Research Article

Evaluation of the Osteoblast Behavior to PGA Textile Functionalized with RGD as a Scaffold for Bone Regeneration

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The new era of biomaterials for repairing bone tissue injury continues to be a challenge in bone tissue engineering. The fiber scaffolds allow for cellular interconnection and a microenvironment close to the bone extracellular matrix. The aim of this study was to evaluate the osteoblast behavior on a 3D textile of PGA (polyglycolic acid) fibers functionalized with the RGD (R: arginine; G: glycine; D: aspartic acid) peptide. The cell morphology, proliferation, and calcium phosphate deposition ability were evaluated on textiles at different time intervals under a confocal laser scanning microscope. The osteoblast viability ranged from 92% to 98%, and cell proliferation was higher in PGA-RGD than control PGA (uncoated). In addition, the osteoblast calcium phosphate deposition was significantly greater on PGA-RGD in osteogenic inductor medium (OIM) in contrast to controls without inducing factors. The PGA-RGD fibers supported proliferation and viability of osteoblast and stimulated bone osteogenesis and mineralization. These results support the adoption of this 3D polymeric textile as a scaffold for bone tissue engineering.

1. Introduction

The healing of damaged or diseased bone is still a challenge issue for orthopedic surgeon and craniofacial surgeon. To overcome reduced bone formation and healing, many have turned to regenerative medicine. Usually, autogenous bone grafting is the first choice for bridging the bone defect. The advantages are free of immunogenic response and completely biocompatible since it comes from patient himself. The disadvantages are limited donor source, donor site morbidities, and variable bone graft survival. Xenograft and allograft are alternative choices for the treatment. However, immunogenic reaction and inadequate bone regeneration due to incomplete resorption result in nonunion or pathologic fracture. Recently tissue engineering becomes a promising matter to improve bone defect reparation [1–4].

The tissue engineering approach is a promising strategy added in the field of bone regenerative medicine, which aims to generate new, cell-driven, functional tissues, rather than just to implant nonliving scaffolds. The manufacture of scaffolds mimicking the extracellular bone matrix, which is composed of collagen fibers, calcium, phosphorus, and other minerals, has not yet been resolved [5–7]. Scaffolds made from polymer fibers have many of the characteristics necessary for the adhesion, proliferation, and differentiation of mesenchymal stem cells and osteoblasts [8–11]. Cell behavior can be influenced by the topography of fiber scaffolds; the organization, alignment, and direction of the fibers allow cells to attach to multiple textile walls due to their close interconnectivity and larger surface-area [12–15].

One of the goals of bone tissue engineering is to reproduce the biological and physiological conditions of bone within the human body by mimicking the in vivo cellular microenvironment, which consists of multiple complex factors. The ideal scaffold for bone engineering should promote mineralization and support the new osteoid matrix to do
this; biomaterial surfaces are modified and functionalized with bioactive factors that stimulate and promote a cellular response [16].

To repair bone defects and regenerate bone, the induction of osteogenesis is key. It is also a key determining the capacity of osteoblasts to deposit extracellular matrix and bone mineralization. The new biomaterials replace not only the mechanical functions but also the biological characteristics of the bone tissue [12, 17, 18].

Previous work demonstrated that the biocompatibility of a 3D polymeric textile intended for use in tissue regeneration was tested by evaluating the viability, proliferation, and adhesion of dental pulp stem cells (DPSCs); morphological parameters were characterized using scanning electron microscopy (SEM), X-ray photoelectron spectroscopy (XPS), and X-ray microtomography (μCT) analyses. The textiles were functionalized with the RGD (R: arginine; G: glycine; D: aspartic acid) peptide, which is thought to be the principal peptide responsible for cell adhesion [19]. Because synthetic polymers do not have bioactive groups on their surfaces, they do not promote cellular adhesion and require surface functionalization [20]. The 3D polymeric textile has been shown to support cell viability; however, it is necessary to perform studies with cell lineages specific to the tissue that needs to be regenerated [19].

