Effects of phosphate ion concentration on in-vitro fibrillogenesis of sturgeon type I collagen

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Abstract

Nonmammalian collagens have attracted significant attention owing to their potential for use as a source of cell scaffolds for tissue engineering. Since the morphology of collagen fibrils controls cell proliferation and differentiation, its regulation is essential for fabricating scaffolds with desirable characteristics. In this study, we evaluated the effects of the phosphate ion (Pi) concentration on the characteristics of fibrils formed from swim bladder type I collagen (SBC) and skin type I collagen (SC) from the Bester sturgeon. An increase in the Pi concentration decreased the fibril formation rate, promoted the formation of thick fibrils, and increased the thermal stability of the fibrils for both SBC and SC. However, the SBC and SC fibrils exhibited different fibril formation rates, degrees of fibrillogenesis, morphologies, and denaturation temperatures for the same reaction conditions. Finally, by regulating the Pi concentration, various types of SBC and SC fibrils could be coated on cell culture wells, and fibroblasts could be cultured on them. The results showed that thin fibrils enhance fibroblast extension and proliferation, whereas thick fibrils restrain fibroblast extension but orient them in the same direction. The results of this study suggest that SBC fibrils, which exhibit diverse morphologies, are suitable for use as a novel scaffold material, whose characteristics can be tailored readily by varying the Pi concentration.
Keywords: Sturgeon type I collagen; Phosphate ion; Fibril formation; Fibril morphology;

Fibroblast morphology
1. Introduction

Type I collagen is the primary component of the extracellular matrix in skin, bone, and tendon [1]. Once its molecules have been secreted from cells, they spontaneously self-assemble into fibrils, which subsequently form thick fibers through a series of enzymatic and chemical reactions [2]. Collagen fibrils and fibers provide support for cell growth and ensure the mechanical resilience of connective tissues [3–4].

Under appropriate reaction-fluid conditions, extracted type I collagen molecules can form fibrils with structural, mechanical, and biological properties similar to those of native fibrils in vivo [5–7]. Because of their low antigenic activity, desirable cell adhesion properties, biocompatibility, and biodegradability, type I collagen fibrils have high potential for use as biomedical materials, such as cellular scaffolds for tissue engineering [8, 9]. When these fibrils are used as collagenous scaffolds, their morphological characteristics, including their diameter, shape, and orientation, have a determining effect on the adhesion, alignment, proliferation, and differentiation of anchorage-dependent cells [1, 10]. Therefore, the optimization of fibril morphology in scaffolds is essential for ensuring desirable cellular performance for various applications in tissue engineering.

Previous studies have shown that the ionic strength of the reaction fluid influences the in-vitro formation of collagen fibrils [11–13]. For example, Wood [14] found that the rate
of fibril formation and fibril width decrease with an increase in the ionic strength of the reaction fluid; the authors used NaCl to regulate the ionic strength. Divalent anions also play a critical role in regulating fibril formation rate [15]; in particular, phosphate ions are known to bind to collagen molecules, significantly affecting the electrostatic interactions between the molecules and, hence, the process of self-assembly of the collagen molecules into fibrils [15–17]. Because in-vitro collagen fibrillogenesis is a complex process affected by several factors, such as the electrostatic and hydrophobic interactions that occur as well as the intermolecular force involved [16, 18], the mechanism through which the ionic strength affects collagen fibrillogenesis remains unknown. Therefore, a better understanding of the effects of the ionic strength on the fibril morphology will allow for the rational design of collagen-based scaffolds.

Currently, the most common collagen sources for biomedical use are bovine skin and tendon, porcine skin, and rat tail [19–21]. However, animal collagen-containing products can potentially include components that cause infectious animal diseases in humans, including bovine spongiform encephalopathy, foot-and-mouth disease, and African swine fever [22]. Religious beliefs also restrict the use of some animal products [23]. Therefore, alternative collagens extracted from marine sources, especially from fish byproducts, such as skin, scales, and bones, attracted increasing attention [20, 21, 24–26]. Our recent research has shown that
mammalian and fish collagen have different biochemical and physicochemical characteristics, including different amino acid compositions, thermal stabilities, and fibril-forming abilities [27]. Therefore, to take advantage of marine collagens as biomedical materials, it is necessary to elucidate their characteristics, especially their fibril-forming ability and the factors that determine the morphological characteristics of the fibrils formed in vitro. Compared to the case of mammalian collagens, however, there is limited knowledge regarding marine collagens and their fibrillogenesis properties.

