Development of bifunctional oriented bioactive glass/poly(lactic acid) composite scaffolds to control osteoblast alignment and proliferation

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Received 3 August 2018; revised 12 October 2018; accepted 6 November 2018
Published online 6 February 2019 in Wiley Online Library (wileyonlinelibrary.com). DOI: 10.1002/jbm.a.36619

Abstract: During the bone regeneration process, the anisotropic microstructure of bone tissue (bone quality) recovers much later than bone mass (bone quantity), resulting in severe mechanical dysfunction in the bone. Hence, restoration of bone microstructure in parallel with bone mass is necessary for ideal bone tissue regeneration; for this, development of advanced bifunctional biomaterials, which control both the quality and quantity in regenerated bone, is required. We developed novel oriented bioactive glass/poly(lactic acid) composite scaffolds by introducing an effective methodology for controlling cell alignment and proliferation, which play important roles for achieving bone anisotropy and bone mass, respectively. Our strategy is to manipulate the cell alignment and proliferation by the morphological control of the scaffolds in combination with controlled ion release from bioactive glasses. We quantitatively controlled the morphology of fibers containing bioactive glasses by electrospinning, which successfully induced cell alignment along the fibers. Also, the substitution of CaO in Bioglass®(45S5) with MgO and SrO improved osteoblast proliferation, indicating that dissolved Mg2+ and Sr2+ ions promoted cell adhesion and proliferation. Our results indicate that the fibers containing developed in this work are candidates for the scaffolds to bone tissue regeneration that enable recovery of both bone quality and bone quantity. © 2019 The Authors. Journal of Biomedical Materials Research Part A Published By Wiley Periodicals, Inc. J Biomed Mater Res Part A: 107A:1031–1041, 2019.

Key Words: biomaterial, bioactive glass, bone anisotropy, electrospinning, bone quality

How to cite this article: Lee S, Matsugaki A, Kasuga T, Nakano T. 2019. Development of bifunctional oriented bioactive glass/poly(lactic acid) composite scaffolds to control osteoblast alignment and proliferation. J Biomed Mater Res Part A 2019: 107A:1031–1041.

INTRODUCTION
Bone is a highly calcified tissue consisting of collagen fibrils and biological apatite (BAp) with several hierarchical levels from nano to microscale.1 Importantly, the multiscale structure of bone tissue exhibits highly anisotropic properties associated with collagen fibril orientation and the direction of the c-axis of BAp crystals.2,3 The anisotropic microstructure of bone tissue is one of the most important “bone quality” indices, which mainly governs the mechanical performance of bone tissue rather than bone mass (“bone quantity”).4 During the bone regeneration process, the recovery of anisotropic bone microstructure is significantly delayed compared to the bone mineral density (BMD) restoration, which induces severe mechanical dysfunction.4 The development of bifunctional biomaterials with controllable “bone quality” and “bone quantity” is therefore necessary for the recovery of highly ordered healthy bone tissue. Control of cell alignment is a valuable strategy for constructing anisotropic bone matrices; collagen/apatite matrix alignment depends on the osteoblast orientation.5 Moreover, the degree of BAp c-axis orientation shows a dependence on the directional distribution of osteoblasts.6 Accordingly, directional and quantitative control of osteoblasts can be determinative for achieving satisfactory bone tissue with both “quality” and “quantity.”

Bioglass® introduced the concept of bioactive materials; chemical cues from the material indicate enhanced metabolism and accelerated healing of damaged bone.7,8 Xynos et al. reported that the dissolved silicate, calcium, and orthophosphate ions from Bioglass® stimulated human osteoblast proliferation by increasing the production of insulin-like growth factor II (IGF-II).9–11 Magnesium ions promote cell adhesion, proliferation, differentiation, and subsequent mineralization.12–15 The expression of various integrin family members, which are a class of adhesion proteins, was increased by Mg2+ ions.13 Strontium ions have several effects on the stimulation of osteoblast proliferation and differentiation and the inhibition of

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Contract grant sponsor: Japan Society for the Promotion of Science; contract grant number: JP16K14403, JP18H03844, JP18H05254, and JP17H06224
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preosteoclast differentiation.\textsuperscript{16–18} Sr\textsuperscript{2+} ions increase the mRNA levels of c-fos and egr-1, which are involved in cell proliferation.\textsuperscript{16} They also promote the metabolism of osteoblasts due to activation of calcium-sensing receptors,\textsuperscript{16,19} and increase alkaline phosphatase (ALP) activity,\textsuperscript{20} which is a marker for osteoblast differentiation. Osteoprotegerin (OPG) was upregulated by Sr\textsuperscript{2+} ions and accompanied the downregulation of receptor activators of nuclear factor kappa B (RANK) ligand expression, which involve differentiation of pre-osteoclasts.\textsuperscript{18}