In this study the behavior of osteoblast on PGA 3D textile functionalized with RGD to determine their ability induced of calcium phosphate deposition by used as scaffold for bone regeneration was evaluated. Additionally, the capacity of the scaffolds to induce osteoid matrix synthesis followed directly by mineralization was examined. The proliferation of osteoblasts on the fibers was evaluated by [3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulphophenyl)-2H-tetrazolium] (MTS) assay, cell viability was evaluated with the LIVE/DEAD Viability/Cytotoxicity Kit assay (Invitrogen) according to manufacturer’s instructions. The cell viability was determined at 3, 9, and 12 days of cell culture with the in vitro LIVE/DEAD Viability/Cytotoxicity Kit assay (Invitrogen) according to manufacturer’s instructions. The cells were observed under a confocal laser scanning microscope (CLSM, Leica Model DMI4000B, Germany) using a laser beam emitted at 488 nm and 532 nm wavelengths. Images were processed and analyzed using the LASAF software (Leica, Germany). Adherent live and dead cells were counted manually based on 10 images (1 x 1 mm) per textile acquired randomly. The percent viability was calculated using the following formula: number of live cells/number of total cells at each time × 100.

2. Materials and Methods

2.1. Textile Fabrication and RGD Textile Functionalization. The textiles were fabricated and characterized as described in previous work. Briefly, the textiles were knitted by Centexbel (Verviers, Belgium) from resorbable thread made from polyglycolic acid (PGA) multifilaments. After cutting, cleaning, drying, and undergoing 2 hours of UV sterilization and generating textile samples of an overall dimension of 0.2 × 0.5 cm (h w d), the textiles were stored under vacuum at −20°C until future use. The RGD (R: arginine; G: glycine; D: aspartic acid, A8052 Sigma-Aldrich) was physically absorbed by the textiles by incubating them in a solution of RGD (1 mg/mL) plus 0.1 M PBS (phosphate saline buffer) for 6 h at 4°C. Next, the textiles were washed with PBS and dried for 1 h under sterile conditions. The μCT and XPS analyses were conducted to analyze the porosity and surface properties of the fabrics, respectively [19].

2.2. Osteoblast Culture. Human osteoblasts (hOB) were isolated from the maxillary tuberosity using the explant technique during third molar surgery. This study was approved by the Institutional Ethics Committee (UASLP CEIFE-032-012). The hOB were cultured in 75 cm² cell culture flasks containing Dulbecco’s Modified Eagle Media (DMEM, Sigma-Aldrich, St. Louis, USA) supplemented with 10% fetal bovine serum (FBS, Biosciences, USA) and antibiotic solution (streptomycin 100 μg/mL and penicillin 100 U/mL, Sigma-Aldrich). The cell cultures were incubated in a 100% humidified environment at 37°C in an environment of 95% O₂ and 5% CO₂. The hOB from passages 2-6 were used for all of the experimental procedures. For in vitro osteogenic differentiation, cells were cultured in an osteogenic inducer medium (OIM) consisting of complete DMEM medium supplemented with 50 μg/mL of ascorbic acid, 10 mM of b-Glycerophosphate, and 10⁻⁷ M of Dexamethasone (all from Sigma-Aldrich St. Louis, USA). During the experiment, the culture medium was changed twice per week.

2.3. Seeding on Textiles. The hOB were cultured in a 75 cm² cell culture flask upon reaching 90% confluence and then washed with PBS. The cell monolayer was detached using 1x Trypsin-EDTA (0.25%) solution (Gibco, Life Technologies, USA). The human osteoblastic cells were seeded at a concentration of 1 x 10⁴ cells/textile onto the upper surface of the PGA textiles and allowed to infiltrate into the textiles within Millicell EZ SLIDE 8-well glass slides (Millipore-USA). The samples evaluated in the study were PGA-RGD and PGA with and without OIM induction. Cells were cultured without agitation at 37°C in a humidified atmosphere under 5% CO₂. Conventional polystyrene 24-well culture plates were used as a control. The medium was changed twice per week [19].

2.4. Cell Viability. The cell viability was determined at 3, 9, and 12 days of cell culture with the in vitro LIVE/DEAD Viability/Cytotoxicity Kit assay (Invitrogen) according to manufacturer’s instructions. The cells were observed under a confocal laser scanning microscope (CLSM, Leica Model DMI4000B, Germany) using a laser beam emitted at 488 nm and 532 nm wavelengths. Images were processed and analyzed using the LASAF® software (Leica, Germany). Adherent live and dead cells were counted manually based on 10 images (1 x 1 mm) per textile acquired randomly. The percent viability was calculated using the following formula: number of live cells/number of total cells at each time × 100.

