Development of a Porous Scaffold-Manufacturing Method by Blending Silk Fibroin and Agarose Polymer Solutions

Seung-Won Park*, HaeYong Kweon, Tae-Won Goo, Seong Ryul Kim, You-Young Jo, and Gwang-Ho Choi
Department of Agricultural Biology, National Academy of Agricultural Science, Rural Development Association, Suwon 441-100, South Korea

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Low-melting-temperature agarose gel solution, as a novel porogen was combined with a silk fibroin solution to generate interconnected porous networks. The porosity of the resultant silk fibroin-agarose scaffolds was greater than that of the scaffolds generated with agarose and deionized water. The porosities of silk fibroin scaffolds containing agarose gel at 0.5%, 1.0%, 1.5%, 2.0% [w/v] were 110.9%, 111.7%, 120.9%, and 123.0%, respectively. Lastly, the internal space generated in scaffolds after dissolution of the agarose gel provides a good environment for cell growth and movement within the scaffold.

Key words: Silk fibroin, Agarose hydrogel solution, Scaffold, Tissue engineering

Introduction

The microenvironment of three dimension (3D) scaffolds is important for cell attachment, proliferation, and penetration within the scaffold. Consequently, much of tissue engineering research has focused on the utilization of porous sponge structures, owing to their microenvironment and adaptability to various complicated morphologies (Makaya et al., 2009). Sofia et al. (2001) first described the microstructure of a silk porous scaffold. Using salt-leaching methods, the authors developed scaffolds varying in porosity by altering salt solution concentrations and salt particle sizes (Kim et al., 2005). Although the procedure was successful in generating larger size porous silk scaffold by using salt particles of 1,000 µm, it was inadequate for use with salt particles less than 500 µm (Makaya et al., 2009).

Furthermore, scaffolds irregular internal pore sizes cause technical difficulties in the establishment of accurate external structures and, subsequently, cell culture systems. Recent studies have explored scaffold-manufacturing, techniques in an attempt to maintain acceptable microenvironment properties. Sarazin et al. (2003; 2004) developed a novel method for the generation of ultra-porous scaffolds by melt blending 2 or 4 immiscible polymers (Virgilio et al., 2010). Roy et al. (2006) deposited polymer/protein layers onto scaffold surfaces to increase the porosity of the materials. Virgilio et al. (2005; 2009; 2010) and Horiuchi et al. (1997) reported partial wetting in a significant number of scaffolds generated by ternary immiscible polymers blends.

In the present study, by melt blending of solutions derived from silk fibroin and agarose polymers, we developed a novel manufacturing method for fully interconnected porous scaffolds. Low-melting-temperature agarose gel solution-, used as a novel porogen, was combined with silk fibroin solution to generate the ultra-porous 3D structure of scaffold. While agarose gel dissolves in warm water, silk fibroin scaffold does not. As a result, in silk fibroin-agarose scaffolds generated using a melt processing technique, the large internal spaces within the scaffolds contain dissolved agarose. Porosity and interconnected porous networks of silk sponge scaffolds can be regulated by altering the ratio of silk fibroin solution to agarose gel solution. Lastly, the internal space generated in scaffolds after dissolution of -agarose gel provides a good environment for cell growth and movement within the scaffold. Since silk and agarose are derived from natural products, they are biocompatible with the human body. Silk fibroin-agarose scaffolds, therefore, are non-toxic,
as they will not cause a foreign body immune response that other synthetic polymers can induce.

Materials and Methods

Materials
Low-melting-temperature agarose, SeaPlaque® Agarose, was provided by Lonza Corp. (Rockland, ME). The agarose melts with distilled water under 65°C and has a gelation temperature from 26°C to 30°C. Cocoons of *Bombyx mori* silkworm were obtained from National Academy of Agricultural Science (Suwon, Korea). Silk was boiled off for 30 min in an aqueous solution of 0.02 M Na₂CO₃ and rinsed thoroughly with water to produce silk fibroin proteins.

Preparation of silk fibroin solution
Purified silk fibroin was dissolved in a CaCl₂/EtOH/water solution for 5 h. The solution was dialyzed in water for 72 h using cellulose membrane (molecular cutoff = 12,000~14,000) at room temperature.

