MICROSTRUCTURE AND PROPERTIES OF NANO-FIBROUS PCL-\textit{b}-PLLA SCAFFOLDS FOR CARTILAGE TISSUE ENGINEERING

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Abstract

Nano-fibrous scaffolds which could potentially mimic the architecture of extracellular matrix (ECM) have been considered a good candidate matrix for cell delivery in tissue engineering applications. In the present study, a semicrystalline diblock copolymer, poly{(\varepsilon-caprolactone)-block-poly(L-lactide)} (PCL-\textit{b}-PLLA), was synthesized and utilized to fabricate nano-fibrous scaffolds via a thermally induced phase separation process. Uniform nano-fibrous networks were created by quenching a PCL-\textit{b}-PLLA/THF homogenous solution to -20°C or below, followed by further gelation for 2 hours due to the presence of PCL and PLLA microcrystals. However, knot-like structures as well as continuously smooth pellicles appeared among the nanofibrous network with increasing gelation temperature. DSC analysis indicated that the crystallization of PCL segments was interrupted by rigid PLLA segments, resulting in an amorphous phase at high gelation temperatures. Combining TIPS (thermally induced phase separation) with salt-leaching methods, nano-fibrous architecture and interconnected pore structures (144±36 μm in diameter) with a high porosity were created for in vitro culture of chondrocytes. Specific surface area and protein adsorption on the surface of the nano-fibrous scaffold were three times higher than on the surface of the solid-walled scaffold. Chondrocytes cultured on the nano-fibrous scaffold exhibited a spherical condrocyte-like phenotype and secreted more cartilage-like extracellular matrix (ECM) than those cultured on the solid-walled scaffold. Moreover, the protein and DNA contents of cells cultured on the nanofibrous scaffold were 1.2-1.4 times higher than those on the solid-walled scaffold. Higher expression levels of collagen II and aggrecan mRNA were induced on the nanofibrous scaffold compared to those on the solid-walled scaffold. These findings demonstrated that scaffolds with a nanofibrous architecture could serve as superior scaffolds for cartilage tissue engineering.

Keywords: Cartilage, tissue engineering, PCL-\textit{b}-PLLA, thermally induced phase separation, nanofibrous scaffold.

Introduction

Cartilage defects resulting from aging and joint injury usually cause joint pain and loss of mobility. However, cartilage repair and regeneration are major challenges despite recent progress in orthopedic surgery, because the lesions do not spontaneously heal and their consequences have a significant social and economic impact (Brooks et al., 2002; Sellards et al., 2002). The tissue engineering strategy provides a promising therapy to repair cartilage defects and restore joint functions (Stoddart et al., 1999; Ochi et al., 2001; Cancedda et al., 2003; Tuli et al., 2003). A three-dimensional (3D) scaffold is employed for delivery or recruitment of reparative cells in an organized manner to bridge voids within cartilage defects, offering considerable promise as repair strategies (Solchaga et al., 2001; Grande et al., 1999). Thus, 3D biodegradable synthetic polymeric systems play a critical role in tissue engineering. The nature of scaffold and the physicochemical properties of the material are critical parameters for tissue formation (Chen et al., 2002).

Polyhydroxyacids, such as polylactide (PLA), polyglycolide (PGA), poly{(\varepsilon-caprolactone)} (PCL) and their copolymers poly(lactide-co-glycolide) (PLGA), poly(lactide-co-caprolactone) (PLC), poly(glycolide-co-caprolactone) (PGC), have been extensively used in tissue engineering due to their advantages of wide availability, ease of processing, adjustable degradation and mechanical properties, and low inflammatory response (Nair et al., 2007). These polymers were also applied in cartilage tissue engineering, and were documented to support the attachment and proliferation of primary animal chondrocytes (Chu et al., 1995; Gugala et al., 2000; Freed et al., 1994). However, PLA, PGA and PLGA suffer from the disadvantages of rigidity which limited their application as tissue-engineered cartilage (Grad et al., 2003). PCL is in a rubbery state at room temperature due to a low $T_g$ (glass transition temperature) and suffers a very slow degradation in vivo. Therefore, PLC or PGC might be promising materials for cartilage reconstruction because the incorporation of PCL could enhance the flexibility of the scaffold, and the degradation could be...
modulated by changing the composition of the copolymer (Jeong et al., 2004; Zhao et al., 2007; Kister et al., 2000; Li et al., 1996). It was also documented that biodegradable polyurethanes based on the PCL and PLLA segments were alternative candidates for cartilage tissue engineering, and that they were degraded to nontoxic by-products in vivo (Gorna et al., 2006).

