Multifunctional PLLA-ceramic fiber membranes for bone regeneration applications

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Abstract
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Multifunctional PLLA-ceramic fiber membranes for bone regeneration applications

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Abstract:
A novel method to process electrospun poly(L-lactic acid) (PLLA) membranes incorporating glass reinforced hydroxyapatite granules (gHA) interspatially between the polymeric fibers is reported, thus increasing the surface area for cellular interactions. gHA granules (≤150 µm) electrospun together with the polymer solution, lead to an average fiber diameter of 550 ± 150 nm for pristine PLLA and 440 ± 170 nm for the composite samples. An increase of the overall porosity was observed, from 79 ± 3 % for the PLLA up to 88 ± 5 % for the hybrid samples, keeping material`s wettability and mechanical properties. Bone-bonding ability showed that both samples induced HA crystal nucleation, but with a distinct pattern of mineral deposition. gHA microcomposite allows a better F-actin cytoskeleton organization during the initial adhesion and spreading, favoring cell-fibers and cell-to-cell interactions and enhanced alkaline phosphatase activity, making them potential candidates for bone healing strategies.

Keywords: Biomaterial; Poly (lactic acid), Glass-reinforced hydroxyapatite, Bone grafts; Electrospun microcomposite fibers, Bone regeneration.
Introduction

People suffering from bone defects arising from trauma, tumor or bone diseases is effectively a reality in the present days and every year several million orthopedic procedures are performed [1]. At present, the standard treatments for bone repair involve, in general, autografts, where bone is transplanted from another part of the recipient's body, and allografts, where bone is transplanted from genetically non-identical members of the same species. These grafting solutions present several limitations, namely donor site scarcity, rejection, disease transmission, need of a separate incision for harvesting and post-operative morbidity [2, 3]. Therefore, there is a need to explore novel biomaterials in bone tissue engineering, so that the treatment strategy can be as quickly and efficient as possible.

In this context, tissue engineering has an important role to provide functional substitutes for the damaged tissues [4, 5]. Man-made manufacture substitutes, also known as scaffolds, capable of mimicking the natural tissue environment to successfully meet or perhaps surpass the original mechanical, structural, and functional properties are constantly being developed [4]. The conception of a scaffold with such features is a demanding task, bearing in mind that it should provide a transitional three dimensional support for cell migration, attachment and proliferation, as well as offer mechanical, biological and chemical cues to guide the cells towards tissue restoration [5]. To satisfy those requirements, electrospinning arises as a versatile technique to produce fibrous membranes with same morphology to the ones found in the extracellular matrix (ECM) [4]. Besides, the electrospun fibers with diameters ranging from micro to nanometer scale offer several advantages such as an extremely high surface-to-volume ratio, tunable porosity, malleability to conform to a wide variety of sizes and shapes and the ability to control the fiber composition [6]. Furthermore, scaffolds consisting of
electrospun fibers can be functionalized for enhanced cellular activities by incorporating bioactive compounds.

The current challenge in bone tissue engineering is to engineer bioartificial bone grafts mimicking the ECM, while allowing effective bone mineralization [1]. Until now, different biomaterials like bioactive ceramics and degradable polymers, have been developed to mimic the mechanical and biological properties required by bone tissue [7]. However, the electroactive properties of these materials, which are important in promoting electrical cues to cells, have been neglected in these advancements.

Electroactive polymers are an emergent technology for tissue and biomedical applications. Piezoelectric materials generate transient surface charges by tiny mechanical deformations of the material under mechanical solicitation and do not require additional energy sources or electrodes [8].

PLLA has generated great interest as one of the most promising biodegradable polymers due to its mechanical properties, thermoplastic processability and biodegradability, becoming highly attractive for biological and medical applications [3, 9]. Furthermore, the degradation products of poly(lactides) are nontoxic, which is fundamental for biomedical engineering applications. PLLA is also known for its piezoelectric properties that are in the core of technological applications as sensors [10] and actuators [11]. Moreover, PLLA presents a piezoelectric constant around 10 pC.N$^{-1}$, similar to the one in human bone, reported to be between 7 - 12 pC.N$^{-1}$ [12, 13].