Aqueous-derived agarose polymer preparation
To prepare a 100 ml stock of 3.0% agarose stock solution, add 3.0 g agarose to 100 ml sterile deionized water in a 250 ml Pyrex media bottle, cap loosely and completely dissolved in the microwave. Mix well as soon as the microwave cycle has finished (being careful to avoid splashing of the hot liquid) and keep with 200 ml of deionized water in the water bath at 50ºC. Before silk fibroin-agarose polymer solution blend, 3.0% agarose stock solution was diluted with deionized water at weight ratios of 0.5, 1.0, 1.5, and 2.0% [w/v].

Preparation of silk fibroin-agarose polymer solution blend scaffold
Aqueous-derived silk fibroin solution (10%) was mixed with sterile deionized water or different concentrations of agarose solution at a ratio of 1:1 [v/v]. To cast the scaffolds, the mixed solutions were poured into a 24-well tissue culture plate (2 ml/well) and allow the plates to cool and keep at 4°C to solidification of agarose. The plates dried at ~80°C in a freeze-drying oven for 72 h. The resultant scaffolds were incubated with absolute ethanol for 1 h in order to induce insolubility of silk fibroin against water. To remove all of residual agarose molecules, a used agarose molecules could be melts with water under 65°C. In the inmersing step, the scaffolds were immersed in the 50 ml cap tube filled with deionized water and incubated to water bath at 60°C for 2 days. The internal deionized water of 50 ml cap tube was changed with new water (total 6 times for 2 days).

Scanning electron microscopy (SEM)
The morphology of the scaffold had been analyzed by a scanning electron microscope (Hitachi S-3500, Japan). SEM was done at Regional Research Center, Myongji University, South Korea.

Swelling test
To determine the water absorption efficiency of scaffolds, swelling studies were performed according to the procedures in the previous study (Mathew *et al.*, 2010). Briefly, scaffolds were immersed in Hexane and temperature of 37°C. The dry weight of the scaffold was noted (Wo). Scaffolds were placed in Hexane and after a predetermined time (24 h) scaffolds were taken out, surface adsorbed water was removed by filter paper and wet weight was recorded (Ww). The ratio of swelling was determined using the following formula: Swelling ratio = (Ww – Wo)/Wo.

Results and Discussion

Development of agarose-silk fibroin scaffolds
For the generation of ultra-porous scaffolds, an aqueous-derived silk fibroin solution was mixed with various concentrations of an agarose hydrogel solution, the porogen, at a ratio of 1:1 (v/v). As shown in the schematic diagram in Fig. 1, the 2 polymer blends were dissolved in an aqueous solution. Water, therefore, was present between silk fibroin and agarose molecules. Because upon water evaporation, the aqueous-derived mixed solution could parti-

![Fig. 1. A schematic diagram representing the development of silk fibroin-agarose scaffold. Silk fibroin and agarose polymer blends were dissolved in an aqueous solution. Water, therefore, was present between silk fibroin and agarose molecules. Fully interconnected porous scaffolds were prepared using the 2 immiscible polymers and a blending technique. Silk fibroin solution derived from *B. mori* cocoons was mixed with different concentrations of agarose hydrogel solution at a ratio of 1:1 (v/v).](image-url)
tion, we performed a freeze-drying step during scaffold preparation. We assumed, however, that agarose molecules would remain attached to the silk fibroin scaffolds. To remove the residual agarose molecules, after freezing-drying treatment, we immersed in warm deionized water for 48 h. The portion of internal space in the resultant scaffolds was greater than that before the immersion step, and the internal space could be regulated by altering the ratio of silk solution to agarose solution.

Morphological analysis of scaffolds

Fully interconnected porous scaffolds were constructed in 24-well tissue culture plates by using 2 different types of polymers and a blending technique. Briefly, a silk fibroin solution derived from *B. mori* cocoons was mixed with varying concentrations of agarose hydrogel solution, the porogen, at a ratio of 1:1 [v/v] (Fig. 1). Prior to agarose leaching, the scaffold was in the shape of a cylinder, with a diameter of over 1.2 cm and a height of approximately 1.0 cm (Fig. 2). There was no change in scaffold size after freeze-drying (Fig. 2). To determine if the presence of water between silk fibroin and agarose molecules could influence the structure of scaffolds, we created sponge-type scaffolds. A agarose leaching and freeze-drying of the scaffolds demonstrated that the agarose gel act as a porogen, creating an internal ultra-porous structure within the scaffold. The sponge like scaffolds absorbed water, their shape did not change (Fig. 3 A and B). Micro-structure characterization of both the silk fibroin-agarose and pure scaffolds was performed with scanning electron microscopy (SEM). SEM image of the scaffold internal microenvironment revealed that the pores in agarose hydrogel mixed silk scaffold were much larger than the pores of pure scaffold (Fig. 3C).