Recently, intensive studies showed that cells recognized nanometric topologies of fibrous or microporous structure (Laurencin et al., 1999). Such a nanoscale structure, which geometrically mimicked the native state of the extracellular matrix (ECM), could selectively enhance protein adsorption and consequentially enhance cell attachment (Woo et al., 2003), and it therefore received much academic attention in medical applications (Ma, 2008). Zhang and Ma pioneered the preparation of PLLA nano-fibrous scaffolds with a fiber diameter ranging from 50 to 500 nm through thermally induced liquid-liquid phase separation (L-L TIPS) (Ma and Zhang, 1999). In a study by Hu et al., MC-4 cells acquired a more rounded shape on a PLLA nano-fibrous matrix with abundant interactions with nanofibers. No long dense stress fibers or typical focal adhesion structures were formed on the nano-fibrous matrix (Hu et al., 2008). Moreover, the nano-fibrous architecture of PLLA enhanced the osteogenic differentiation and mineralization of embryonic stem cells compared to the solid-walled architecture in both two and three dimensional cultures (Smith et al., 2009). In this regard, the nano-fibrous architecture may provide an environment closer to the ECM than the solid-walled architecture. Due to the mechanism of nanofiber fabrication by TIPS, semicrystalline homopolymers or copolymers which would crystallize and form microcrystalline domains during the TIPS process, were indispensable. PLLA/PCL blends were also utilized to prepare 3D nano-fibrous scaffolds by L-L TIPS from a THF solution (Zhao et al., 2004). However, PLLA with a high molecular weight could usually not be mixed with PCL, and the adhesion between PCL and PLLA became weak, resulting in poor mechanical properties (Kim et al., 2001).

In view of the facts above, we designed and synthesized a semicrystalline diblock copolymer, poly(ε-caprolactone)-block-poly(L-lactide) (PCL-b-PLLA). Both the PCL and the PLLA segments have the capability of crystallization, thus the nano-fibrous structure could be probably prepared by TIPS. For an ideal scaffold for cartilage tissue engineering, an interconnected pore network is a prerequisite to load cells and facilitate tissue ingrowth (Buma et al., 2004). Klompmaker’s studies revealed that implants with macropore sizes of at least 100-300 μm allowed fibrocartilage formation in the implant (Klompmaker et al., 1992; Klompmaker et al., 1993). Consequently, combining TIPS with the salt-leaching method, a high porosity with an interconnected pore structure (144 ± 36μm) and nano-fibrous architectures were created from PCL-b-PLLA copolymer for in vitro culture of chondrocytes.

**Materials and Methods**

**Materials**

ε-Caprolactone (ε-CL, 99%, Aldrich, St. Louis, MO, USA) was distilled from CaH2 in vacuum under dry N2. L-Lactide (Purac Biochem, Gorinchem, the Netherlands) was purified by recrystallization from ethyl acetate twice, and dried in vacuum at room temperature. All reagents were used without further purification.

**Synthesis and characterization of polymers**

PCL-b-PLLA (W<sub>PCL</sub>/W<sub>PLLA</sub> = 50/50) block copolymer was synthesized according to Fig. 1. Firstly, PCL pre-polymer (PCL-OH) with a pre-designed molecular weight of 5.0×104 was polymerized in bulk with Sn(Oct)2 as catalyst and dodecanol as initiator at 130°C under vacuum for 30 h. The ring-opening polymerization of L-LA with the pre-designed molecular weight of 5.0×104 was then carried out at 130°C under vacuum for 30 h, using PCL-OH as macromolecular initiator and Sn(Oct)2 as catalyst. All crude products were purified by dissolution-precipitation in chloroform/alcohol. PLLA homopolymer with a molecular weight of 5.0×104 was obtained via polymerization of L-LA using Sn(Oct)2 as catalyst and dodecanol as initiator for 30h at 130°C under vacuum.
The structure of polymers was confirmed by 1H NMR and 13C NMR performing on a Mercury-Plus 300 spectrometer (Varian, Inc., Pala Alto, CA, USA), using CDCl3 as solvent. IR spectra were recorded on a Fourier transform infrared (FTIR) spectrometer (Nexus 670, Nicolet, GMI, Ramsey, MN, USA).

The molecular weight and its distribution were determined at 30°C by a gel permeation chromatography (GPC) system consisting of a Waters 1525 binary high pressure liquid chromatography (HPLC) pump, three ultrastyragel columns, and a Waters 2414 refractive index detector (Waters Alliance GPC2000, Waters Corporation, Milford, MA, USA).

**Fabrication of nano-fibrous scaffolds**

The nano-fibrous scaffold was fabricated by liquid-liquid phase separation from the solution of PCL-b-PLLA in tetrahydrofuran (THF). Briefly, a 5% (w/v) clear solution was obtained by dissolving a certain mount of PCL-b-PLLA into THF at 50°C. Then 3mL of polymer solution was quickly poured into a 25mL glass beaker followed by quenching to a chosen temperature (-40ºC, -20ºC, 4ºC, 8ºC, 12ºC, 16ºC) and maintained for 2h. Soon during this period, the solution became a gel. After that, the gel was immersed into 4ºC double-distilled water for 2 days to leach out the solvent by changing the water four times per day. The scaffold was obtained after lyophilization under 0.940mbar at 0°C for five days (ALPHA2-4 freeze dryer, Martin Christ Gefriertrocknungsanlagen GmbH, Osterode am Harz, Germany).

PLLA nano-fibrous scaffolds were also prepared by phase separation adopting the same steps above. The chosen Tgel was 4ºC, 8ºC, 12ºC and 16ºC.