2.5. Cell Proliferation. Proliferation of hOB cells seeded onto PGA at a concentration of 1 x 10⁵ cells{textile was quantified at 3, 5, and 7 days of culture using the Cell Titer 96™ Solution Cell Proliferation Assay (Promega), also known as the MTS assay. This assay is based on the ability of mitochondrial dehydrogenases in living cells to oxidize the MTS tetrazolium compound forming a colored formazan product that is soluble in tissue culture medium. The concentration of colored product is directly proportional to the number of metabolically active cells. The hOB seeded at the indicated times onto PGA textile material and PGA textile material functionalized with RGD were washed with PBS, incubated with fresh culture medium containing 20 μL of the stock solution, and incubated for 3 h at 37°C. After incubation, the absorbance was quantified by spectrophotometry at 490 nm.
with a plate reader (Epoch, BioTek). During the experiment, the culture medium was exchanged for fresh medium every two days. All MTS experiments were conducted in triplicate and repeated at least three times.

2.6. Mineralization Analysis and Calcium Phosphate Deposition. An osteogenesis quantization assay kit (Millipore) was used to determine the degree of mineralization, which is related to calcium content, during osteogenesis of human osteoblasts cultured onto the PGA textile for 4, 14, and 21 days with and without OIM. This assay is based on the quantification of Alizarin Red Stain (ARS), which binds selectively to calcium phosphate deposits (mineralization nodules). After incubation, the PGA textiles were carefully washed with PBS, fixed with 10% formaldehyde for 1 h at room temperature, carefully washed with distilled H₂O three times (5–10 minutes), and then stained with ARS solution (40 mM) for 30 min at room temperature. After several washes with distilled H₂O to remove excess dye, the PGA textiles were examined under an optical microscope (Leica DMIL LED), and images were processed and analyzed using the LAS EZ software (Leica, Germany). After microscopic analysis of ARS staining, the dye was extracted from the PGA textile by treatment with 10% acetic acid for 30 min. After neutralization with 10% ammonium hydroxide, the optical density of ARS was measured at 405 nm with a spectrophotometer (Epoch, BioTek). The concentration of ARS was determined by correlating the absorbance of the experimental samples with a standard curve of known ARS dye concentrations.

2.7. Statistical Analysis. Significant differences between experimental groups were determined using the Kruskal-Wallis and Wilcoxon tests, with \( p < 0.05 \) considered significant. Data were analyzed using SigmaPlot Ver. 11.0 statistical software (SPSS Inc., Chicago, IL, USA).

3. Results

3.1. Cell Viability. The viability of human osteoblast cells isolated using the explant technique was evaluated by LIVE/DEAD Viability/Cytotoxicity Kit assay. The results show more than 92% viability with well-defined polygonal, spindle shaped cell morphology (Figures 1(a)–1(c)). The cell morphology was observed at different intervals throughout the viability assays (Figures 1(d)–1(f)). Our results showed that the viability of human osteoblasts cultured on PGA textile fibers functionalized with RGD peptide in the presence of OIM at 3, 9, and 12 days was significantly greater for all time intervals than viability of osteoblasts cultured on control PGA (Figures 2(a)–2(f)). However, no differences in biocompatibility were found when human osteoblasts were cultured onto PGA-RGD and PGA textile in normal complete medium.
Figure 2: Images taken using a CLSM on PGA textiles at 3, 9, and 12 days. Osteoblasts were stained using the LIVE/DEAD Viability/Cytotoxicity Kit assay (10x). PGA-RGD-OIM at 3 days (a), 9 days (b), and 12 days (c). PGA-OIM at 3 days (d), 9 days (e), and 12 days (f). PGA-RGD control at 3 days (g), 9 days (h), and 12 days (i); control PGA at 3 days (j), 9 days (k), and 12 days (l).
3.2. Cell Proliferation. Proliferation of osteoblasts cultured on RGD (Figures 2(c) and 2(f)). To the flattened morphology of osteoblasts cultured on PGA-control PGA textiles were round to oval in shape compared to the flattened morphology of human osteoblasts cultured on polymeric textile scaffolds of PGA-RGD-OIM and control PGA-OIM have an enhanced calcium phosphate deposition in the extracellular matrix compared with cultures without OIM (Figure 5).