Morphologically, we observed the porous structure of silk fibroin-agarose blended scaffolds. The volume of the two polymer blended scaffolds were measured and compared to the pure silk scaffolds. The described silk fibroin-agarose scaffold development technique is novel scaffold-manufacturing procedure with 2 benefits: (1) economically, it is useful because of the inexpensive nature of the materials used, and (2) immunologically, because warm water substitutes the need for other, harmful solvents in scaffold generation, the resultant scaffold will not stimulate an immune reaction.

Internal capacity analysis of scaffolds

Liquid intrusion analysis was used to determine the porosity of scaffolds, containing various concentrations of hydrogel solution (Soliman et al., 2011). Aqueous-derived silk fibroin solution was mixed with deionized water or agarose hydrogel solution, the porogen, at a ratio of 1:1 [v/v]. The porosity was greater in the silk fibroin solution mixed scaffold than in the deionized water mixed scaffold (Fig. 4A). The porosity of the silk fibroin-agarose scaffolds increased with increasing hydrogel solution concentration [w/v] (Fig. 4B). The scaffolds were arbitrarily assigned to 4 groups differing in agarose hydrogel solution concentration [w/v]. The porosity of hydrogel silk scaffolds was greater than that of silk pure scaffold. Based on normalization to pure silk scaffold porosity, the poros-
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Fig. 4. Porosity analysis of silk fibroin-agarose scaffolds. A. The porosity shows a higher increase with the silk fibroin solution mixed scaffold than deionized water mixed scaffold. B. The scaffolds were arbitrarily assigned into four groups based on the concentration of agarose hydrogel solution [w/v]. Compared with pure silk scaffold, porosity of hydrogel mixed scaffolds were shown higher.

The porosity of scaffolds containing agarose gel were 110.9% on 0.5% [w/v] agarose, 111.7% on 1.0% [w/v] agarose, 120.9% on 1.5% [w/v] agarose, and 123.0% on 2.0% [w/v] agarose. Therefore, 1.5% agarose was sufficient to generate ultra-porous silk scaffold.

The microenvironment of 3D scaffolds is important for cell attachment, proliferation, and penetration within the scaffold. Previously, regulation of pore size within scaffolds was induced using salt leaching methods (Makaya et al., 2009). Although the regulation of porous structure by using salt particles of varying size has its advantages, it does not overcome issues pertaining to the integration of fully interconnected porous networks with symmetric channels into scaffolds, which are required to guide the migration of cells into internal pores. Recently, alternatives to salt were used as porogens in the development of scaffolds. Sarazin, Favis (2003) and Virgilio et al. (2010) developed a new method for generating porous Poly-L-Lactide Acid (PLLA) scaffold by blending immiscible polymers. The resultant porous scaffold structure had a fully interconnected porosity and a highly controlled morphology (Virgilio et al., 2010). Although this method can be successfully applied to the development of porous 3D scaffolds, we argue that it is necessary to reduce chemical scaffold manufacturing, including the use of chemical solvents, as a protective measure against immune response.

Our novel silk fibroin-agarose scaffold-manufacturing method uses only biomaterials derived from natural products, thus eliminating the probability of stimulating an immune reaction. Furthermore, additional steps in scaffold development are unnecessary, because the agarose-leaching treatment essentially replaces salt leaching.

Lastly, the observed increase in interconnectivity between pores in our silk fibroin-agarose scaffolds will like aide in guiding the migration of cells into internal pores. In summary, the main benefits of our novel scaffold-manufacturing method are as follows: (1) the 2 polymers used are biocompatible with the human body, (2) the technique for the removal of agarose molecules is relatively simple, (3) the interconnected internal space within scaffolds is uniform and can be regulated by altering the ratio of silk solution to agarose solution, and (4) the combination of salt and agarose hydrogel solution can aid in the development of highly porous and ultra-interconnected scaffold structures.

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