To prepare a macroporous scaffold, NH4HCO3 particles (125-200μm) that were 5 times heavier than the copolymer were added to the 5% (w/v) polymer solution in THF, and followed by the same procedures as mentioned above. The scaffold was referred as nano-fibrous scaffold (NF scaffold). Solid-walled scaffold (SW scaffold) was fabricated as control by solid-liquid (S-L) phase separation from 5% (w/v) PCL-b-PLLA/dioxane solution. In this approach, the weight ratio of NH4HCO3 particles to polymer was still 5:1 (w/v), and the temperature of S-L phase separation was -40ºC. All samples were dried through lyophilization under 0.940mbar at 0°C for five days, and the NH4HCO3 particles were removed by leaching in double-distilled water. The scaffold was finally obtained by lyophilization.

**Characterization of PCL-b-PLLA scaffolds**

The morphology of scaffolds was observed by scanning electron microscopy (SEM) in a JSM-6380LA Analytical Scanning Electron Microscope (JEOL Ltd., Tokyo, Japan).

Thermal properties were analyzed with a Differential Scanning Calorimeter (MDSC2910: TA Instruments, Wilmington, DE, USA) in a temperature range from 20ºC to 220ºC with a heating rate of 10ºC/min. The first heating was performed for the analysis.

Porosity of PCL-b-PLLA macroporous scaffolds was measured by a gravimetric method. Dry samples were weighed, and their weight was denoted mdry. The samples were filled with ethanol under vacuum, and subsequently weighed, and their weight was denoted mwet. The volume of pores, Vpore, was calculated according to Eq. (1):

\[
V_{\text{pore}} = \frac{(m_{\text{wet}} - m_{\text{dry}})/\rho_{\text{ethanol}}}{V_{\text{scaffold}}} \quad (1)
\]

where \( \rho_{\text{ethanol}} \) was the density of ethanol which was 0.789g/cm³.

The volume of the whole scaffold was measured as the volume of double-distilled water (Vscaffold) by immersing the scaffold in water, assuming that the amount of water absorbed in the scaffold was negligible during the short time of the experiment. Therefore the porosity of scaffold was calculated as the quotient of the volume of pores and the total volume of the scaffold according to Eq. (2). Five measurements were carried out for each sample and the results were expressed as mean ± standard deviation (SD).

\[
\text{Porosity} = \frac{V_{\text{pore}}}{V_{\text{scaffold}}} \quad (2)
\]

The nitrogen (N2, 99.99%) adsorption measurement at 77K was conducted using a porosimetry analyzer (ASAP 2020, Micromeritics, Norcross, GA, USA). The specific surface area was calculated from the Brunauer-Emmett-Teller (BET) plot of the N2 adsorption isotherm.

The mechanical properties of the macroporous PCL-b-PLLA (50/50) scaffold were tested using a universal mechanical properties tester (Instron 3345, Norwood, MA, USA) with a 100N load-cell at a compression speed of 1mm/min. A cylindrical specimen with a diameter of 10mm and a length of 5mm was used. Fully dried samples were compressed between parallel plates at 25°C and 70% humidity.

**Protein adsorption to scaffolds**

Nano-fibrous-walled and solid-walled PCL-b-PLLA (50/50) macroporous scaffolds of equal weight were first wetted by soaking in ethanol for an hour and subsequently in phosphate buffered saline (PBS) for 30 min thrice. The scaffolds were then incubated in PBS overnight at room temperature. The wetted scaffolds were incubated in 0.5% (w/v) bovine serum albumin (Shanghai Bo’ao Biological Technology Co., Ltd. Shanghai, China) solutions in PBS for 4h. After the incubation, the scaffolds were removed from the protein solution and then washed in PBS thrice (1×1h) to remove free and loosely adsorbed proteins. The total amount of protein adsorbed to the scaffolds was measured using the MicroBCA assay (Pierce, Rockford, IL, USA) and was calculated from the standard curve of BSA.

**Chondrocyte isolation and culture**

Fresh chondrocytes were isolated from 3 separate articular cartilages harvested from the knee joint of a 1-year-old dog. Circa 1mm³ pieces of fresh canine articular cartilage were digested in 0.25% trypsin (2mL/g tissue) for 1h at 37°C, followed by 0.2% type II collagenase for 16h at 37°C. Digested suspension was filtered through a 100-mm mesh nylon filter and washed 2 times in PBS supplemented
with penicillin (100U/mL) and streptomycin (100mg/mL). Chondrocytes were seeded at an initial density of 3.5×10^4 cells/cm² at each passage in Dulbecco’s modified Eagle’s Medium (DMEM) containing 4.5mg/mL D-glucose, 0.1mM nonessential amino acids, 1mM sodium pyruvate, 100mM HEPES buffer, 100U/mL penicillin, 100mg/ml streptomyein and 0.29mg/mL L-glutamate and supplemented with 10% of fetal bovine serum at 37°C with 5% CO₂ in T-25 flasks. Culture medium was changed every 3 days. Cells were passaged with trypsin/EDTA (0.25%/0.1mM).

**SEM observation of cell morphology in scaffolds**

The NF and SW scaffolds were sterilized with 75% alcohol for 24h and then washed in PBS for three times. Chondrocytes of passage 3 were seeded on the scaffolds of 6mm in diameter and 2mm in height (SW scaffold and NF scaffold) at a density of 1×10⁶ cells/scaffold in a 48-well plate. At culture time points of day 3, day 6 and day 12, the scaffolds were washed once with PBS, and then fixed with 2.5% glutaraldehyde fixative and osmium tetroxide at 4°C for 2h each. After dehydration with a series of graded alcohols, the cells kept in a fume hood for air-drying. The scaffold was then gold-sputtered in vacuum and examined by SEM.