Electrospinning is a useful method for fabricating fibrous scaffolds, which can be applied to biomimetic templates for cell adhesion, proliferation, differentiation, and mineralization of damaged tissue. Obata et al. reported the use of poly (\textalpha;L-lactic acid) (PLLA) micro-fibermats containing silicon-doped vaterite for guided bone regeneration (GBR) membranes.\textsuperscript{21} Proliferation of osteoblasts on the fibermats was improved by dissolved silicate ions. \textit{In vivo}, newly formed bone was observed over 4 weeks, and the defect was covered after 12 weeks. Fibroblasts (NIH3T3) on the oriented nanofiber scaffold were elongate and aligned parallel to the fibers, and their gene expression upregulated associated with actin production, action polymerization, and focal adhesion formation than random one.\textsuperscript{22} Human mesenchymal stem cells (hMSC) on PLLA aligned nano-fibermats were highly oriented in the direction of the collector, where the cells were stretched along the long axis of the nanofibers.\textsuperscript{23,24} Subsequently, the collagen fibril bundles produced by hMSC were aligned in the direction of the cell adhesion (i.e., the nano-fiber direction). Tujunen et al.\textsuperscript{25} reported that mouse osteoblast-like cells on PLLA/siloxane-doped vaterite aligned micro-fibermats were oriented in the fiber orientation direction and elongated along the microfiber.

The aim of this study is to create a novel bifunctional biomaterial for bone tissue regeneration, which achieve recovery of both bone quality (oriented bone microstructure) and bone quantity (bone mass). The oriented fibermats were prepared with bioactive glass/PLLA composites by electrospinning an anisotropic scaffold to control cell alignment. PLLA, which is the most widely used biodegradable polymer in the biomaterials field, was chosen for fabrication of the oriented fibermat. The glasses in the composites were prepared by substituting the CaO in Bioglass\textsuperscript{40} (45S5) with MgO and SrO to improve bone regeneration with the dissolved ions from the glasses. Herein, a fundamental investigation on the design of oriented bioactive glass/PLLA fibermats for biomedical applications is reported, evaluating their anisotropic morphology, ion-releasing ability, and cell behavior on the fibermats.

**MATERIALS AND METHODS**

**Preparation of the bioactive glasses**

Glasses with compositions of 46.1SiO\textsubscript{2}:24.4Na\textsubscript{2}O:26.9MgO:2.6P\textsubscript{2}O\textsubscript{5} (mol%, M = Ca, Mg, or Sr, denoted by BGM) were prepared by melt quenching. Glass batches were prepared by mixing SiO\textsubscript{2} (99.0%), Na\textsubscript{2}CO\textsubscript{3} (99.5%), CaCO\textsubscript{3} (99.5%), MgO (99.0%), SrCO\textsubscript{3} (98.0%), and NaH\textsubscript{2}PO\textsubscript{4} (99.0%). All the reagents were purchased from Kishida Chemical Co. The batches were melted in a platinum crucible at 1500°C for 30 min and quenched by pressing with two stainless steel plates. The glasses were examined using laser Raman spectroscopy in between 220 and 1300 cm\textsuperscript{-1} (NRS-5100, JASCO). The resulting glasses were pulverized using an automatic alumina mortar, and the powders were stored in a desiccator. The resulting powders were observed by field emission gun electron microscopy (SEM, JSM-6500, JEOL) with an accelerating voltage of 15 kV after coating the samples with an amorphous osmium layer using an osmium coater (Neoc CS, MeiwaFosisis Co. Ltd.). Particles diameter were measured using the ImageJ software (NIH).

