E). They indicate that the vibrational bands at 470 and 780 cm⁻¹ decrease further and the vibrational band at 590 cm⁻¹ increases.

**Cell culture experiments**

Growth analyses indicated that there were no significant differences in cell number at days 4 and 11, while more cells were adhered on Biosilicate® parent glass at day 7 (Table I). There was no difference in cell viability between Bioglass® 45S5, Biosilicate® parent glass, and Biosilicate® at all time points (Table I). At day 1, phalloidin staining revealed that adherent cells were spread and randomly distributed throughout the surfaces [Fig. 3(A–F)]. On control glass coverslips, cells exhibited fusiform or polygonal shapes [Fig. 3(A)]. Some cells showed typical features of directional cell movement, with leading and trailing edges [Fig. 3(D)]. On such surface, actin cytoskeleton was characterized by bundles of stress fibers...
throughout the cytoplasm [Fig. 3(D,G,H,K)]. Strikingly, at days 1 and 3, the majority of adherent cells on Biosilicate® and on control bioactive glasses exhibited typical features of actin disassembly, with disruption of stress fibers in varying degrees [Fig. 3(B,C,E,F,I,J,L–N)]. Because of that, cell outlines could not be readily assessed on bioactive surfaces. At day 5, while reassembly/rearrangement of actin stress fibers took place in areas of cell multilayering [Fig. 3(Q,R,T–V)], cells adhering directly to the substrate still exhibited evidences of actin cytoskeleton disruption [Fig. 3(Q,R,T,U)]. Except for glass coverslips, cells with morphological features suggestive of chondrocytic differentiation were occasionally observed in early multilayering nodule formation [Fig. 3(T,U), arrowheads]. Straightforward observation revealed that mitotic figures increased from day 1 to day 5.

At days 1–5, OPN labeling was mainly perinuclear and also detected as punctate deposits throughout the cytoplasm [Fig. 3(A–F,O–R)]. Extracellular OPN accumulations were detected in cultures grown on Biosilicate® [Fig. 3(C,F)], and only occasionally on Bioglass® 45S5 [Fig. 3(B)], in a much lesser extent. Such accumulations were frequently found in association with intensely immunoreactive cells that exhibited morphological features of directional cell movement [Fig. 3(B,C,F)]. No extracellular OPN was evident on Biosilicate® parent glass and on control bioactive glasses [Fig. 3(L)]. At day 3, FN labeling was predominantly localized extracellularly and associated with cell outlines in cultures grown on glass coverslips [Fig. 3(H)]. It is worthy of note that only rarely was FN labeling observed on Biosilicate® [Fig. 3(I)] and on control bioactive glasses [Fig. 3(L)]. At day 5 on all surfaces, cells associated with initial multilayered nodule formation exhibited cytoplasmic OPN labeling [Fig. 3(O–R)]. In addition, only on glass cover-
Figure 2. FTIR transmission spectra of Biosilicate® glass-ceramic, Bioglass® 45S5, and Biosilicate® parent glass surfaces stored in isopropyl alcohol at time 0 (A), exposed to SBF for 24 h (B), and to the culture conditions in the absence of cells for 12 (C), 24 (D), and 72 h (E). (F) The band assignments and their corresponding wavenumbers.
The results of the present study showed that (1) all osteogenic materials evaluated supported cell attachment and proliferation in vitro, (2) the presence of bone-like matrix formation occurred on all surfaces, and (3) although the osteoblastic cells were detected for the fully crystallized glass-ceramic (Biosilicate®), no significant differences were observed in cell attachment and proliferation between the control glasses Bioglass® 45S5, 17S, and Biosilicate® parent glass. The presence of bone-like matrix formation was evident on all surfaces, and the osteoblastic cells were detected in areas of cell multilayering. The red-stained areas contained osteocyte-like cells (Fig. 4(G–I)]. At day 17, for all surfaces, Alizarin red-stained areas exhibited numerous lacunae, which were mainly in areas of cell multilayering (Fig. 4(J)]. The Alizarin red-stained areas exhibited numerous lacunae, which were mainly in areas of cell multilayering (Fig. 4(J)]. The Alizarin red-stained areas were stained with Alizarin red at day 7 (not shown).