In a previous study [27], we showed that the sturgeon contains a large amount of type I collagen in its byproducts, including its swim bladder and skin. The pepsin-treated collagen (atelocollagen) from these tissues reassembled into fibrils more quickly than mammalian collagen under specific conditions, suggesting that the former may be suitable for fabricating biomedical materials [27]. We also found that increasing the ionic strength of the reaction solution by adding NaCl sped up the fibrillogenesis process and promoted the formation of thicker fibrils. Notably, the swim bladder collagen assembled into a unique fusiform structure. These findings suggest that sturgeon collagens can potentially be used to form fibrils of different morphologies by controlling the reaction-fluid environment.

The objective of the present research is to clarify the regulation ability of the fibril morphology of sturgeon collagens to evaluate their potential as scaffold materials for tissue
engineering. Specifically, we elucidated the effects of the phosphate ion (Pi) concentration of
the reaction fluid on the in-vitro formation of fibrils of swim bladder type I collagen (SBC)
and skin type I collagen (SC). Other ion species, including cations, and the pH of the reaction
fluid may also affect the electrostatic interactions between the collagen molecules and, hence,
their self-assembly to fibrils. However, Pi is the first choice, since it has been proven to bind
to collagen molecules and regulate fibril formation [28]. As a first step, we evaluated the
effects of Pi concentration on the fibrillogenesis of SBC and SC because, to date, there has
not been any research on the influences of Pi on the fibrillogenesis of fish collagen. Based on
our findings, we successfully coated different forms of SBC and SC fibrils on cell culture
wells by regulating the Pi concentration of the reaction fluid and evaluated the proliferation of
fibroblasts on them along with the morphology of the fibroblasts.

2. Materials and methods

2.1. Isolation and purification of collagen

A Bester sturgeon (*Huso huso* × *Acipenser ruthenus*; body weight of 6.0 kg) was
procured from the Nanae Fresh-Water Laboratory, Field Science Center for Northern
Biosphere, Hokkaido University, Japan. Its swim bladder and skin were dissected, washed
with chilled tap water, lyophilized, and stored at -30 °C until use. Before the collagen
extraction process, the skin fat was removed during a period of 24 h using 99.5% ethanol (two solution changes) in a sample/solution ratio of 1/10 (w/v). Then, the swim bladder and defatted skin were cut into small pieces (ca. 0.5 × 0.5 cm) for collagen extraction. Extraction, purification, and freeze-drying were performed as per the procedures reported by Meng et al. [29].

2.2. In-vitro collagen fibrillogenesis

Collagen fibrillogenesis was performed according to the method of Meng et al [29], with modifications of the Na-phosphate buffer (PB) concentrations. Freeze-dried SBC and SC were dissolved in a HCl solution (pH 2.0) at a concentration of 0.3% (w/v). Subsequently, these SBC and SC solutions were mixed with a Na-phosphate buffer (pH 7.4) solution, having a Na-phosphate concentration of 22.5, 45, 90, or 180 mM, respectively; the collagen solution/PB ratio was set to 1/2 (v/v), which resulted in a final PB concentration of 15, 30, 60, or 120 mM, respectively. The mixtures were kept at 21 °C, and fibril formation was monitored during a period of 1 h in terms of the changes in turbidity based on the absorbance at 320 nm; this was done using a spectral monitor (Hitachi High-Technologies Corporation, Tokyo, Japan).
2.3. Degree of collagen fibrillogenesis

The degree of collagen fibrillogenesis was measured using the method proposed by Zhang et al. [27]. In brief, the reacted solution was centrifuged at 20,000×g for 20 min to precipitate the fibrils, and the protein content of the supernatant was measured based on the method reported by Lowry [30]; bovine serum albumin was used as the standard. The degree of collagen fibrillogenesis was defined as the percentage of decrease in the collagen concentration. In other words, it was defined as the percentage of collagen molecules that formed fibrils.