**Protein content measurements**

Chondrocytes were seeded at a density of 2×10⁶ cells/scaffold in a 48-well plate on the PCL-β-PLLA macroporous scaffolds (SW scaffold and NF scaffold). At culture time points of day 3, day 6 and day 12, the composites were lysed by RIPA (Radio Immunoprecipitation Assay) Lysis Buffer (150mM NaCl, 0.1% SDS, 1% sodium deoxycholate, 1% triton X-100, 50mM Tris-HCl (pH 7.5), 2mM EDTA) (Biocolor BioScience&Technology Co., Shanghai, China) on ice for 30 minutes. The suspension was centrifuged in the refrigerated microcentrifuge at maximum speed for 10 min and the supernatants were harvested. The content of protein was determined by BCA™ protein assay kit (Pierce) 0.1mL sample was mixed with 2.0mL work reagent and incubated at 37°C for 30 minutes. Then absorbance measurements were taken at 562nm after the samples were cooled to room temperature (FLUO star OPTIMA; BMG Labtech, Offenburg, Germany). The protein content was calculated from the standard curve of BSA.

**DNA content measurements**

Chondrocytes were seeded at a density of 2×10⁶ cells/scaffold in a 48-well plate on the PCL-β-PLLA macroporous scaffolds (SW scaffold and NF scaffold). At culture time points of day 3, day 6 and day 12, the composites were digested overnight at 60°C in a solution of protease K (0.5mg/mL). The 100μL of lysate was mixed with 1ml of Hoechst 33258 (Sigma, St Louis, MO, USA) working solution (0.1μg/ml) in TEN (10mM Tris, 1mM Na₂EDTA, 0.1M NaCl, pH 7.4). Fluorescence was measured at excitation (EX) 365 and emission (EM) 458nm in the fluorometer. The DNA content was calculated from the standard curve of DNA from calf thymus (Sigma).

**Gene expression analysis for Collagen type II and aggrecan**

Chondrocytes were seeded at a density of 2×10⁶ cells/scaffold in a 48-well plate on the PCL-β-PLLA macroporous scaffolds (SW scaffold and NF scaffold). At day 3, day 6 and day 12, the collagen type II and aggrecan gene was assessed by RT-PCR (Reverse transcription polymerase chain reaction). Briefly, total RNA was extracted from cells using the RNeasy Mini RNA isolation kit (Qiagen, Crawley, UK). The cDNA was reverse transcribed from 1μg of total cellular RNA using ThermoScript RT-PCR system (Invitrogen, Carlsbad, CA, USA). The amplification primer pair for collagen type II was selected as 52 - CGA GTG GAC AAA GGA GAA GC -32†and 52 - GTC TCC GCT CTT CCA TTC AG -32, aggrecan was 52- TGG AGC AAC TGA CTC CAG TG -32†and 52- GGA CGC TAG TGC GTA ATGA CA -32. PCR reactions were performed using aliquots of the cDNA template with Platinum PCR Supermix kit (Invitrogen).

Pre-denaturation was 94°C for 5min; The cDNA was amplified by 30 cycles of denaturation (40s at 94°C), annealing (collagen II 40s at 60°C, aggrecan 40s at 60°C), and elongation (60s at 72°C), followed by a final step at 72°C for 10 min. PCR products were separated electrophoretically on a 2.0% agarose gel (Sigma) and the products were visualized by staining in ethidium bromide (Invitrogen).

**Results**

**Synthesis of PCL-β-PLLA copolymer**

PCL-β-PLLA block copolymer was synthesized in two steps. Firstly, hydroxyl-terminated PCL (PCL-OH) pre-polymer was synthesized as shown in Fig. 1, and its structure was confirmed by 1H NMR in Fig. 2(A). The molecular weight of PCL-OH was calculated from the integrals of the characteristic peaks at 2.3ppm and 0.87ppm (Guan et al., 2007). When ring-opening polymerization of L-LA was carried out using the macromolecular monomer PCL-OH as initiator, new signals attributed to the PLLA segment at 5.16ppm and 1.60ppm were detected in Fig. 2B. The resulting molar ratio and the molecular weight of block copolymer determined by 1H NMR were listed in Table 1. The resulting molar ratio of PCL segment to PLLA segment in copolymer was close to the feed molar ratios. The copolymer had one glass transition temperature and two melting point attributed to PCL and PLLA segment respectively.