**DISCUSSION**

The results of the present study showed that (1) all osteogenic materials evaluated supported cell attachment and proliferation in vitro, (2) the presence of bone-like matrix formation occurred on all surfaces, and (3) although the osteoblastic cells were detected for the fully crystallized glass-ceramic (Biosilicate®), no significant differences were observed in cell attachment and proliferation between the control glasses Bioglass® 45S5, 17S, and Biosilicate® parent glass. The presence of bone-like matrix formation was evident on all surfaces, and the osteoblastic cells were detected in areas of cell multilayering. The red-stained areas contained osteocyte-like cells (Fig. 4(G–I)]. At day 17, for all surfaces, Alizarin red-stained areas exhibited numerous lacunae, which were mainly in areas of cell multilayering (Fig. 4(J)]. The Alizarin red-stained areas were stained with Alizarin red at day 7 (not shown).

**TABLE I**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Time Points (Days)</th>
<th>Bioglass® 45S5</th>
<th>Biosilicate® Parent Glass</th>
<th>Biosilicate®</th>
<th>Kruskal–Wallis Test</th>
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<tr>
<td>Total cell number ($\times10^4$)</td>
<td>4</td>
<td>5.8 ± 1.5 (4)</td>
<td>8.1 ± 2.4 (4)</td>
<td>7.2 ± 2.2 (4)</td>
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<tr>
<td></td>
<td>7</td>
<td>21.2 ± 1.9 (4)</td>
<td>35.9 ± 3.3 (4)</td>
<td>22.6 ± 2.6 (4)</td>
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<tr>
<td></td>
<td>11</td>
<td>8.9 ± 2.9 (4)</td>
<td>11.6 ± 3.7 (4)</td>
<td>11 ± 2.3 (4)</td>
<td>NS</td>
</tr>
<tr>
<td>Cell viability (%)</td>
<td>4</td>
<td>69.3 ± 7.9 (4)</td>
<td>77.9 ± 5.5 (4)</td>
<td>71 ± 8.2 (4)</td>
<td>NS</td>
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<td></td>
<td>7</td>
<td>81.9 ± 4.7 (4)</td>
<td>79.5 ± 2.2 (4)</td>
<td>78.8 ± 0.7 (4)</td>
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<td>73.3 ± 3.1 (4)</td>
<td>65.9 ± 14.7 (4)</td>
<td>71.6 ± 4.6 (4)</td>
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<td>Total protein content (µg/mL)</td>
<td>4</td>
<td>153.9 ± 14.5 (5)</td>
<td>140.4 ± 10.5 (5)</td>
<td>141.4 ± 9.6 (5)</td>
<td>NS</td>
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<td></td>
<td>11</td>
<td>19.5 ± 3.2 (5)</td>
<td>6.6 ± 3.3 (5)</td>
<td>5.7 ± 0.8 (5)</td>
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<td>ALP activity (µmol thymolphthalein/h/mg)</td>
<td>4</td>
<td>13.8 ± 5.4 (5)</td>
<td>19.5 ± 3.2 (5)</td>
<td>15.5 ± 5.9 (5)</td>
<td>NS</td>
</tr>
<tr>
<td>Alizarin red-stained areas (scores 0–3)</td>
<td>11</td>
<td>13.8 ± 5.4 (5)</td>
<td>19.5 ± 3.2 (5)</td>
<td>15.5 ± 5.9 (5)</td>
<td>NS</td>
</tr>
<tr>
<td>Alizarin red-stained areas (%)</td>
<td>17</td>
<td>13.8 ± 5.4 (5)</td>
<td>19.5 ± 3.2 (5)</td>
<td>15.5 ± 5.9 (5)</td>
<td>NS</td>
</tr>
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</table>