**Preparation of the composite pellets**

BGM powders were mixed with PLLA (LACEA, molecular weight of 140 kDa, Mitsui Chemical) by a melt-blending method using a kneader (PBV-0.1, Irie Shokai) at 190°C for 10 min, resulting in BG\textsubscript{M}/PLLA composite pellets. The volume ratios of BG\textsubscript{M} powders in the composites were set to 10 and 30 vol.%. The volumes of PLLA and BG\textsubscript{M} powders were calculated from their density. The densities were measured by an Archimedes’ method using acetone and water as immersion fluid for BG\textsubscript{M} and PLLA, respectively, at 25°C (n = 3). Molecular weight distributions of the composites were determined by gel permeation chromatography (GPC, Prominance, Shimadzu) using a KP-806L column (Shodex). For detection, a Shimadzu refractometer RID-10A was used. Chloroform (99.7%, HPLC grade, Wako Pure Chemical) was used as the eluent flowing at 1 mL min\textsuperscript{-1} at 35°C. The composites were manually injected (20 μL) at a concentration varying between 10 and 15 mg mL\textsuperscript{-1}. Average molecular weights and distributions were determined against a linear polystyrene calibrant.

**Preparation of the oriented fibermats**

The oriented fibermats with the composites were prepared by an electrospinning method, and that with PLLA was prepared as a control for cell proliferation test. The composite pellets and PLLA were dissolved in chloroform (99.0%, Wako Pure Chemical) at 14 wt.% PLLA to prepare the solution for electrospinning. In our preliminary experiments, this ratio was found to be optimal for preparing the oriented fibermats. In case of 10 vol.% BGM composite, the pellet was dissolved in chloroform at 10 wt.% PLLA, since the 14 wt.% solution could not be ejected from the syringe. The viscosities of the prepared solutions were measured using a vibration-type viscometer (VM-10A-M, Sekonic Co.). Subsequently, the prepared solutions were loaded into a syringe pump (FP-1100, Melquest, Japan) with an 18 gauge syringe needle, which was set at a discharge rate of 0.15 mL min\textsuperscript{-1}. A high-voltage supply (HARb-40P0.75, Matsusada Precision Inc.) was used to apply voltages of 16 kV at the needle tip. The distance between the needle tip and the drum collector was maintained at 200 mm. The drum collector (diameter 30 mm) was rotated at 3000 rpm (4.7 m·s\textsuperscript{-1}). The obtained fibermats were denoted by BGM\textsubscript{x}, where BGM is sample code for the bioactive glass and x (x = 10 or 30) is the vol.% of BGM in the composite. The electrospinning was carried out at room temperature (approximately 25°C) and approximately 40% relative humidity.
**Morphology of the fibermats**

The morphology of the prepared fibermats was observed by SEM with an accelerating voltage of 3 kV after coating the samples with an amorphous osmium layer using an osmium coater. Fiber diameter and the angle (θ) between the fiber and collector rotation direction were measured using the ImageJ software (NIH).

**Ion-releasing behavior of the fibermats**

To characterize the ion-releasing behavior from the fibermats, samples of 14 mm diameter and 120–160 μm thickness were soaked in 10 mL of 50 mM Tris buffer solution (TBS, pH 7.40, 37°C) for 9 days. The concentrations of Si, P, Ca²⁺, Mg²⁺, and Sr²⁺ ions in the TBS were measured by inductively coupled plasma optical emission spectroscopy (ICP-OES, Agilent 720 ICP, Agilent Technologies). The fraction of weight released of various elements in TBS were calculated as follows:

\[
\text{Release percentage (\%) = } \frac{a \times V_{\text{solution}}}{M_{\text{sample}} \times W_{\text{glass}} \times \text{Frac}_a}
\]

where \(a\) is the concentration of the element of interest in mg L⁻¹, \(V_{\text{solution}}\) is the volume of soaked solution in L, \(\text{Frac}_a\) is the nominal weight fraction of the element in the glass, \(M_{\text{sample}}\) is weight of the sample in mg, and \(W_{\text{glass}}\) is the wt% of the glass in the sample. After soaking in TBS for 9 days, the fibermats were analyzed by X-ray diffraction (XRD, X-pert PRO, Phillips) using Cu Kα radiation.

**Osteoblast proliferation on the fibermats**

Fibermats with 8 mm diameter were prepared for osteoblast tests. A PLLA oriented fibermat was used for the control. The samples were soaked in 70% ethanol for 30 s and subsequently dried under UV light for 30 min for sterilization. The cells were cultured in alpha-minimum essential medium (α-MEM, containing 10% fetal bovine serum (FBS, Invitrogen)) and 1% normal goat serum (NGS) for 30 min to block nonspecific antibody binding sites. Subsequently, the cells were incubated with mouse monoclonal antibodies against vinculin (Sigma-Aldrich) at 4°C for 12 h. The cells were incubated with Alexa Fluor® 546-conjugated anti-mouse IgG (Invitrogen) and Alexa Fluor® 488-conjugated phalloidin (Invitrogen). Finally, the cells were washed and mounted in Fluoro-KEEPER Antifade Reagent with DAPI (Nacalai Tesque). Fluorescent images were taken using a fluorescence microscope (BZ-X700, Keyence). The cell orientation angle (θ) against the collector rotation direction was analyzed using the Cell Profiler software (Broad Institute Cambridge).