Synthetic glass reinforced hydroxyapatite (gHA) was sintered in the presence of CaO-P$_2$O$_5$ based glass using a patented process [14]. HA - Ca$_{10}$(PO$_4$)$_6$(OH)$_2$ - is the major mineral constituent of the bone matrix, but its poor mechanical properties and low reabsorption rate by the organism, compromise the recovery of bone normal strength, causing the bone to remain brittle and easily prone to fractures [15, 16]. gHA particles
have a highly controlled chemical composition of $\beta$ and $\alpha$ tricalcium phosphates secondary phases known to accelerate biodegradation over HA [17, 18]. The content of ionic species (such fluoride and sodium, among others) along with its micro and macroporous structure uphold a positive effect in osteoconduction and osteointegration in bone regeneration [16, 19].

Bone is a hybrid system of HA and collagen type I fibers, that assembles in a complex and organized porous structure [20]. Electrospun fiber membranes with inorganic HA can mimic the ECM morphology, chemical composition and improves the scaffold mechanical properties [21]. Nevertheless, electrospinning of polymer matrix composites with inorganic fillers are mainly based on the incorporation of HA nanoparticles, which raises issues concerning the immobilization of the ceramic filler on the surface of the fibers. When nanoparticles are used, they often are immobilized in the core of the fiber, and the contact between the active HA filler and the cells is hindered [22]. To overcome this issue and to increase the exposed area of the ceramic to cells, new processing strategies are required.

In this work, we intended to produce hybrid microcomposite electrospun membranes using PLLA as a matrix and gHA micrometer particles as a filler material, and create a topography where the ceramic “islands” appear entrapped in a “sea” of polymeric fibers, increasing the amount of ceramic in direct contact with the cells and the surface-to-volume ratio of the microcomposite materials. The influence of the gHA in fiber average diameter, wettability and biocompatibility was assessed. The suitability of such composite membrane composition for bone healing and regeneration was explored.
Experimental:

gHA synthesis: Ceramic powder was synthetized as per the method described elsewhere [23, 24]. In this work, using standard milling and sieving techniques, gHA granules with particle size $\leq 150$ µm were obtained.

Electrospun membrane preparation: Poly(L-lactic acid) (PLLA, Purasorb PL18, $M_w = 217 – 225$ kDa) from Corbion (Netherlands) was dissolved in a DMF (Merck):DMC (Sigma-Aldrich) solution (3:7 v/v) to achieve a polymer concentration of 10 wt% of the solution. gHA microcomposite suspensions were prepared by adding the ceramic filler to the DMF:DMC (3:7 v/v) solvent mixture, followed by dispersing in an ultrasound bath (Bandelin, Model Sonorex Super RK106) for 6 h, to promote a good dispersion of the particles. Then, the polymer was added to the mixture and stirred at room temperature until complete dissolution. The concentration of filler related to polymer was 80 wt%.

The neat PLLA and gHA - PLLA solutions were placed in a commercial glass syringe fitted with a metallic needle with 0.5 mm inner diameter. Electrospinning was conducted at 1.25 kV.cm$^{-1}$ with a high voltage power supply from Gamma High Voltage. A syringe pump (from KDScientific) was used to feed the polymer solution into the needle tip at 0.5 mL.h$^{-1}$. The electrospun fibers were collected in ground collecting plate placed at 20 cm apart from the needle (random aligned fibers). All experiments were conducted at $21 \pm 2$ ºC and a relative humidity of $43 \pm 5$%.