In order to prove that the synthetized copolymer was block copolymer rather than random copolymer, 13C NMR and FTIR were used to confirm its structure. As shown in...
Table 1. Synthesis results for PCL-\(b\)-PLLA (50/50)

<table>
<thead>
<tr>
<th>Polymer</th>
<th>CL/L-LA Molar Feed Ratio</th>
<th>CL/L-LA Molar Resultant Ratio</th>
<th>(M_n,\text{NMR} \times 10^4)</th>
<th>(M_n,\text{GPC} \times 10^4)</th>
<th>(M_w/M_n,\text{GPC})</th>
<th>(T_g/\degree C)</th>
<th>(T_c/\degree C)</th>
<th>(T_m/\degree C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCL-OH</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>6.20(^a)</td>
<td>-</td>
<td>-</td>
<td>-65.5</td>
</tr>
<tr>
<td>PCL-(b)-PLLA</td>
<td>1.26</td>
<td>1.44</td>
<td>11.60(^b)</td>
<td>9.0</td>
<td>1.22</td>
<td>-59.7</td>
<td>29.3(^c)</td>
<td>57.1(^d)/177.6</td>
</tr>
<tr>
<td>PLLA</td>
<td>-</td>
<td>-</td>
<td>4.80</td>
<td>5.0</td>
<td>1.12</td>
<td>60.8</td>
<td>-</td>
<td>177.2</td>
</tr>
</tbody>
</table>

\(^a\) Determined by \(^1\)H NMR: \(M_n,\text{PCL-OH}=I_{\delta=2.3}/I_{\delta=0.87}\times114\times1.5+114\times185\)
\(^b\) Determined by \(^1\)H NMR: \(M_n,\text{PCL-}b\)-PLLA = \(M_n,\text{PCL-OH} + I_{\delta=5.1}/I_{\delta=4.05}\times2\times72\timesn_{\text{PCL}}, n_{\text{PCL}}\) was the number of repeating segments in PCL-OH
\(^c\) the crystallization temperature of PCL segment
\(^d\) the crystallization temperature of PLLA segment
\(^e\) the peak melting temperature of PCL segment
\(^f\) the peak melting temperature of PLLA segment

Table 2. Thermal properties of PCL-\(b\)-PLLA (50/50) scaffolds prepared at different \(T_g\)\(^s\).

<table>
<thead>
<tr>
<th>(T_g) (\degree C)</th>
<th>(T_m,\text{PCL}) (\degree C)</th>
<th>(T_p,\text{PCL}) (\degree C)</th>
<th>(\Delta H_m,\text{PCL}) (J/g)</th>
<th>(T_m,\text{PLLA}) (\degree C)</th>
<th>(T_p,\text{PLLA}) (\degree C)</th>
<th>(\Delta H_m,\text{PLLA}) (J/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>-40 °C</td>
<td>50.1</td>
<td>63.1</td>
<td>40.8</td>
<td>164.1</td>
<td>177.2</td>
<td>27.5</td>
</tr>
<tr>
<td>-20 °C</td>
<td>52.4</td>
<td>63.6</td>
<td>39.5</td>
<td>163.9</td>
<td>178.0</td>
<td>29.0</td>
</tr>
<tr>
<td>4 °C</td>
<td>51.9</td>
<td>63.5</td>
<td>26.7</td>
<td>163.0</td>
<td>176.2</td>
<td>26.3</td>
</tr>
<tr>
<td>8 °C</td>
<td>46.7</td>
<td>62.6</td>
<td>29.5</td>
<td>163.7</td>
<td>175.4</td>
<td>28.2</td>
</tr>
<tr>
<td>12 °C</td>
<td>46.0</td>
<td>61.3</td>
<td>24.5</td>
<td>165.0</td>
<td>176.3</td>
<td>28.1</td>
</tr>
<tr>
<td>16 °C</td>
<td>44.6</td>
<td>62.5</td>
<td>25.2</td>
<td>163.4</td>
<td>176.4</td>
<td>33.5</td>
</tr>
</tbody>
</table>

\(^a\) the initial melting temperature
\(^b\) the peak melting temperature

Table 3. Properties of macroporous PCL-\(b\)-PLLA (50/50) scaffolds fabricated with different wall structures. \(p<0.05\), \(n=3\).

<table>
<thead>
<tr>
<th>sample</th>
<th>porosity</th>
<th>pore size ((\mu)m)</th>
<th>Specific surface (m(^2)/g)</th>
<th>Amount of protein adsorbed ((\mu)g)</th>
<th>Compression modulus (kPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCL-(b)-PLLA SW scaffold</td>
<td>95.5% ± 1.5%</td>
<td>168±69(^a)/40±15</td>
<td>2.90±0.45</td>
<td>249±43</td>
<td>90±15</td>
</tr>
<tr>
<td>PCL-(b)-PLLA NF scaffold</td>
<td>93.0% ± 1.0%</td>
<td>144±36</td>
<td>8.09±0.23</td>
<td>808±67</td>
<td>108±20</td>
</tr>
</tbody>
</table>

\(^a\) The diameter of the macropores created by the porogen.
\(^b\) The diameter of micropores created by evaporation of solvent.
Figure 2. $^1$H NMR spectra of PCL-OH (A) and PCL-$b$-PLLA (B) at 298K in CDCl$_3$.

Figure 3. $^{13}$C NMR spectrum of PCL-$b$-PLLA diblock copolymer (50/50) at 298K in CDCl$_3$. The two single resonance peaks at $\delta=169.7$ ppm and $\delta=173.6$ ppm were assigned to C=O of PLLA and PCL segments in the PCL-$b$-PLLA diblock copolymer (50/50).

Figure 4. FTIR spectrum of PCL-$b$-PLLA diblock copolymer. The two absorptions at 1757 cm$^{-1}$ and 1724 cm$^{-1}$ corresponded to the stretching vibration of ester carbonyl groups for PCL and PLLA blocks.