Data represent mean values ± standard deviation (n). NS, nonsignificant ($p > 0.05$); S, significant ($p < 0.05$). Multiple comparisons procedure: Bioglass® 45S5 = Biosilicate® ($p > 0.05$); Biosilicate® parent glass > Biosilicate® ($p < 0.01$); Bioglass® 45S5 < Biosilicate® parent glass ($p < 0.05$).
Figure 3. Fluorescence labeling preparations of osteogenic cultures grown on Biosilicate® glass-ceramic (C, F, I, J, N, R, and V), control bioactive glasses Bioglass® 45S5 (B, L, P, and T), and Biosilicate® parent glass (E, M, Q, and U), and control glass coverslips (A, D, G, H, K, O, and S) at days 1 (A–F), 3 (G–N), and 5 (O–V). Green fluorescence (Alexa Fluor 488-conjugated phalloidin) reveals actin cytoskeleton (A–N and Q–V; H in pale white), while blue fluorescence (DAPI DNA stain) highlights cell nuclei (D and G–V). At day 1, the majority of cells on bioactive surfaces exhibit disassembly of actin cytoskeleton, whereas all cells on control glass coverslips show bundles of stress fibers throughout their cytoplasm (cf. B and C with A, and E and F with D). Osteopontin (OPN) labeling (red fluorescence) on control glass coverslips is mainly cytoplasmic, in a region suggestive of the Golgi apparatus (A and D) and in some granules. Noteworthy, on Biosilicate® (C and F) and in a less extent on Bioglass® 45S5 (B), OPN labeling is also characterized by extracellular deposits adjacent to cells exhibiting strong cytoplasmic labeling and typical morphology of directional cell movement. At day 3, almost all cells on bioactive surfaces exhibit a significant decrease in actin labeling, suggestive of disruption of stress fibers, while cells on glass coverslips appear more spread, showing bundles of stress fibers (cf. I with G, and L–N with K). Fibronectin (FN) labeling (red fluorescence) is predominantly extracellular and associated with cell outlines for cultures grown on glass coverslips (H). Conversely, FN labeling is only rarely detected in cultures grown on bioactive surfaces (I and L). At day 5, reassembly of actin cytoskeleton takes place in areas of cell multilayering on all bioactive surfaces (Q, R, and T–V). OPN labeling is detected on all surfaces (O–R), whereas bone sialoprotein (BSP) labeling (red fluorescence) is only detected on glass coverslips (cf. T–V with S). Focal areas of group of cells exhibiting typical aspects of chondrocytic differentiation are observed on control bioactive surfaces (T and U, arrowheads) and on Biosilicate®. [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
Figure 4. Epifluorescence of osteogenic cultures grown on Biosilicate® glass-ceramic (C, F, I, and L), on control bioactive glasses Bioglass® 45S5 (A, D, G, and J), and Biosilicate® parent glass (B, E, H, and K) at days 11 (A–C), 14 (D–I), and 17 (J–L). At day 11, nodular areas on all surfaces exhibit ALP activity, as revealed by Fast red staining (A–C). At day 14, BSP labeling (red fluorescence) is predominantly detected within the cells in nodular areas on Biosilicate® (F) and on control bioactive glasses (D and E). In areas of cell monolayer, large amounts of apoptotic bodies are observed on all bioactive surfaces (G–I). At day 17, Alizarin red-stained areas exhibit lacunae containing osteocyte-like cells (J). Noteworthy, larger areas of calcified matrix are observed on Biosilicate® (cf. L with K). Green fluorescence (Alexa Fluor 488-conjugated phalloidin) reveals actin cytoskeleton (D–I), while blue fluorescence (DAPI DNA stain) highlights cell nuclei (D–J). [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
It has been generally agreed that surface topography and chemistry are key factors to control tissue-implant interactions. In the present study, surface topography was qualitatively evaluated by light microscopy (reflected light) and SEM, whereas surface chemistry was analyzed by FTIR. While only Biosilicate\textsuperscript{1} exhibited a fully crystallized surface, no relevant changes in topographic features due to surface preparation were detected for all surfaces stored in isopropyl alcohol. Concerning chemical composition, the Biosilicate\textsuperscript{1} glass-ceramic, the control bioactive glasses Bioglass\textsuperscript{1} 45S5, and Biosilicate\textsuperscript{1} parent glass belong to the quaternary P\textsubscript{2}O\textsubscript{5}–Na\textsubscript{2}O–CaO–SiO\textsubscript{2} system. Although no major changes in terms of FTIR spectra were detected between both glass controls, the vibrational bands of Bioglass\textsuperscript{1} 45S5 was slightly shifted to smaller wavenumbers. Indeed, the increase of the modifiers Na\textsuperscript{+} and Ca\textsuperscript{2+} causes a shift of the vibrational bands of SiO\textsubscript{2} towards smaller wavenumbers, resulting from the depolimerization of the silicate framework of the glass. These shifts have already been demonstrated for other glass systems\textsuperscript{24} and support the fact that Biosilicate\textsuperscript{1} parent glass presents reduced concentration of glass modifiers compared to Bioglass\textsuperscript{1} 45S5. The FTIR spectrum for Biosilicate\textsuperscript{1} was significantly different from both control glasses. In glasses above the glass transition temperature, the covalent bond angles have some freedom to vary in a small range within the random network. The resulting infrared spectrum of the material is a sum of the contributions of the vibrations in this range. Conversely, in crystallized materials the bonding angles and lengths are well defined, which account for the original vibrational bands of the glass to split, shift, and sharpen in the spectrum of the crystallized material, as observed for Biosilicate\textsuperscript{1}.