2.4. Morphology of fibrils formed

The microstructures of the SC and SBC fibrils were observed using a scanning electron microscopy (SEM) system (JSM6010LA, JEOL Ltd., Tokyo, Japan). The collagen fibrils were formed as described in Section 2.2 and obtained by centrifugation as described in Section 2.3. The precipitated fibrils were fixed with 2.5% (v/v) glutaraldehyde in 0.1 M PB (pH 7.4) for 3 h at room temperature, rinsed with PB to remove the fixative, dehydrated using a graded series of ethanol solutions, and soaked in a t-butyl alcohol solution for two 30-min intervals. Finally, the samples were freeze-dried in a t-butyl alcohol solution using a freeze-drying device (JFD-320; JEOL Ltd.) and coated with gold-platinum, using an auto fine
coater (JFC-1600; JEOL Ltd.). After the SEM observations, digital images of the samples were taken, and the diameters of 100 randomly selected fibrils were determined using the software ImageJ. The measurements were performed three times using different photographs.

2.5. Thermal stability of collagen fibrils

The maximum transition temperatures (Tm) of the reassembled fibrils of SBC and SC were measured by differential scanning calorimetry (DSC; EXSTAR DSC6100, SII Nano Technology Inc., Chiba, Japan). The collagen fibrils were prepared as described in Section 2.4, placed in an aluminum pan (70 µl), and DSC measurements were performed from 20 to 70 °C at a rate of 3 °C/min. An empty aluminum pan was used as reference. The Tm value was taken to be the temperature corresponding to the endothermic peak in the DSC curve.

2.6. Coating of collagen fibrils on cell culture wells

A 200 µl aliquot of a 1% (w/v) SBC or SC solution (pH 2.0) was placed in each cell culture well (24-well cell culture plate; FALCON, Durham, NC) and kept there for 30 min. Then, the solution in each well was replaced with 200 µl of PB (15, 30, 60, or 120 mM; pH 7.4), and fibril formation was induced for 1 h. After removing the PB solution, a PB solution with the same Na-phosphate concentration but containing 0.1 mM genipin as a crosslinker
was added, and the wells were stored for 12 h. Finally, the genipin-PB solution was removed, and the wells were washed twice with Hanks balanced salt solution, containing 100 U/ml penicillin and 100 μg/ml streptomycin, and immediately used for cell culture experiments. A diluted HCl solution (pH 2.0) was used instead of the SBC or SC solution as the control. All procedures were performed under aseptic conditions at 21 °C.

2.7. Morphology of collagen fibrils formed on cell culture wells

After the formation of the collagen-fibril coating, the bottom surface of each well was fixed, dehydrated, and freeze-dried by removing the solutions as described in Section 2.4. After the completion of the freeze-drying process, the bottom of each well was removed and cut into a 1×1 cm piece using a heated scalpel. The surface was coated with gold-platinum and observed using SEM.

2.8. Cell culturing

To study the cell morphology and proliferation on the SBC and SC collagen fibrils, we used L929 mouse fibroblastic cells maintained in Eagles minimum essential medium (Life Technologies Corporation, Grand Island, NY, USA) supplemented with a 5% fetal bovine serum (Thermo Fisher Scientific, Carlsbad, CA, USA) and a 1% penicillin/streptomycin
mixture (Thermo Fisher Scientific, Grand Island, NY, USA). The cells were cultured for 5 days in a humidified incubator at 37 °C and 5% CO₂; the medium was renewed after 3 days.

2.9. Cell morphology and proliferation

The morphology of the cells was observed over the cell-culture period using a phase-contrast microscope (DMI600B, Leica, Wetzlar, Germany). The microstructure of the cells 2 days after seeding was observed by SEM as described in Section 2.7.