**Primary osteoblast isolation and culture on the fibermats**

Primary osteoblasts were isolated from newborn mouse calvariae as described in our previous report.²⁶ Briefly, calvariae from newborn C57BL/6 mice were excised under aseptic conditions, placed in ice-cold α-MEM, and then the fibrous tissues around the bone were gently removed. The calvariae were then subjected to a series of collagenase (Wako Pure Chemical)/trypsin (Nacalai Tesque) digestions at 37°C for 15 min each. The first two digests were discarded, since fibroblasts were mixed.²⁷ The supernatants of digests 3–5 were neutralized with α-MEM and pooled. The pooled solutions were filtered using a 100 μm mesh. The filtrates were centrifuged, and the resulting pellets were resuspended in α-MEM containing 10% FBS. The population of obtained cells was verified by real-time reverse transcription polymerase chain reaction (RT-PCR, Step-one, Applied Biosystems). The positive expression of typical osteoblastic markers, collagen type I, ALP, and bone sialoprotein were confirmed, indicating the successful isolation of osteoblastic cell population in the present method. Fibermats with 14 mm diameter were prepared. The samples were soaked in 70% ethanol for 30 s and subsequently dried under UV light for 30 min for sterilization. The samples were placed into 24 well plates (n = 4), and primary osteoblast cells were seeded by adding 1.0 mL of the culture medium containing cells at a concentration of 3 × 10⁴ cells mL⁻¹. The culture medium was replaced after 1 day.

**Fluorescence imaging of primary osteoblast**

Primary osteoblasts were cultivated for 3 days on the samples, and the cells were fixed with 4% formaldehyde in PBS for 20 min. After washing three times with PBS-0.05% Triton X-100 (PBST), the cells were incubated in PBST containing 1% normal goat serum (NGS) for 30 min to block nonspecific antibody binding sites. Subsequently, the cells were incubated with mouse monoclonal antibodies against vinculin (Sigma-Aldrich) at 4°C for 12 h. The cells were incubated with Alexa Fluor® 546-conjugated anti-mouse IgG (Invitrogen) and Alexa Fluor® 488-conjugated phalloidin (Invitrogen). Finally, the cells were washed and mounted in Fluoro-KEEPER Antifade Reagent with DAPI (Nacalai Tesque). Fluorescent images were taken using a fluorescence microscope (BZ-X700, Keyence). The cell orientation angle (θ) against the collector rotation direction was analyzed using the Cell Profiler software (Broad Institute Cambridge).

**Quantitative analysis for the degree of fiber and cell orientation**

To evaluate the degrees of fiber and cell arrangement, the orientation order parameters FD and CD were calculated, where FD and CD are the degrees of fiber and cell alignment, respectively.²⁸ This system was derived using a distribution function \(n(θ)\), which is defined as the number of measured fibers or cells at the angle θ. The expected value of the mean square cosine \(<\cos^2θ>\), FD, and CD were calculated as follows:

\[
<\cos^2θ> = \frac{\int_0^{\pi/2} \cos^2θ \cdot n(θ) \, dθ}{\int_0^{\pi/2} n(θ) \, dθ}
\]

\[
\text{FD or CD} = 2( <\cos^2θ> - 0.5 )
\]

FD and CD take values ranging from −1 (fiber or cell completely aligned perpendicularly to the collector rotation
direction), to 0 (fiber or cell oriented randomly), to 1 (fiber or cell completely aligned parallel to the collector rotation direction).

Statistical analysis
Statistical significance was assessed by one-way ANOVA, followed by Tukey’s post hoc test. A significance of $p < 0.05$ was required for rejection of the null hypothesis.