The bioactivity of Biosilicate\textsuperscript{1} and of the control Bioglass\textsuperscript{1} 45S5, and Biosilicate\textsuperscript{1} parent glass (i.e., the bone-bonding ability of such materials) was assayed according to the ability of HCA to form on their surfaces in SBF with ion concentrations nearly equal to those of human blood plasma.\textsuperscript{16} FTIR revealed that all materials developed a HCA layer formation after 24 h in SBF-K9 at 36.7°C with $R_{SA/VS} = 0.1$ cm$^{-1}$ and that full crystallization did not alter such layer formation. However, when the materials were exposed to the culture conditions in the absence of cells, with $R_{SA/VS} = 1$ cm$^{-1}$, only silica-gel and calcium phosphate deposits with no crystallization were detected at 12, 24, and 72 h, with a progressive reduction of silica-gel and increase of the phosphate-related vibrational bands. Such results were supported by the SEM observation of a globular surface pattern at the submicron scale on all surfaces as early as 12 h, suggestive of precipitated amorphous calcium-phosphate onto the silica-gel layer, representing the in vitro surface reaction stages 3 and 4 of class A bioactive materials.\textsuperscript{2,3,25} That the silica-gel layer was formed was revealed by both SEM and FTIR analyses, by the presence of surface cracks due to the shrinkage of the fragile silica-gel layer during fast drying of the samples and the vibrational bands at 470, 800, and 870 cm$^{-1}$, respectively. The differences in the kinetics of the surface reactions for samples exposed to the culture medium compared to the ones exposed to SBF-K9 could be attributed not only to the solution itself, with diverse ionic concentrations and the presence of other substances and pH conditions, but also to variations in the ratio $R_{SA/VS}$. The latter may affect the pH of the solution, the rate of dissolution of the material, the saturation of the solution with the released ions, and

Figure 5. Macroscopic images of osteogenic cultures grown on Biosilicate\textsuperscript{1} glass-ceramic, on control bioactive glasses Bioglass\textsuperscript{1} 45S5, and Biosilicate\textsuperscript{1} parent glass, stained with Alizarin red at day 17. Biosilicate\textsuperscript{1} supports larger amounts of bone-like matrix formation (yellowish areas in a reddish background). [Color figure can be viewed in the online issue, which is available at www.interscience.wiley.com.]
the concentration of ions necessary to the HCA formation by total consumption. Finally, the possibility that molecules in the culture medium could inhibit crystal nucleation and HCA layer formation should not be ruled out. Irrespective of the exact mechanism, the slower surface reactions under the culture conditions would result in interactions of the cells with the reactive surface over a longer period of time.