The percent cell viabilities determined by image analysis of the PGA textiles ranged from 92% to 98%. Moreover, the morphology of human osteoblasts cultured on control PGA textiles were round to oval in shape compared to the flattened morphology of osteoblasts cultured on PGA-RGD (Figures 2(g) and 2(h)). The percent cell viabilities determined by image analysis of the PGA textiles ranged from 92% to 98%.
fiber densities and patterns; therefore, the growth patterns depend on cell-cell communication and cell alignment [14].

The cell morphologies observed under CLSM include elongated and polygonal morphology and the presence of flattened and cytoplasmic extensions between the multilamellar. Cell infiltration is possible due to the porosity and the morphological parameters of the fiber. Cell morphology is one way to determine if the substrate is suitable for the cells. The osteoblasts cultured on textiles had an elongated morphology with abundant cytoplasmic cell processes, filopodia, and lamellipodia, connected to the fiber matrix and to neighboring cells. These morphological characteristics are a key factor in many biological processes [23].

The proliferation assay demonstrated that all cells could proliferate after they adhered to the textiles. The fiber scaffold structures were shown to promote cell attachment;
depends on the biomaterial surface roughness, topography, and chemistry, and functionalization. The PGA textiles functionalized with RGD in inductive conditions tend to deposit calcium phosphate, which is an indicator of matrix formation. The osteoblast viability ranged from 92% to 98%, and cell proliferation was higher in PGA-RGD than control PGA. Our data showed that the adhesion and proliferation of osteoblasts cultured on PGA-RGD-OIM were significantly increased relative to control. In addition, the calcium phosphate deposition from osteoblasts was significantly greater on PGA-RGD-OIM. The PGA-RGD fibers supported proliferation and viability of osteoblast. These results suggest that this 3D polymeric textile can be employed as a scaffold for bone tissue regeneration. Future in vivo studies should be performed as well.

**Conflicts of Interest**

The authors declare that they have no conflicts of interest and any financial interest related to this study.

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**References**


**Figure 5:** Quantitative Alizarin Red Staining after 4, 14, and 21 days of cell culture. There were significant differences on PGA-RGD and PGA with osteogenic inductor medium at 4, 14, and 21 days *(p < 0.05).*

nevertheless, surface functionalization stimulated protein adsorption, which represents the first step in cell adhesion. Studies have provided evidence that synthetic fibers may exhibit certain properties that are comparable to natural collagen fibers; thus, the nanostructured architecture may be superior to the solid-walled architecture for promoting osteoblast differentiation and biomineralization [24–28].

Calcium phosphate deposition was evaluated by ARS staining, which is used to visually detect mineralization in bone tissue. In the textiles functionalized with RGD in the presence of OIM, the presence of ARS demonstrates calcium phosphate deposition throughout the multifilament polymeric textiles. In the case of control PGA without functionalization, but with osteoblasts cultured in OIM, lower amounts of ARS were observed. We supposed that RGD promotes the growth of osteoblasts and produces a greater amount of mineral nodules deposition on PGA-RGD. The growth rate of osteoblasts on control PGA textiles was slower than on functionalized textiles. This result may be related to the different degrees of cell spreading on these two surface types.

Studies have demonstrated that culturing osteoblast cells on nanofibers made of poly(L-lactic acid) results in much higher levels of alkaline phosphatase in cells than culturing on flat films in complete induction medium [29], while growth on polycaprolactone/gelatin electrospun scaffolds in medium with osteogenic supplementation promoted bone osteogenesis and mineralization [30]. In collagen/hydroxyapatite composite nanofibrous scaffolds, the mineralization reflects the functional activity of osteoblasts [31].

**5. Conclusions**

The interaction between osteoblasts and biomaterials depends on the biomaterial surface roughness, topography,