RESULTS
Glass and composite characterization
Densities of $\text{BG}_{\text{Mg}}$, $\text{BG}_{\text{Ca}}$, $\text{BG}_{\text{Sr}}$, and PLLA were 2.59, 2.72, 3.04, and 1.25 g/cm$^3$, respectively. Pulverized $\text{BG}_{\text{Mg}}$, $\text{BG}_{\text{Ca}}$, and $\text{BG}_{\text{Sr}}$ diameters were approximately 2.7, 3.0, and 2.8 μm, respectively, and their distribution were shown in Figure 1. The particle diameter distribution between $\text{BG}_{\text{Mg}}$ showed no significant difference. Laser Raman spectra of $\text{BG}_{\text{Mg}}$ are shown in Figure 2A. The following Raman bands corresponding to the silicate $\text{QSi}_n$ ($n = 0–3$) groups$^{29,30}$ and orthophosphate ($\text{QP}_0$) group$^{31,32}$ were observed: the symmetric stretching mode of $\text{QSi}_3$ (~1030 cm$^{-1}$), symmetric stretching mode of $\text{QSi}_2$ (~970 cm$^{-1}$), symmetric stretching mode of $\text{QSi}_1$ (~910 cm$^{-1}$), symmetric stretching mode of $\text{QSi}_0$ (~850 cm$^{-1}$), Si-O stretching linkages (~640 cm$^{-1}$), and symmetric stretching mode of the non-bridging oxygen in $\text{QP}_0$ (~950 cm$^{-1}$). $\text{BG}_{\text{Mg}}$ may contain low amount of $\text{QSi}_4$.

![FIGURE 2. (A) Raman spectra for $\text{BG}_{\text{Mg}}$ and (B) integrated portion of the $\text{QSi}_n$ groups in $\text{BG}_{\text{Mg}}$.](image)

![TABLE I. Molecular Weights and Polydispersity Indices ($M_w/M_n$) of $\text{BG}_{\text{Mg}}$/PLLA Composites with 10 or 30 vol.% of $\text{BG}_{\text{Mg}}$ and Viscosities of their Solutions with 14 wt.% of PLLA in Chloroform](table)
(< 2%), which simulated by molecular dynamics. However, the band corresponding to the asymmetric stretching of $Q_{Si}^n$ ($\sim 1160$ cm$^{-1}$) was not observed for BGM in this work. The spectra between 800 and 1200 cm$^{-1}$ were fitted with Gaussian functions, and integrated portions of the $Q_{Si}^n$ ($n = 0$–3) groups are shown Figure 2B. The integrated portion of $Q_{Si}^0$ for BGM was 2.4%, while those of BGCa and BGSr were 8.3 and 8.5%, respectively. The percentage of

FIGURE 3. SEM images of (A) BGMg10, (B) BGCa10, (C) BGSr10, (D) BGMg30, (E) BGCa30, and (F) BGSr30. Arrows indicate the collector rotation direction. Fiber orientation angle histograms for (G) BGMg10, (H) BGCa10, (I) BGSr10, (J) BGMg30, (K) BGCa30, and (L) BGSr30. Error bars represent the standard deviation.
non-bridging oxygen (NBO) in the silicate groups of BG were calculated using the following equation:

\[
NBO_{\text{BG}}(\%) = \frac{\sum_{n=0}^{3} \left(4 - n\right) \times [Q_{n}^{3+}]_{\text{BG}}}{4 \times \sum_{n=0}^{3} [Q_{n}^{3+}]_{\text{BG}}} \times 100
\]

where \( n \) is number of bridging oxygen in \( Q_{n}^{3+} \) group, and \([Q_{n}^{3+}]_{\text{BG}}\) is integrated portions of the \( Q_{n}^{3+} \) groups in BG, which shown in Figure 2B. The values of BG, BG, and BG were 35.7%, 42.7%, and 44.0%, respectively. The NBO percentage indicates that the oxygen was in a \( \text{SiO}_{4} \) tetrahedron, not connected to other \( \text{SiO}_{4} \) tetrahedra.

Number-average molecular weights (\( M_n \)), weight-average molecular weights (\( M_w \)), and polydispersity indices (PDI, \( M_w/M_n \)) of PLLA and the composites are shown in Table I. The viscosities of the solutions for electrospinning are also shown in Table I, where the solutions were prepared with concentrations of 14 wt.% of PLLA in chloroform. The 10 vol.% BG composite was dissolved in chloroform at 10 wt.% of PLLA for electrospinning, with a viscosity of 1.4 Pa s. The composites containing BG were found to have larger \( M_n \) values compared to those of the composites containing BG and BG. The composites containing 10 vol.% of glass powders had larger \( M_n \) values and solution viscosities, and smaller PDI values than those of the 30 vol.% samples.