The in vitro five-stage surface reactions of class A bioactive materials have been described to alter the gene expression profile of osteoblast lineage cells through the release of ionic products and the generation of an alkaline pH at the material surface, promoting osteoblast differentiation and function. In addition, surface reactions most likely affect early interactions of cells with matrix/serum proteins and substrate topography and chemistry, such as cell adhesion, spreading, and protein adsorption, which have also been demonstrated to influence interfacial tissue formation. In the present study, all bioactive surfaces affected actin cytoskeleton assembly as early as 24 h post-plating, resulting in disruption of actin stress fibers with a significant reduction in phalloidin labeling. Remarkably, cell-mediated fibronectin fibrillogenesis was also significantly reduced on such surfaces, compared to control glass coverslips, where cells exhibited well-developed actin stress fibers and an associated fibronectin fibrillar matrix. It is generally agreed that surface effects on cells are often mediated through integrins that bind the Arg–Gly–Asp (RGD) sequence of cell adhesion-associated serum/matrix proteins. Moreover, it has been demonstrated that the assembly of F-actin depends on integrin binding to RGD sequence and focal adhesion assembly, and that the tension generated by the actin cytoskeleton contributes to the assembly of a fibronectin fibrillar matrix. This could explain the differences observed in fibronectin labeling between bioactive and bioinert surfaces. Actin disassembly has already been demonstrated in cells grown on smooth and rough 45S5 monoliths at early time points, and the maintenance of such phenotype on rough surfaces has been associated with eventual enhanced bone nodule formation. Because the dynamics in actin assembly/disassembly were similar for Biosilicate and control bioactive glasses, such biologic process could not explain the differences noticed between surfaces in terms of amount of bone formation. In our study, reassembly of stress fibers took place in areas of initial cell multilayering by day 5 on all bioactive surfaces. In addition, F-actin-associated fibronectin matrix assembly also took place in such areas (not shown), consistent with the fact that fibronectin is essential for osteoblast differentiation and bone-like nodule formation in vitro.

Biomaterial surfaces, including those of bioactive glasses, have been coated with RGD peptides aiming to improve initial cell-substrate interactions. Early matrix deposition of OPN, a RGD-containing matrix-cellular protein, among other serum/matrix molecules, has been considered a key event in the development of the bone–material interface. In the present study, initial secretion of OPN took place mainly for cultures grown on Biosilicate and was associated with cells exhibiting typical morphological aspects of directional cell movement. Enhanced extracellular accumulations of OPN has been observed at early time points in calvaria-derived osteogenic cultures grown on bioactive nanostructured titanium surfaces, which eventually support increased bone-like nodule formation compared to machined surfaces. Such biologic event should be taken into consideration in strategies to design novel biomaterial surfaces modified with bioactive peptides/prteins with cell adhesion capacity.