Membranes Characterization: Electrospun fiber membranes were coated with a thin gold layer using a sputter coater (Polaron, SC502) and their morphology was analyzed using a scanning electron microscope (NanoSEM FEI Nova200, from FEI) with an accelerating voltage of 10 kV. An energy-dispersive X-ray (EDS) was used in
conjunction with SEM for elemental analysis of the deposited mineral crystals. Fibers
average diameter and its distribution was calculated over approximately 50 fibers using
the Image J software [25].
Contact angle measurements (sessile drop in dynamic mode) were performed at room
temperature in a Data Physics OCA20 device using ultrapure water as test liquid. A
sessile drop (3 µL) of ultrapure water (Milli-Q, Millipore) was deposited on the surface
of the samples and the contact angle was measured using a DataPhysics OCA 35
goniometer and SCA20 software. For the same sample, contact angles were measured
on 8 different spots and the result is expressed as their average and standard deviation.
XRD analysis was performed on powder samples of gHA and on PLLA/gHA fiber
membranes using a Phillips Analytical X-Ray model PW 1710, employing Cu Kα
monochromatic radiation (40 kV, 30 mA, Kα=1.541838 Å). Data was collected at room
temperature, for 2θ values ranging from 5 to 70º in increments of 0.02º and a counting
time of 2 s/step. Thermogravimetric analysis (TGA) was performed in a Q500 apparatus
from TAINstruments at heating rate scan of 20 ºC.min⁻¹, under a nitrogen atmosphere.
Mechanical properties were performed in a Shimadzu Universal Testing Machine (AG-
IS with a 10 N load cell) in tensile mode, at a strain rate of 0.5 mm.min⁻¹. Rectangular
stripes of 10x40 mm² were measured with a caliper (Mitutoyo) and a thickness of ~250
µm was measured with a DUALSCOPE® MPOR (Fischer). From the stress-strain data,
elasticity modulus was calculated in the linear zone, between 0 and 2 % of strain. The
values presented were calculated through the average and standard deviation of five
individual measurements performed for each sample.

Samples porosity: The porosity of PLLA and gHA-PLLA membranes was measured
using the pycnometer method [26]. First, a pycnometer filled with ethanol was weighted
and labeled as $W_1$. Then, the sample (with a weight of $W_{\text{sample}}$) was immersed in ethanol and, when it was saturated by the ethanol, additional ethanol was added to complete the volume of the pycnometer. The weight of this system was labeled as $W_2$. Finally, the sample saturated with the ethanol was taken out of the pycnometer and the residual weight of the ethanol and the pycnometer was labeled as $W_3$. The porosity of each membrane was calculated according to equation 1. The porosity of each membrane was obtained as the mean value of the porosity determined in three samples.

$$
\varepsilon = \frac{W_2 - W_3 - W_s}{W_1 - W_3}
$$

(1)

**Biomimetic mineralization of fiber samples:** The simulated body fluid (1.5xSBF) was prepared according to the procedure proposed by Oyane et al. [27] in order to mimic ion blood plasma concentrations. Briefly, NaCl, NaHCO$_3$, Na$_2$CO$_3$, KCl, K$_2$HPO$_4$, MgCl$_2$.6H$_2$O, HEPES, CaCl$_2$.2H$_2$O and Na$_2$SO$_4$ were accurately weighted and dissolved in ultra-pure water (Milli-Q, Millipore) at 36.5 ± 0.5 °C, with the help of a magnetic bar. All chemicals were reagent grade from Sigma Aldrich. Final pH was adjusted to 7.40 at 36.5± 0.5 °C using a 1M NaOH solution. 1.5xSBF freshly prepared solutions were stored in a refrigerator (5-10 °C) in closed plastic containers and used within 30 days after preparation. Solution stability was evaluated by periodic control (7/7 days) of salt precipitation and deposition after 48h at 36 ± 0.5 °C, according to the protocol explained elsewhere [28]. Electrospun PLLA and microcomposite gHA - PLLA membranes were cut into 10 mm diameter specimens and placed into conical plastic containers. The volume of SBF was calculated according to Kokubo and Takadama method [28] and using the following equation (2):
where $V_s$ is the volume of SBF (mL) and $S_a$ is the apparent surface area of specimen (mm$^2$). Immersed samples were kept at 36 ± 0.5 °C, and for each time point, samples were collected (in triplicates), gently washed with ultra-pure water and dried in a desiccator until SEM and XRD analysis. The SBF solution was renewed every 48 h to ensure sufficient ion concentrations.