**Morphology of the fiber mats**

SEM images and fiber orientation angle histograms of the fiber mats are shown in Figure 3. The fibers were aligned with the collector rotation direction (parallel to the yellow arrows), and the fiber orientation angles were distributed about a center of zero. The calculated FD values and diameters of the fiber mats are shown in Figure 4. The FD of BG was significantly larger than the others (\( p < 0.01 \)), whereas the FD values showed no significant differences between BG, BG, and BG. The fiber diameters of BG, BG, BG, BG, BG, BG, and BG were 6.6, 4.8, 4.9, 3.4, 6.9, and 3.3 \( \mu \)m, respectively. The diameters of the fibers containing 10 vol.% of BG were larger than those of 30 vol.% fibers. The FD and fiber diameter of the PLLA fiber mat were 0.97 and 9.6 \( \mu \)m, respectively.

**Ion-releasing behavior of the fiber mats**

Ion-releasing behaviors of BG in TBS are shown in Figure 5A–D. The released amount of Si, divalent (\( \text{Mg}^{2+}, \text{Ca}^{2+}, \) and \( \text{Sr}^{2+} \)), and Na ions showed increasing trends with increased soaking time. The P ion releasing behaviors of the samples were significantly different. The amount of P ions released for BG increased to almost 100% with increased soaking time, whereas that of BG increased to approximately 65%, and those of BG and BG were approximately 30%, irrespectively the soaking time. Notably, the P ion releasing behavior of BG decreased to 0% with increased soaking time. The amounts of divalent ions released for BG and BG were smaller than those of the other BG. XRD patterns of BG after soaking in TBS for 9 days are shown in Figure 5E. BG and BG showed XRD peaks corresponding to \( \text{Ca}_{10}\left(\text{PO}_{4}\right)_{6} (\text{OH})_{2} \) (HA, ICCD card: 74–0566) and \( \text{Sr}_{5}\left(\text{PO}_{4}\right)_{3} \cdot \text{OH} \) (Sr-HA, ICCD card: 33–1348), respectively. The XRD peak intensities of BG and BG were larger than those of BG and BG, respectively.

**Cell behavior on the fiber mats**

The cell numbers on BG are shown in Figure 6. BG showed significantly larger cell numbers than PLLA at all sampling times, and those of BG were significantly larger at 3 days. BG and BG showed significantly larger values than PLLA at 3rd and 1st day, respectively. Fluorescence images of the cells and cell orientation angle histograms are shown in Figure 7. The cells were aligned in the fiber oriented direction (i.e., the collector rotation direction), and the cell orientation angles were distributed about a center of zero. Calculated cell orientation degree (CD) values on
the fibers are shown in Figure 8. The CD values of BG_{Mg10}, BG_{Ca10}, and BG_{Sr10} were larger than those of BG_{Mg30}, BG_{Ca30}, and BG_{Sr30}, respectively; that is, the CD values of the fibers containing 10 vol.% of BG_M were larger than those of the 30 vol.% fibers.

**DISCUSSION**

The Raman band intensity of BG_{Mg} was smaller than those of BG_{Ca} and BG_{Sr}, which was also noted by Karakassides *et al.*\(^{31}\) and Morikawa *et al.*\(^{35}\) Raman bands corresponding to \(Q_{Si}^n\) and \(Q_p^0\) were red-shifted (moved to lower frequencies) in
which is the simplified Coulomb’s force of the ions in glass, is defined as follows:\textsuperscript{36,37}

\[ F = \frac{Z^+}{d^2} \left( \text{valance}/\text{Å}^2 \right) \quad (4) \]

where \( Z^+ \) is the ionic charge and \( d \) is the interatomic distance between the cation and oxygen. The \( F \) values of Mg, Ca, and Sr were 0.53 or 0.46 (four-fold or six-fold coordination), 0.33, and 0.28, respectively.\textsuperscript{36} The bonding strength decreased in the order Mg-O, Ca-O, and Sr-O, causing the observed differences in the silicate and phosphate stretching vibration frequencies.\textsuperscript{31,35} The NBO contents in \( \text{BG}_{\text{Mg}} \) was larger than \( \text{BG}_{\text{Ca}} \) and \( \text{BG}_{\text{Sr}} \). According to Dietzel, Mg can be classified by the intermediates, which can switch role network modifier and former.\textsuperscript{36,37} Watts et al. reported that Mg can enter a silicate network as \( \text{MgO}_4 \) tetrahedral units, which can act as a network former.\textsuperscript{38} Thus, Mg in \( \text{BG}_{\text{Mg}} \) acts as a network former to enter the silicate network; which means that Mg is less effective at breaking the silicate chains than Ca and Sr. Consequently, \( \text{BG}_{\text{Mg}} \) showed a smaller NBO content than those of \( \text{BG}_{\text{Ca}} \) and \( \text{BG}_{\text{Sr}} \).