Figure 5. SEM pictures of PCL-$b$-PLLA (50/50) nano-scaffolds fabricated from a solution of 5.0% (w/v) in THF at different $T_{gel}$s: (A) -40°C; (B) -20°C; (C) 4°C; (D) 8°C; (E) 12°C and (F) 16°C.
Figure 6. SEM pictures of PLLA nano-scaffolds fabricated from a solution of 5.0% (w/v) in THF at different $T_{gel}$s: (A) 4°C; (B) 8°C; (C) 12°C and (D) 16°C.

Figure 7. The fiber diameter of PCL-b-PLLA (50/50) and PLLA nano-fibrous scaffolds at different $T_{gel}$. $p$ <0.05, n=3.

Figure 8. DSC thermogram of scaffolds prepared from 5% (w/v) PCL-b-PLLA (50/50)/THF solution at different $T_{gel}$: a) -40°C; b) -20°C; c) 4°C; d) 8°C; e) 12°C; f) 16°C. Two melting peaks at around 63°C and 177°C corresponded to the melting point ($T_m$) of PCL segment and PLLA segment.
Figure 9. SEM pictures of PCL-b-PLLA (50/50) nano-fibrous scaffolds with macroporous structure fabricated from a solution of 5.0 % (w/v) in dioxane (A, B) and THF (C, D). Scale bar in A, C = 500μm; in B = 50μm and in D = 5μm.

Figure 10. SEM micrographs of cartilage cells on the PCL-b-PLLA (50/50) porous scaffolds with solid wall (A, C, E) and nano-fibrous wall (B, D, F). Cells were cultivated for: (A, B) 12 hours; (C, D) 1 day; (E, F) 6 day. Scale bars for A, B, C, D: 2μm, E: 50μm, F: 30μm.

Figure 11. The contents of protein (A) and DNA (B) of the cell cultured on different scaffolds. NF scaffold: nano-fibrous scaffold, SW scaffold: solid-walled scaffold. *p < 0.05, n=7.
especially the freeze-drying. Among these steps, the gelation process, dissolving, gelation, solvent exchanging, freezing and fibrous scaffold by L-L TIPS, including polymer five characteristic steps were involved in fabricating nano-scaffolds at different PCL-al.

Relative higher (Fig. 6D) was observed instead of smooth pellicle structure diameter changing little. However, platelet-like structure T 5F). These fibers were thicker than those formed at low pellicles among the nano-fibrous network at 16°C (Fig. 5D and 5E) and even more continuous smooth knot-like structures (Fig. 5C), coating-like wrapped fibers increasing nanofibers were 95±39nm and 98±45nm, respectively. and -20°C (Fig. 5A and 5B). The diameters of the nano-fibrous network was only obtained at network was observed in all matrixes, but the uniform gels (4~12°C, Fig. 6A-6C), with the fiber gelation temperature (Tgel) on the morphologies and crystallization of PCL-b-PLLA scaffolds

Five characteristic steps were involved in fabricating nano-fibrous scaffold by L-L TIPS, including polymer dissolving, gelation, solvent exchanging, freezing and freeze-drying. Among these steps, the gelation process, especially the Tgel was critical in influencing the architecture of the scaffold (Ma and Zhang, 1999; He et al., 2009). Fig. 5 displays the morphologic evolution of PCL-b-PLLA scaffolds as a function of Tgel. These scaffolds were fabricated from a 5% (w/v) polymer solution in THF at different Tgel-s (-40°C, -20°C, 4°C, 8°C, 12°C, 16°C) with aging for 2h in the gel status. The nano-fibrous network was observed in all matrixes, but the uniform nanofibrous network was only obtained at Tgel-s of -40°C and -20°C (Fig. 5A and 5B). The diameters of the nanofibers were 95±39nm and 98±45nm, respectively. Increasing Tgel concomitantly tended to generate many knot-like structures (Fig. 5C), coating-like wrapped fibers (Fig. 5D and 5E) and even more continuous smooth pellicles among the nano-fibrous network at 16°C (Fig. 5F). These fibers were thicker than those formed at low Tgel-s of -40°C and -20°C (Fig. 7). Compared with pure PLLA scaffolds, nano-fibrous structures were obtained at relative higher Tgel-s (4~12°C, Fig. 6A-6C), with the fiber diameter changing little. However, platelet-like structure (Fig. 6D) was observed instead of smooth pellicle structure as shown in Fig. 5F.

DSC analysis was performed to study the crystallization behaviors of PCL-b-PLLA in nano-fibrous scaffolds fabricated at different Tgel-s (Fig. 8). Two melting peaks were observed at around 63°C and 177°C for all samples, which corresponded to the melting points (Tm) of PCL segment and PLLA segment. A little decrease in both Tm (Tm,PCL, Tm,PLLA), compared to homopolymer of PCL and PLLA, meant that the crystallization of PCL and PLLA segments were limited by one another. For PLLA segment, the initial melting temperature (Tm,PLLA), Tm,PLLA, and the melting enthalpies (∆Hm,PLLA) changed little within a broad scale of Tgel (-40~12°C). Whereas, ∆Hm,PCL increased obviously at the Tgel of 16°C. In the case of PCL segment, in spite of no obvious decrease in Tm,PCL was observed as a function of Tgel, a decrease in Tm,PLLA, a broad melting range and a definitely decline of ∆Hm,PCL were detected.