**Cytocompatibility assays**

**Cell culture**: MG 63 osteoblastic-like cells (ATCC number CRL-1427TM, passage 25) were cultured in MEM-α supplemented with fetal bovine serum (FBS, 10% v/v), 50 μg mL$^{-1}$ ascorbic acid, 100 IU mL$^{-1}$ penicillin, 2.5 μg mL$^{-1}$ streptomycin and 2.5 μg mL$^{-1}$ amphotericin B, and maintained at 37 °C in a 5 % CO$_2$ humidified atmosphere. Cells were seeded in 25 cm$^2$ T-flasks until 70 – 80% confluence is reached. Detachment of adherent cells was achieved by a 5 min incubation in 0.05% trypsin – 0.25 % EDTA solution, at 37 ºC. Cell counting was performed using a 1:1 proportion of Trypan Blue and the colorless (viable) cells were counted in a Neubauer haemocytometer chamber. Electrospun PLLA and microcomposite gHA - PLLA membranes were cut out with punch (13-mm in diameter), sterilized under ultraviolet (UV) light for 30 min each side, and pre-incubated with the α-MEM for 2 h at 37 °C in a humidified atmosphere of 95 % air and 5 % CO$_2$. Specimens were then placed onto a 24-well culture plate, being held at the bottom by individual 7.8 mm diameter Teflon inserts. Cells were seeded over the fiber mats and on tissue culture polystyrene wells (TCPS controls) at a density of $1.5 \times 10^4$ cells/well. Cultures were incubated for periods up to 7 days, and characterized for cell viability, alkaline phosphatase (ALP) activity and observation by SEM and
confocal laser scanning microscopy (CLSM) for cell adhesion, morphology and pattern of cell growth.

**Cell viability/proliferation:** Cell viability was estimated by the resazurin assay, a nontoxic alamar blue dye (resazurin) that is reduced by intracellular enzyme activity to resofin, a fluorescent form of alamar blue [29, 30]. After 1, 3 and 7 days in culture, the medium was carefully removed from the wells and fresh media with 50 µL of 10% (v/v) of resazurin solution was added to each well. Samples were then incubated for 3.5 h at 37 ºC in humidified atmosphere (95 % air / 5 % CO₂). The fluorescence intensity was measured in a Power Wave XS2 spectrophotometer (Biotek) at 530 and 590 nm for excitation and emission wavelength, respectively.

**ALP Activity and Total Protein Content:** ALP synthesis by MG 63 osteoblastic-like cells cultured on the fiber membranes and TCPS controls was determined after 3 and 7 days, using p-nitrophenyl phosphate (pNPP) solution as the reaction substrate. For this purpose, cultured membranes were gently washed with PBS and with a lysis solution (0.1% Triton X-100) and then samples were stored at -20 ºC until ALP and total protein measurements. Thawed samples were then incubated with pNPP in alkaline buffer for 1 h at 37 ºC. The reaction was stopped with NaOH (5 M) and the absorbance of the hydrolysis product (p-nitrophenol) was immediately measured at 400 nm using a plate reader (Power Wave XS2 spectrophotometer, Biotek). ALP activity results were normalized by total protein content, quantified by Lowry’s method.
SEM and CLSM observation: For SEM analysis, samples were fixed in 2.5% glutaraldehyde for 1 h at room temperature, rinsed with distilled water and dehydrated through immersion for 30 minutes in a series of successive ethanol:water solutions (55, 70, 80, 90, 95 and 100 % v/v of ethanol). Samples were then dried at room temperature and coated with gold prior to SEM analysis. The average cell area was measured using the Image Pro Plus 4.01 analysis software (Media Cybernetics), on five representative fields of each sample.

For CLSM assessment, samples were fixed in 3.7% paraformaldehyde. Cell cytoskeleton filamentous actin (F-actin) was visualized by treating cells with Alexa Fluor 488-conjugated phalloidin (1:20 dilution; 1 h) and counterstaining with propidium iodide (1 µg/mL; 10 min), for cell nuclei labelling. Stained samples were mounted in Vectashield® and examined in a Leica SP2 AOBS microscope.

Statistics and Data Analysis: All quantitative data is presented as mean± standard deviation. Significant differences in statistical analysis were determined using one-way ANOVA test with Tukey’s HSD post hoc analysis. Statistical differences are found when p<0.05.

Results and Discussion
It was reported that the cells can detect and respond to the mechanical properties of the extracellular environment, by altering cytoskeletal organization, cell-substrate adhesion and other important processes for regulating cell behavior [31]. Figure 1 illustrates the fabrication process of the microcomposite gHA - PLLA electrospun membrane. gHA granules were dispersed in a polymer solution and placed in a glass syringe and electrospun according to the previously described in the experimental section. The
amount of ceramic present in the polymer solution was chosen by trial and error, and
the maximum content of filler possible was added to the electrospun solution. Further,
the purpose of using of large particles was to immobilize them on the outside of the
polymer fibers. This was done to increasing the contact area between the ceramic filler
and the cells, and the overall surface overall surface roughness and the local stiffness of
the membrane, due to the presence of a soft polymeric matrix and the hard-ceramic
filler, which could resemble the native ECM features.