The \( M_n \) of the composite pellets decreased from that of PLLA, since absorbed water in the glass powder break the PLLA polymer chain during the melt-blending process. The absorbed water in the glass powder was bonded to the NBO in the glass structure. Accordingly, \( \text{BG}_{\text{Mg}} \) contained less absorbed water than \( \text{BG}_{\text{Ca}} \) and \( \text{BG}_{\text{Sr}} \) did, since the NBO content of \( \text{BG}_{\text{Mg}} \) was less than that of the others. Thus, the \( M_n \) of the composite containing \( \text{BG}_{\text{Mg}} \) was larger than those of \( \text{BG}_{\text{Ca}} \) and \( \text{BG}_{\text{Sr}} \). Similarly, the composites containing 10 vol.% of the glass powder had larger \( M_n \) values than those of the 30 vol.% composites, because they contained less glass powder (i.e., containing less amount of NBO). Generally, when a polymer with a larger \( M_n \) is dissolved in a solvent, its viscosity will be higher than a solution of the same polymer with a smaller \( M_n \).\textsuperscript{39} Correspondingly, the viscosity of the solutions had larger values for \( \text{BG}_{\text{Mg}} \) composites than those of the \( \text{BG}_{\text{Ca}} \) and \( \text{BG}_{\text{Sr}} \) solutions. Also, the composites containing 10 vol.% of the glass powder showed larger viscosities than the 30 vol.% composites.

In electrospinning, polymer fibers are formed by the creation and elongation of an electrified fluid jet.\textsuperscript{40} The velocities of the jets measured using a high framerate video camera were in the range from 0.5 to 5 m s\(^{-1}\).\textsuperscript{40} A collector speed of 4.7 m s\(^{-1}\) was used in this work; accordingly, the fibers could be collected while elongating in the collector rotation direction. If the solution formed a stable jet during the electrosprinning conditions, the resulting fibers showed oriented morphologies. The composites containing 10 vol.% of \( \text{BG}_{\text{M}} \) formed stable jets during electrospinning, whereas the 30 vol.% composites formed branched fluid jets. Generally, jet instability induces a branched fluid jet;\textsuperscript{39} the composites with 30 vol.% \( \text{BG}_{\text{M}} \) formed unstable jets more easily than the 10 vol.% composites due to their larger PDI values and because they contained larger amounts of glass powder. Branched fluid jets can be induced by the random angle distribution of the fibers, since there is an angular difference from the primary jet. This may also lead to larger \( FD \) values for the fibrermats containing 10 vol.% of \( \text{BG}_{\text{M}} \) than those of the 30 vol.% composites. A branched jet forms by ejecting smaller jets from the surface of the primary jets, which have smaller diameters than the primary jet.\textsuperscript{39} Thus, \( \text{BG}_{\text{M}30} \), \( \text{BG}_{\text{Ca}30} \), and \( \text{BG}_{\text{Sr}30} \) had smaller fiber diameters and larger \( FD \) values than those of \( \text{BG}_{\text{M}10} \), \( \text{BG}_{\text{Ca}10} \), and \( \text{BG}_{\text{Sr}10} \), respectively.

The released amounts of divalent and P ions from \( \text{BG}_{\text{M}30} \) and \( \text{BG}_{\text{Sr}30} \) were smaller than that of \( \text{BG}_{\text{Ca}30} \), owing to precipitation of HA and Sr-HA, respectively. No XRD peaks were observed for \( \text{BG}_{\text{M}30} \) since the \( \text{Mg}^{2+} \) ions inhibited the precipitation of apatite.\textsuperscript{41} The released amounts of Si ions from \( \text{BG}_{\text{M}30} \) and \( \text{BG}_{\text{Sr}30} \) were smaller than those of the others, due to formation of an Si-OH gel layer,\textsuperscript{42–44} which induced apatite formation.\textsuperscript{45} The released amounts of P and divalent ions from \( \text{BG}_{\text{Ca}30} \) and \( \text{BG}_{\text{Sr}30} \) were smaller than those of \( \text{BG}_{\text{Ca}10} \) and \( \text{BG}_{\text{Sr}10} \), respectively.