A cell counting kit (CCK-8, DOJIN, Mashiki, Japan) was used to evaluate the cell-proliferation activity. After a 5-day culture, the culture medium was removed, and the culture was washed with PBS twice. Then, 540 μl of the culture medium and 60 μl of the CCK-8 solution were added to each well, and the cells were cultured for 30 min. The supernatant (100 μl) was transferred to a 96-well plate, and the optical density at 450 nm was determined using a microplate reader (infiniteF50R, Wako, Osaka, Japan). Six parallel replicates were read for each sample.

2.10. Statistical analysis

The data on the degree of fibril formation and the proliferation of L929 cells are expressed as mean ± standard error. Significance was set to p < 0.01. The data were analyzed
using ANOVA and Tukey–Kramer post-hoc tests for multiple comparisons, which were performed using a statistical add-in for Microsoft Excel (SSRI, Tokyo, Japan).

3. Results

3.1. Collagen fibrillogenesis

The effects of the Pi concentration on the progression of SBC and SC fibril formation in vitro are shown in Fig. 1. When the Pi concentrations of the PB solution were 15 and 30 mM, the initial turbidity in the case of SBC was higher than 0.5 and increased sharply without a lag phase. As the Pi concentration increased to 60 mM, however, a short lag phase appeared before the growth phase. At a Pi concentration of 120 mM, the turbidity curve was sigmoidal and contained an extended lag phase and a more gradual growth phase. The turbidity in the plateau phase was also affected by the Pi concentration: it was highest at 60 mM, followed by the turbidity at 120, 30, and 15 mM. The same trend was observed in the turbidity curves for SC. However, no increase in the turbidity was observed at a Pi concentration of 120 mM. In addition, for SC and a Pi concentration of 60 mM, the turbidity in the plateau phase was lower than that for SBC.

3.2. Degree of fibrillogenesis
The degree of fibrillogenesis was assessed after 1 h (Fig. 2). For both SBC and SC, the degree was higher than 95% when the Pi concentration was less than 30 mM. Further, it decreased to 89% for SBC and 60% for SC when the Pi concentration increased to 60 mM. At a concentration of 120 mM, the fibrillogenesis degree decreased to 64% in SBC, while no collagen fibrils were formed in SC, which conformed with the SC turbidity curve.

3.3. Morphology of fibrils

SEM images of the SBC and SC fibrils formed after incubation for 1 h are shown in Fig. 3, while the diameter distributions of the fibrils are shown in Fig. 4. Hereafter, the fibrils are referred to on the basis of the origin of the collagen used (swim bladder or skin) and the Pi concentration at the time of fibrillogenesis (15–120 mM). Hence, SBC-15 refers to the fibrils formed using SBC and a Pi concentration of 15 mM. At the lowest phosphate concentration (15 mM), thin fibrils curled together in high density were formed in both SBC and SC (Fig. 3). The mean diameters were 77.7 nm for SBC-15 and 78.8 nm for SC-15, and the fibril diameter distributions were narrow (Fig. 4). As the Pi concentration increased, thicker fibrils appeared for both SBC and SC (Fig. 3). As a result, the range of the fibril diameters also increased (Fig. 4). The largest mean diameters were observed in the cases of SBC-60 (397 nm) and SC-60 (238 nm). In addition, the SBC-30 preparation included thick but short fusiform structures.
At a Pi concentration of 60 mM, the fusiform structures became larger and were characterized by a transverse periodic banding pattern (Fig. 3). These fusiform structures were interlaced with thin fibrils, with a maximum diameter of more than 1600 nm (Fig. 4). In SBC-120, the fusiform structures disappeared. However, the mean fibril diameter was higher than that of SBC-60 (Figs. 3 and 4). In the SC fibrils, no fusiform structures were observed, but thick fibril bundles were present in SC-30 (Fig. 3). In contrast to the case for the fusiform structures in SBC, these bundles were probably formed by the spiral assembly of the thin fibrils (Fig. 3). Finally, these bundles were not present in SC-60. However, the diameter of the fibrils was higher than that of the fibrils in SC-30 (Figs. 3 and 4).