**Microcomposite sample characterization**

Figure 2 shows the morphology of the synthetized ceramic filler, pristine polymer fibers
and microcomposite electrospun fiber membranes. The ceramic filler particles presented
irregular shape and size (≤150 µm) due to the milling procedure performed during the
material synthesis (figure 2a). It was observed that the PLLA electrospun fibers showed
a porous and interconnected network of smooth and defect free fibers (figure 2b).
Furthermore, when the ceramic filler is added to the polymeric solution and the mixture
electrospun, a microcomposite with morphologic features like the PLLA membranes is
obtained, containing micro sized gHA granules placed in the free spaces between the
polymeric cylindrical fibers, producing a configuration like “islands in the sea” (figure
2c and d). The purpose of this special configuration is to create a membrane surface
with homogeneously distributed ceramic filler, generating a topography with heights
and valleys, increasing both surface area and roughness. This mechanically challenging
substrate may indeed potentiate cell behavior, since it had been related to the increasing
of focal adhesions and cytoskeleton organization, migration and differentiation [31-34].
Additionally, the presence of bioactive gHA granules exposed on the outside of the fiber
further increases their exposure to cells during culture, as opposed to when they are
completely wrapped by the polymer or placed in the core of the fiber, which is often observed when nanoparticles are used as fillers [22, 35]. Interestingly, despite the broad size of gHA particles used in this work, small ceramic particles (less than \( \leq 100 \) nm) were also immobilized in the surface of the individual fibers (inset of figure 2d). In addition to increasing the roughness of the fibers compared to the neat PLLA fibers, this feature also provides nanotopographic decoration of individual PLLA fibers (hydrophobic) with hydrophilic ceramic nanoparticles. Both micro- and nano patterning associated with stiffness change are typical features of mechanical environment of native ECM, which ultimately modulates cellular responses [31, 36].

Average fiber diameter was calculated over 50 individual measurements with the help of the image J software. It was observed that PLLA pristine nanofibers have a mean diameter of 510 ± 150 nm, while the polymer fibers in the microcomposite membranes have an average diameter of 440 ± 170 nm (table 1). This decrease in the average fiber diameter is probably due to the charge distribution on the polymer droplets induced by ceramic particles. gHA can be ionized and carry more electrical charges, which will stretch the droplet even further, resulting in a decrease on the final average fiber diameter.

Membrane wettability is an efficient and simple method to evaluate surface properties on polymer fiber surface. Pristine PLLA and microcomposite electrospun membranes present a water contact angle (WCA) around 132 ± 3º, which suggests a strong hydrophobic behavior for all prepared samples, and the incorporation of the ceramic filler does not change the membrane surface properties (Table 1). The area of the water droplet in contact with the membrane is bigger than the individual gHA microparticles and the WCA is a result of the wettability of the polymer and ceramic all together, and
due to the high surface area of the polymer fibers, probably their contribution dominates the WCA for the microcomposite samples.

Membranes average porosity was assessed by gravimetric method and applying equation 1. The average porosity calculated for the pristine PLLA membranes was around 79 ± 3 %, while an increase of the overall porosity up to 88 ± 5% was observed for the microcomposite samples (Table 1). This is probably related to the incorporation of micro sized ceramic fillers that disturb packing of the polymer fibers, creating a less dense fiber membrane when compared with pristine PLLA one. In figure 2d, one can observe that around the gHA microparticles the pore size is bigger between fibers, when compared to the pristine PLLA membrane.

Figure 3 shows the XRD patterns of the synthetized gHA microparticles and the microcomposite polymer matrix membranes. The ceramic filler was composed by hydroxyapatite, and β and α-tricalcium phosphate (TCP) secondary phases (figure 3). XRD pattern of the microcomposite membranes presents only the main diffraction peaks characteristic of the ceramic filler, without any evidence of the crystalline diffraction peaks of the polymeric matrix. The XRD results suggest that when the solvent evaporates during the path between the needle tip and the ground collector, PLLA chain reorganization is fast and the polymer chains do not suffer any rearrangements towards an ordered fashion, thus remaining amorphous. Previous works reported that the electrospinning process leads to the amorphous PLLA chains [35, 37], which is in accordance with our XRD results.