The released amounts of divalent and P ions from \( \text{BG}_{\text{M}30} \) and \( \text{BG}_{\text{Sr}30} \) were significantly larger than those of the control after 1 and 3 days of culturing. This was caused by the dissolved \( \text{Mg}^{2+} \) ions from the \( \text{BG}_{\text{M}30} \), which improved cell adhesion and proliferation.\textsuperscript{12–15} \( \text{BG}_{\text{Sr}30} \) showed
a significantly larger cell number after 3 days of culturing compared to that of the control, since the dissolved Sr\(^{2+}\) ions from BGS\(_x\) improved proliferation.

In our previous report, primary osteoblasts on oriented collagen substrate were aligned parallel to the collagen fiber orientation.\(^6\) Cell produced collagen matrix oriented in the

**FIGURE 7.** Fluorescence images of osteoblasts cultured on (A) BGM\(_{10}\), (B) BG\(_{Ca10}\), (C) BGS\(_{10}\), (D) BGM\(_{30}\), (E) BG\(_{Ca30}\), and (F) BGS\(_{30}\). Arrows indicate the collector rotation direction. Green: F-actin, blue: nuclei, and red: vinculin. Cell orientation angle histograms for (G) BGM\(_{10}\), (H) BG\(_{Ca10}\), (I) BGS\(_{10}\), (J) BGM\(_{30}\), (K) BG\(_{Ca30}\), and (L) BGS\(_{30}\). Error bars represent the standard deviation.
direction of cellular elongation, and the c-axis of depositedapatite crystals showed preferential alignment along the
direction of the newly synthesized collagen fibers. Thus, control of the osteoblast alignment can construct the bone
tissue anisotropy depending on the alignment of the cells themselves: that is, strongly orientated osteoblasts can pro-
duce the anisotropic bone tissues. Sun et al. reported that the
cells adhered to a single fiber when the diameter of fiber
was larger than 10 μm, whereas the cells adhered to several
fibers and spread when the diameter of the fibers was
<10 μm. In this study, the osteoblasts adhered on the fab-
cricated microfibermats elongate their stress fiber along the
fibermats, because the dynamics of actin organization is
strictly regulated by the spatial geometry involving the scaffold curvature. In microfibermats, the fiber diameter with
>10 μm could helpful to improve cell alignment by inhibiting
cells spread between fibers. That is, the orientation degree of
cells adhered to a single fiber can be easily controlled by
controlling the morphology of the fibermats. BG_{Mg}10, BG_{Ca}10, and BG_{Sr}10 showed larger CD values than those of
BG_{Mg}30, BG_{Ca}30, and BG_{Sr}30, respectively, since the fiber diameters of the 10 vol% BG fibers were larger than those of the 30 vol% fibermats. Notably, the cells on
BG_{Mg}10 and BG_{Sr}10, whose fiber diameters were larger than 6 μm, adhered to some single fibers, resulting in CD values
that were larger than those of the other samples. These results indicate that we can manipulate the cell orientation and proliferation freely by controlling the fiber diameter and the ionic species released from the bioactive glasses.

**CONCLUSION**
A novel bifunctional biomaterial, which can control osteoblast orientation as well as cell proliferation, was established.

The oriented BG_{Mg}/PLLA composite fibermats enabled cell
alignment along the fibers and promoted cell proliferation
due to the released ions from the bioactive glasses. The degree of fiber orientation was successfully controlled by modulating the content of the bioactive glasses and fiber diameters. The cell proliferation was significantly upregulated by the release of Mg^{2+} and Sr^{2+} ions from the bioactive glasses. The cell ori-
entation was determined by the cell recognition of the fiber orientation and adherence along single fibers or the formation of cell branches protruding across multiple fibers. The obtained results not only indicated that the fabricated com-
posites could control the cytoskeletal arrangement of osteo-
blasts along the fiber direction, but also, the controlled release of ions successfully improved osteoblast proliferation. These findings can lead to the development of innovative multifunc-
tional biomaterials suitable for tissue regeneration treatments by the optimization of the structural properties of the scaffold and inorganic ion element release.

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**ACKNOWLEDGMENTS**
This work was supported by Grants-in-Aid for Scientific Research from the Japan Society for Promotion of Science (Grant Numbers JP16K14403, JP18H03844, JP18H05254, and JP17H06224).