Parameters to evaluate quantitatively the growth and differentiation phases of the primary osteogenic cultures did not show any relevant differences between the bioactive materials tested. Noteworthy, despite such results, the total area of bone-like formation was enhanced for Biosilicate, as detected histochemically. Because the pattern of matrix mineralization prevented accurate quantification of calcified areas, bone-like formation at day 17 was evaluated by two different semiquantitative methods. Both analyses by scores and percentage of surface area stained with Alizarin red revealed significantly larger areas of calcified matrix on Biosilicate than on control bioactive glasses. Trypsin/collagenase digestion of newborn rat calvarial bone allows the isolation of a mixed, heterogeneous cell population composed of osteoprogenitors, preosteoblasts, differentiated osteoblasts, osteocytes, and fibroblasts. Under the osteogenic conditions used in this study, calvaria-derived primary cultures generate woven bone-like nodules, which have been demonstrated to derive from the clonal expansion of osteoprogenitors and their entry into the osteoblast differentiation sequence. The role of isolated active osteoblasts in the process of matrix mineralization in vitro is still unclear. The rate of bone formation will mostly be determined by the rate of replication of osteoprogenitors, the number of active osteoblasts, and their life-span. Additionally, the enhanced production of bone-like matrix by osteogenic cells grown on Biosilicate could result from the selective recruitment of progenitor cells from the mixed cell population obtained following enzymatic digestion of calvarial bone and the stimulation of osteoblast activity by ionic products distinctively released from Biosilicate into the culture medium. For all bioactive surfaces, areas of fibroblastic cell monolayer with no microscopic signs of matrix mineralization.
exhibited large amounts of apoptotic cells, which has also been described for Bioglass® 45S5.3

Bioglass® 45S5 and Biosilicate® parent glass have a homogeneous glass matrix and the concentration of each chemical element is supposed to be uniform all over the material, thus the dissolution rate is the same all over the surface. On the other hand, during the crystallization process of Biosilicate®, selective attachment of atoms on the crystal growth front may occur, resulting in the production of a chemical gradient from the crystal centre to its border. Indeed, during crystallization of glasses of the Na2O–CaO–SiO2 system having compositions close to the stoichiometric Na2O–2CaO–3SiO2, Fokin et al.49,50 provided strong evidence for a continuous variation in both glass and crystal compositions. Partially crystallized samples show Na-enriched crystals, compared to the glassy matrix, while the glassy matrix exhibit a slightly higher concentration of Ca. The authors attributed the occurrence of such phenomenon to the measured variations of crystal growth velocity, nucleation rate, crystal lattice parameters, and other properties with increasing degree of crystallization. Similarly, the centre of each crystal in Biosilicate® could be enriched in Na, while its border would be depleted. The concentration of Ca would then vary in the opposite direction. In the first step of HCA formation on bioactive glasses of this family, there is an exchange of Na+ of the glass surface with H+ of the solution. If the above described compositional variations really occur, this exchange reaction would then take place near the crystal border with a slower rate compared to the parent glass. Therefore, the local pH increase would be higher near each crystal centre than on its border, causing the breakage of Si—O—Si bonds and release of Si(OH)4 in the solution at a higher rate in the central region. Thus, despite their identical (average) chemical composition, Biosilicate® and its parent glass may have different kinetics of dissolution depending on each specific region of the interface with bone tissue. Even if the overall surface concentration of Na in both materials is the same, primary calvarial cells could be sensitive to the Na enriched surface spots and their local pH, which would catalyze the release of ionic products in the solution that are known to regulate osteoblast differentiation and function.3–5 Such phenomenon gives a preliminary explanation for the different results observed on bone-like tissue formation for the fully crystallized Biosilicate® glass-ceramic compared to the control Bioglass® 45S5 and Biosilicate® parent glass surfaces.

CONCLUSIONS

We have demonstrated that the strategy used to improve the mechanical performance of the Biosilicate® parent glass by crystallization may also alter other important properties of the material, such as the dissolution rate. Although no major differences between all bioactive materials tested have been observed during the growth and differentiation phases of primary osteogenic cultures, the fully crystallized Biosilicate® glass-ceramic supported a significant enhancement of calcified tissue areas. The remarkable early changes in actin cytoskeleton and fibronectin matrix assemblies are most likely determined by the dynamic changes that take place on the reactive bioactive glass/glass-ceramic surfaces, and therefore could not explain the difference between Biosilicate® and the control bioactive glasses in terms of amount of bone-like tissue formation. The results presented herein open the doors for the development of a novel bioactive scaffold for bone tissue engineering applications.

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References
IN VITRO BONE FORMATION ON A HIGHLY BIOACTIVE GLASS-CERAMIC