Thermal gravimetric analysis allowed the study of the thermal stability of the processed membranes and to quantify the amount of the gHA particles that were effectively added to the microcomposite fiber membrane. It was observed that the PLLA polymer does not have adsorbed water, which corroborates the strong hydrophobic behavior obtained
from the water contact angle (figure 2). Further, the degradation of the polymer matrix occurs in a single step, with the major weight loss process observed in the range of 300 – 400 °C (figure 4). The comparison between the residual weight of the neat PLLA sample and microcomposite one showed that the amount of ceramic present in the fibrous membrane was around 60 % (figure 4).

The representative quasi-static behavior recorded in tensile mode, for the PLLA and gHA - PLLA samples is presented in figure 5. PLLA electrospun samples presented a maximum stress around 2.8 ± 1.2 MPa and a strain at break of 38 ± 7 %. However, the microcomposite samples, showed a 5-fold decrease of the maximum stress down to 0.57 ± 0.2 MPa and an increase of the strain at break up to 41 ± 8 %. Moreover, the samples are organized in a non-woven fashion, which suggests that only a small amount of fibers can resist to the applied mechanical load, especially for the microcomposite samples, where the increase of the overall porosity decreases the chain entanglement per unit of mass, lead to a decrease of the maximum stress, but without compromising the material stretchability.

_Biomimetic mineralization studies_

Biomineralization with simulated body fluids provides a useful _in vitro_ tool when manufacturing biomimetic materials for bone injury repair, as it can provide evidence on material’s ability to mimic native topography and chemical composition of bone extracellular matrix, eventually improving scaffold osteointegration [38]. SEM images of mineralized electrospun PLLA and gHA - PLLA microcomposite fibers are depicted in figure 6.
After being soaked in 1.5xSBF for 1 week, pristine PLLA membranes exhibited a flat sheet of mineral apatite throughout fibers surface. gHA-PLLA membranes presented more mineralization for the same time point, homogenously dispersed throughout individual fibers and seemed to enwrapped each fiber (insets), even those within the mat. In a time-dependent fashion, mineral deposition increased after 4 weeks for both membranes, presenting however, distinct topography. For pristine PLLA membranes, the development of large spherical minerals covering all the surface was observed, contrasting with the sheet-like morphology observed at week 1. By opposition gHA-PLLA microcomposites were represented by smaller and thicker spherulites wrapping the individual fibers (figure 6c and 6d). In the composite membranes, at the breaking points of the individual fibers, it was possible to observe the single polymeric fibers filling the core of a mineral cylinder and the overall thickness could reach up to 20 microns in diameter. It was confirmed that the biomineralization process in the microcomposite occurs not only on the surface, as it was observed for the pristine PLLA membranes, but also throughout the bulk of the membranes; this effect could be explained by the cumulative effect of an increase in porosity of the microcomposite, which allows a highly efficient ion transport into the inner pores [39] and the transiently supersaturation of $\text{Ca}^{2+}$ and $\text{PO}_4^{3-}$, due to gHA dissolution and that may provide nucleation points for HA growth [40].

The nature of the mineral phase was studied by energy dispersive spectroscopy (EDS), performed on the individual fibers, where the spectra showed the presence of Ca (calcium), P (phosphorus), and O (oxygen) as major elements (figure 7a). By using the intensity of the carbon element as a reference, the intensities from phosphorous and calcium seemed to increase with the increasing of the incubation time in the SBF solution. The Ca:P ratio (table 2) obtained for the samples was smaller when compared
to the stoichiometric HA (Ca:P = 1.67), which suggests the formation of Ca-deficient hydroxyapatite [41]. These results indicate that both PLLA and gHA-PLLA microcomposite membranes promote apatite formation on their surfaces, the main feature to improve osteoinductive properties and osteointegration [28, 42].

The XRD spectra obtained for the pristine and microcomposite samples incubated during different times showed the presence of HA with a broad peak at $2\theta = 32^\circ$ and small ones at $2\theta = 26^\circ, 28^\circ$ and $47^\circ$ (figure 7b).

**Cytocompatibility studies**

Cell viability/proliferation was estimated by the resazurin assay for up to 7 days exposure using an osteoblastic cell line MG-63 and it is reported in figure 8a. Cell density increased gradually with time in culture, as seen by the results observed at days 1, 3 and 7. Values were similar in the three surfaces, i.e. no significant differences were observed between membranes or with controls ($p>0.05$).