Properties of porous PCL-b-PLLA scaffolds with different architecture

For culture of chondrocytes in vitro, the salt-leaching method was incorporated into the phase separation technique to fabricate 3D scaffolds from PCL-b-PLLA. As shown in Fig. 9, large pores in scaffolds were created by leaching out NH4HCO3 particles, small pores and the texture of the porous wall were determined by the phase separation process, i.e., S-L TIPS or L-L TIPS. In spite of the large pores in the resultant scaffold, the diameter of which was 168±69μm, a ladder-like structure with a solid wall via S-L TIPS from a PCL-b-PLLA/dioxane/NH4HCO3 system (Fig. 9B) was observed. The size of ladder-like pores was measured as 40±15μm. For simplicity, this scaffold was defined as the SW scaffold. Fig. 9D shows the nano-fibrous networks in the large pores wall of the scaffold prepared via L-L TIPS from PCL-b-PLLA/THF/ NH4HCO3 system (defined as NF scaffold), the addition of salt particles did not destroy the nano-fibrous network. However, the diameter of a fiber was about 151±56nm, larger than those formed without the incorporation of salt leaching technique (Fig. 5A and 5B).

Though there was no statistical difference between the porosities of these two kinds of scaffolds (Table 3), it was noteworthy that the specific surface and the amount of

Figure 12. Expression of cartilage-specific genes in NF-scaffold and SW-scaffold chondrocyte cultures for different time analyzed by RT-PCR. The mRNA was analyzed for collagen II and aggrecan which showed bands of 549bp and 1025bp, respectively.

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protein adsorbed in NF scaffold were 2.8 and 3.2 times higher than those of the SW scaffold, respectively.

PLLA porous nano-fibrous scaffold prepared by the same process had a higher compression modulus (218±34KPa) than PCL-b-PLLA SW and NF scaffolds. As PLLA had a similar $M_f$ to that of the PLLA segment in the copolymer, the rigidity of the PCL-b-PLLA NF scaffold was decreased compared with the PLLA NF scaffold. However, the compression modulus of PCL-b-PLLA NF scaffold was still higher than that of the SW scaffold.

**Influence of the architecture of PCL-b-PLLA scaffolds on the chondrocyte morphologies and proliferation in vitro**

The morphology of chondrocytes cultured on NF and SW PCL-b-PLLA scaffolds were observed by SEM as shown in Fig. 10. In the early period, after 12h and 24h of culture, chondrocytes cultured in SW scaffolds showed a spherical shape with smooth surface, and then elongated, with few long slender cell processes. In contrast, cells on NF scaffolds assumed a rounded morphology and displayed a rough surface with abundant long slender cell processes or filopodia extensions. Meanwhile, chondrocytes secreted some matrix in the NF scaffolds. After 6 days, chondrocytes flattened and secreted ECM in SW scaffolds. In comparison, cells still maintained a “spherical” condrocyte-like phenotype, and abundant chondral ECM was observed covering the entire surface of the NF scaffold, which was hardly distinguishable from nano-fibrous matrix.

In the studies of cell proliferation and biosynthetic activity, chondrocytes were seeded onto both types of scaffolds and cultured up to 12 days. At 3, 6, 12 day intervals, the total protein and DNA contents on the NF scaffolds were 1.3-1.4 times and 1.2-1.4 times higher than those on the SW scaffolds, indicating that the cells proliferated more rapidly on the NF scaffolds than on the SW scaffolds.

**Influence of the architecture of PCL-b-PLLA scaffolds on the expression of cartilage-specific ECM genes**

RT-PCR analysis was employed to analyze the mRNA expression of cartilage-specific genes from chondrocytes cultured on the NF scaffold and the SW scaffold up to 12 days (Fig. 12). The expression of collagen II and aggrecan, which were the mRNA expression of cartilage-specific ECM genes, were positively induced in both NF-scaffold culture and SW-scaffold culture, indicating that no dedifferentiation trend of chondrocytes occurred during the culture period. Notably, both collagen II and aggrecan were expressed at a higher level in NF-scaffold culture than in SW-scaffold culture. The differences became more significant at day 12. These results indicated that the NF scaffold could serve as an environment more favorable for maintaining chondrocyte phenotype than the SW scaffold.

Many studies have reported that the transmission of mechanical stimulation signals throughout the scaffolds could effectively promote chondrogenic differentiation and significantly enhance chondral ECM deposition. Biodegradable polymers such as poly(L-lactide-co-caprolactone) (PLCL, 5:5) (Jung et al., 2008), poly(3-hydroxy- butyrate-co-3-hydroxyhexanoate) (PHBHx) (Wang et al., 2008), and poly(1,8-octanediol citrate) (POC) (Kang et al., 2006) have been utilized as scaffold materials for cartilage tissue engineering. One point in common of these materials was their complete rubber-like elasticity. Therefore, the semicrystalline PCL-b-PLLA block copolymer was chosen to support chondrocyte growth in our study.