3.4. Thermal stability of collagen fibrils

The DSC curves of the SBC and SC fibrils are shown in Fig. 5. The $T_m$ value of the SBC fibrils increased as the Pi concentration increased to 60 mM but decreased once the Pi concentration was 120 mM. The values for SBC-15, SBC-30, SBC-60, and SBC-120 were 42.9, 43.3, 44.1, and 40.4 °C, respectively. Further, for the same Pi concentration, the $T_m$ values of the SC fibrils were lower than those of the SBC fibrils, with the values of SC-15, SC-30, and SC-60 being 37.7, 40.7, and 44.7 °C, respectively.
3.5. Collagen-fibril coating on cell culture wells

In this study, a two-dimensional collagen-fibril coating was formed in cell culture wells to study the cell growth and proliferation characteristics of the different collagen fibrils. SEM images of the SBC- and SC-fibril coatings are shown in Fig. 6. Both the SBC and SC fibrils were aligned along the same direction on the bottoms of the culture wells. Although the trend in the Pi-dependent changes in the fibril diameter was the same as that for the fibrils formed in solution, the fibrils coated on the wells were thinner. In addition, the SBC fibrils exhibited greater variation in their diameter with changes in the Pi concentration of the buffer. The fusiform structures were observed only in SBC-60. Finally, the bundle structures were not observed in the SC fibrils.

3.6. Cell morphology

Figure 7 shows the morphologies of the L929 cells grown on the SBC and SC fibrils. Six hours after seeding, the cells on all the fibrils were highly stretched, except for those on the SBC-60 and -120 fibrils. In the cases of the SBC-60 and SBC-120 fibrils, the cells were attached firmly but were round and did not spread. Twenty-four hours after seeding, the number of cells increased. This was true for all the cases. Further, the cells spread randomly on the control surface and the fibrils, apart from those in the SBC-60 and -120 samples. On
the SBC-60 and -120 fibrils, in contrast, cells began to spread in the same direction, parallel
to the fibrils.

To further examine cell morphology, the cells were imaged using SEM 2 days after
seeding (Fig. 8). The cells on the control surface were flat and irregular. In contrast, the cells
on both the SBC and SC fibrils had a higher cell height (thickness). Furthermore, many of the
cells exhibited two long protrusions into the counter direction. All the cells on the SC fibrils
were stretched far more than those on the SBC fibrils and irregularly oriented. Similarly, the
cells on the SBC-15 and -30 fibrils were also misaligned, while those on the SBC-60 and -120
fibrils displayed a higher degree of alignment along the fibril direction.

3.7. Cell proliferation

Cell proliferation was assessed on day 5, using a CCK-8 kit (Fig. 9). The cells on the
fibril-coated wells proliferated at a significantly lower rate than those on the control wells.
The cells on the SC fibrils showed a higher proliferation activity than those on the SBC fibrils.
Moreover, no significant differences were observed between the SC-15, -30, and -60 groups.
Cell proliferation on the SBC fibrils decreased with an increase in the Pi concentration from
15 to 60 mM and was lowest on the SBC-60 fibrils and highest on the SBC-120 fibrils.
4. Discussion

The results obtained in this study clearly showed that the Pi concentration in the reaction fluid significantly affects the rate of *in-vitro* fibril formation as well as the morphology of the fibrils formed. This inference is true for both SBC and SC. There have been a few studies on the fibrillogenesis of type I collagen and the morphology of the fibrils formed *in vitro* in fish collagens, such as those from salmon skin [26], shark skin [31], red stingray skin [32], Bester sturgeon skin, and swim bladder [27]. However, these studies focused on the influence of monovalent ions only, such as Na\(^+\) and Cl\(^-\). Because divalent anions have a higher order in the Hofmeister series, they have a greater effect on macromolecules and larger ions [33]. A few studies on fibrillogenesis in mammalian collagens found that divalent anions have a more pronounced effect than monovalent anions [15, 34]. To the best of our knowledge, the present study is the first to elucidate the effects of the Pi concentration on the fibrillogenesis process and the morphology of the fibrils formed *in vitro* in fish collagens. As known from previous studies, ionic strength also affects collagen fibrillogenesis [11–14]. Therefore, Pi concentration and ionic strength may work together to change collagen fibrillogenesis in the present study.