The potential of the developed microcomposite membranes for orthopedic tissue engineering was performed by monitoring the ALP activity, which is an important molecule in hard tissue formation and a hallmark of osteogenesis [43, 44]. ALP activity of MG-63 cells cultured on PLLA and gHA-PLLA fiber membranes, after 3 and 7 days, is shown in figure 8b. A statistically significant increase ($p<0.05$) was found after 7 days for gHA-PLLA membranes when compared to PLLA ones. ALP, a membrane bound enzyme, is synthesized during the early matrix formation and maturation period, and is essential in the onset of mineral deposition. Mineralization initiates in small vesicles that are formed from the cytoplasmic membrane of the osteoblast by a budding process. The membrane vesicles have a high activity of ALP that hydrolyses organic phosphate compounds, providing appropriate concentrations of phosphate ions that concentrate near the membrane. The vesicles also contain proteins and phospholipids.
with calcium-binding ability, allowing the localized accumulation of calcium and phosphate ions. Once the levels of these ions overcome the point of solubility, deposition of calcium phosphate occurs within the vesicles. Upon filling the vesicles, inorganic crystals contact with the extra-vesicular space which, in normal conditions, is rich in calcium and phosphate ions [45-47]. During the extravesicular mineralization, crystals of hydroxyapatite fill the inter-collagen fibrils spaces of the osteoid and, upon the formation of stable mineral “critical nucleus”, crystal growth becomes a faster physic-chemical driven process [43, 45, 48]. Due to its role in the osteoblast-mediated mineral deposition, ALP is a widely used biochemical marker of osteoblast differentiation. Back to the present work, the higher ALP activity observed in the gHA-PLLA fiber membrane and the expected increased availability of calcium and phosphate ions in the fibrillar matrix due to the presence of the ceramic filler would suggest the possibility of a faster and more efficient matrix mineralization in the microcomposite membrane compared to the PLLA membrane.

Membranes seeded with osteoblastic cells were stained for the F-actin cytoskeleton and nucleus and observed after 24 h by CLSM (figure 9). The F-actin cytoskeleton is composed of linear actin polymer microfilaments and is an important structure for a wide range of cellular functions. It provides structural stability having an essential role in the establishment of the cell shape and intercellular junctions. F-actin cytoskeleton is highly dynamic to cope with cellular internal needs and response to the surrounding environment, thus being essential in functions such as cell division, intracellular transport, signaling pathways and gene expression, as well as in cell motility and mechano-transduction mechanisms [49-51]. As such, assessment of the re-organization of the F-actin cytoskeleton during the cell adhesion and spreading to new substrates is an early indicator of the subsequent cell behavior. Images on figure 9 (a, c) show that
cells were randomly distributed on both membranes and presented a round-elongated 
morphology. On the PLLA membrane, cells showed a poor defined F-actin 
cytoskeleton, with a diffuse globular appearance and little evidence of a filamentous 
organization on high magnification images (figure 9b); also, some cells presented a 
nucleus with compacted chromatin and cytoplasm with dense F-actin (figure 9b, 
bottom). Figure 9c shows that on the gHA-PLLA membranes, cells exhibited better-
declared shape and cytoskeleton organization and well-evident nucleus; additionally, F-
actin staining was more intense in the cell limits, which is expected as this structure is 
highly concentrated beneath the plasma membrane. The high magnification images 
(Figure 9d) show evidence of filamentous actin organization with the actin stress fibers 
stretching towards the cell ends. These observations suggest a better substrate 
adaptation on the gHA - PLLA membranes compared to the PLLA membranes, which 
 might be predictive of an improved cell response.