In addition, chondrocytes tend to dedifferentiate, exhibiting profound phenotypic changes while growing in vitro for long periods (Huch et al., 2002; Stokes et al., 2001). Therefore, maintaining a chondrocytic phenotype is indeed challenging. It has been reported that nanoporous or nano-fibrous polymer matrices played a critical role in providing support and anchorage for cells, while maintaining chondrocyte phenotype and enhancing chondrocyte growth and matrix synthesis (Li et al., 2003; Li et al., 2006). The fabrication technologies used for scaffold fabrication in cartilage tissue engineering included electrospinning (Wise et al., 2009), particulate leaching combined with chemical etching (Pattison et al., 2005; Park et al., 2005) and 3-D printing techniques (Yen et al., 2009). L-L TIPS is an easy and effective technique to fabricate 3D nano-fibrous scaffolds (Ma, 2008; Zhang and Ma, 1999). A homogeneous polymer-solvent system would spontaneously undergo a liquid-liquid phase separation when thermodynamically quenched into the unstable zone under the spinodal curve, resulting in a separated polymer-rich phase and a polymer-poor phase. Nano-fibrous scaffolds can be obtained when the gelation of the polymer solution system occurs due to the crystallization of polymer-rich phase and solvent removal. Ma et al. have done intensive studies on the fabrication of PLLA NF scaffolds by TIPS and their influence on cell morphology and function in bone tissue engineering (Woo et al., 2003; Hu et al., 2008; Smith et al., 2009; Wei and Ma, 2008). Even now, no one reported the application of PCL-b-PLLA scaffold with nano-fibrous structure fabricated by TIPS in cartilage tissue engineering.

Based on the above discussion, the key to fabricate PCL-b-PLLA nano-fibrous scaffolds was to choose proper solvents and optimize gelation parameters. On one hand, nano-fibrous structure could result from the crystallization of PCL and PLLA segments independently in the TIPS process at $T_m \leq 4^\circ C$ using THF as a solvent. On the other hand, the crystallization behavior of these two blocks is confined to each other, especially for the PCL segment at high $T_m (\geq 4^\circ C)$. Li et al. reported that the crystallization ability of the PCL segment in the PLLA-PCL-PLLA block copolymer reduced in the presence of PLLA block (Zhao et al., 2007). Combined with the thermal analyses results, it was clear that the knot-like structure and smooth pellicles
as shown in Fig. 5 indicated the presence of the PCL amorphous phase other than PLLA crystals with platelet-like structure.

We also investigated the L-L TIPS behavior of PCL homopolymer in THF solution. When $T_{gel} > 4^\circ C$, no gelation phenomena were observed in the PCL ($M_w$: 80k)/THF system (5%, w/v). Zhao et al. (Zhao et al., 2004) also documented that the gelation time of the PCL/THF system was 60min ($T_{gel}$: -18°C), longer than the gelation time of the PLLA/THF system at the same $T_{gel}$, which was 22min. In this study, we found that increasing $T_{gel}$ would accelerate molecular movement which would not favor the microphase separation of the PCL domain from the PLLA domain in the PCL-b-PLLA/THF system. Thus more PCL molecular chains would be entrapped in the PLLA crystalline regions, which resulted in the decrease in PCL crystallinity. Meanwhile, PLLA crystallization was also interrupted by the PCL domain to some extent. This may explain why there were nanofibers other than the platelet-like structures as in the PLLA/THF system, when the PCL-b-PLLA/THF solution gelled at 16°C.

Although a porous structure can be formed by the L-L TIPS technique, the pore size is quite small for chondrocyte ingrowth as it is mainly determined by the polymer concentration. Moreover, previous studies revealed that a pore size of 100-300μm is essential for chondrocytes to infiltrate (Klompmaker et al., 1992; Klompmaker et al., 1993). Hence, we combined the salt leaching technique with the TIPS method to incorporate macro pores into NF and SW scaffolds. The pore size and the interconnection between pores can easily be modulated by controlling the size and quantity of the porogen.

Chondrocyte culture in vitro showed that the cells could infiltrate in the porous scaffolds with both pore morphology types as shown by SEM, implying that the pore size was appropriate for chondrocyte ingrowth. The chondrocyte evaluation based on the morphology, and quantitative assessment such as the total protein and DNA, the semi-quantitative comparison of the expression of collagen II and aggrecan mRNA on NF scaffolds and SW scaffolds, indicated that PCL-b-PLLA NF scaffolds had the greatest potential in maintaining chondrocyte phenotype and accumulating more cartilage ECM contents. It was because that the nano-fibrous structure mimicked the physical structure of natural ECM, which had a high specific surface area, high protein adsorption capacity and an appropriate mechanical strength.

Conclusions

In this study, we documented a 3D nano-fibrous scaffold fabricated from PCL-b-PLLA (50/50), a semicrystalline diblock copolymer, by L-L TIPS. Uniform networks were created at low gelation temperatures, while high gelation temperatures led to knot-like structures and even continuous smooth pellicles among the nano-fibrous network. Moreover, macroporous structures were created by combining with salt-leaching technique with TIPS. Compared with SW scaffolds, the NF scaffolds had a larger specific surface area, higher protein adsorption capacity and a higher compression modulus. Chondrocytes exhibited a rounded shape, enhanced viability and proliferation on the NF scaffold, compared with the SW scaffold. Further studies are underway to investigate the effects of the scaffold morphology on chondrocyte function as well as the influence of the mechanical stimulation of the PCL-b-PLLA nano-fibrous scaffold on chondrocyte function.

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References