SEM observation also shows that both membranes enabled cellular adhesion and 
spreading on their surfaces (figure 10) displaying a typical flat phenotype [29] with 
extended cytoplasm, appearing to interact and associate with surrounding fibers. 
Comparatively, at day 1, an increased cell spreading was verified on gHA-PLLA 
membranes with cells adopting a more polygonal morphology and a significant higher 
cell surface area, whether those grown on PLLA membranes presented a more fusiform 
structure. In addition, cells with more lamellopodia connecting to the fiber structure of 
the membrane were clearly identified on gHA-PLLA (Figure 10, inset). Cells 
proliferated actively and, at day 7, the establishment of intercellular contacts were 
identified within both formulations, whether increased cell-to-cell contact was observed 
for gHA-PLLA, further supporting the relevant role of gHA on the adhesion, 
proliferation and establishment of intercellular contacts.
Conclusions

A novel processing method to incorporate gHA microparticles in fiber membranes was developed, which placed a higher number of particles in the porous between the PLLA fibers creating a structure like “islands in a sea”, thus increasing both surface area and the roughness. Biomineralization studies showed that gHA – PLLA composites also induced HA crystal nucleation and growth, but with a distinct pattern of mineral deposition. Biological evaluation results indicate that the presence of the gHA microparticles allowed a better F-actin cytoskeleton organization during the initial cell adhesion and spreading, enhanced the cell-fibers and cell-to-cell interactions favoring the formation of a continuous cell layer and, also, increased ALP activity, which make these materials potential candidates for bone healing and repair strategies. The simplicity of the developed processing method creates a unique combination of stiff microcomposite fiber membranes and makes them potentially suitable for biomedical applications such as bone repair.

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References


[42] Zhang C, Cao M, Lan J, Wei P, Cai Q, Yang X. Regulating proliferation and differentiation of osteoblasts on poly(l-lactide)/gelatin composite nanofibers via timed


Figures

Figure 1. Illustration of the fabrication electrospinning setup used to obtain the gHA - PLLA membranes.
Figure 2. Representative SEM images: a) gHA granules, b) As-spun neat PLLA fibers, c) As-spun gHA - PLLA fibers and d) Detail of a gHA microparticle entrapped in the free space between polymer fibers. Inset: gHA nanoparticles immobilized on the surface of PLLA electrospun fibers.
Figure 3. XRD pattern of gHA and PLLA-gHA microcomposite electrospun membranes.

Figure 4. Thermal gravimetric analysis of gHA and PLLA-gHA microcomposite fiber membranes.
Figure 5. Representative stress-strain data obtained for the neat PLLA and gHA - PLLA microcomposite samples.
Figure 6. SEM images of electrospun samples submitted to in vitro biomineralization studies: a) PLLA sample after 1 week, b) PLLA sample after 4 weeks, c) gHA-PLLA sample after 1 week and d) gHA-PLLA sample after 4 weeks. Biomineralization studies were performed at pH = 7.4 and at 36.5 ± 0.5 ºC in 1.5xSBF.
Figure 7. EDS analysis of depositions formed on electrospun fibers after mineralization studies for 1 and 4 weeks in pristine PLLA and gHA - PLLA microcomposite.
Figure 8. (a) Cell proliferation estimated by resazurin assay and (b) ALP activity of MG-63 cells cultured on PLLA and gHA-PLL A membranes for 1, 3 and 7 days. Controls were performed on TCPS. Data plotted as mean ± standard. Statistical analysis: *p<0.05.

Figure 9. CLSM representative images of MG-63 osteoblastic cells cultured on PLLA (a, b) and gHA-PLL A membranes (c, d) for 24 h. Scale bar: 40 µm (a, c) and 20 µm (b, d).
Figure 10. Morphology of MG-63 cells cultured on PLLA and gHA – PLLA, for 7 days. Graph shows the cell surface area determined at 1 day of culture.
Tables

**Table 1.** Physical properties (fiber diameter, porosity and water contact angle) of PLLA and gHA-PLLA fiber membranes.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Fiber average diameter (nm)</th>
<th>Porosity (%)</th>
<th>Water contact angle (º)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLLA</td>
<td>510 ± 150</td>
<td>79 ± 3</td>
<td>131 ± 2</td>
</tr>
<tr>
<td>gHA-PLLA</td>
<td>440 ± 170</td>
<td>88 ± 5</td>
<td>132 ± 3</td>
</tr>
</tbody>
</table>

**Table 2.** The Ca:P ratio for pristine PLLA and gHA-PLLA immersed during 4 weeks in 1.5xSBF.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Time (weeks)</th>
<th>Ca/P</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLLA</td>
<td>1</td>
<td>1.40±0.98</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.35±0.59</td>
</tr>
<tr>
<td>gHA-PLLA</td>
<td>1</td>
<td>0.95±0.06</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.20±0.22</td>
</tr>
</tbody>
</table>