Collagen fibrillogenesis is a complex process of molecular assembly, which can be studied based on the changes in turbidity corresponding to three phases: a lag phase, wherein
there is no change in the turbidity; a growth phase, which involves a rapid increase in the turbidity; and a plateau phase, during which the turbidity again remains unchanged.

Ultrastructural studies on fibrillogenesis using atomic force microscopy have revealed that the lag and growth phases consist of several multistep processes [35, 36]. During the lag phase, multiple coassemblies of several collagen molecules (nucleation) form and their nuclei subsequently undergo linear aggregation to produce subfibrillar units [37]. Then, these subfibrillar units aggregate laterally to form fibrils, which then grow into bundles during the growth phase [38]. Silver et al. [39] reported that fibril formation occurs at the end of the lag phase and that only the mass of the fibrils increases during the growth phase. The subfibrillar units have a determining effect on the mechanical properties of collagen fibers [37].

In this study, the addition of Pi significantly prolonged the lag phase of the fibrillogenesis process in both SBC and SC. At Pi concentrations of 15 and 30 mM, the initial turbidities were higher than those at 60 and 120 mM; this was true for both SBC and SC, suggesting that the lag phase was shorter than the time taken for the first absorbance measurement (less than 5 s after the mixing of the collagen solution and PB). As the Pi concentration increased, the lag phase became longer. Previous studies on mammalian collagens have indicated that phosphate ions direct the linear and lateral growth of collagen molecules [15, 38, 39]. Therefore, we can conclude that one of the primary effects of the Pi
concentration on fibril formation in SBC and SC is related to subfibrillar unit formation during the lag phase. A shorter lag phase was observed at lower Pi concentrations, suggesting that the nucleation and linear aggregation of the nuclei into subfibrillar units was accelerated in these cases. Under such conditions, numerous small subfibrillar units may form within a short period. As the Pi concentration increased, the lag phase became longer, and a few larger subfibrillar units were produced slowly. Hayashi [15] reported that the lag phase during polymerization is a unique feature of the assembly mechanism. Hence, the Pi concentration probably has a determining effect on these molecular interactions of collagen.

Collagen fibril formation is a complex process operating under many secondary forces, such as electrostatic interaction, hydrogen bonding, or salt bridging. Some studies on mammalian collagens suggested that the Pi concentration has a significant effect on the electrostatic interactions that occur between collagen molecules [15–17] and that the assembly of subfibrillar units is primarily controlled by these electrostatic interactions [15]. For example, Li et al. [16] found a strong influence of Pi on the surface charge of collagen molecules, which would change the electrostatic repulsion and regulate the aggregation of such molecules. This process mainly occurred in the lag phase. Thus, the Pi concentration probably affects the lag phase during SBC and SC fibril formation through electrostatic interactions occurring between SC and SBC molecules, respectively.
The growth phase involves the lateral aggregation of the subfibrillar units to form higher-molecular-weight fibrils. The slope of the turbidity curve reflects the rate of lateral aggregation, while the final turbidity is reflective of the fibril diameter [11]. Zhu and Kaufman [40] reported that the increase in turbidity during the growth phase is primarily attributable to an increase in the fibril diameter. In the present study, as the Pi concentration increased, the slope corresponding to the growth phase decreased, and the turbidity was high. These results suggest that the larger subfibrillar units that are formed at higher Pi concentrations require more time to aggregate and form thicker fibrils. The results of the SEM observations of the fibrils supported this hypothesis, namely, that a higher Pi concentration in the growth environment induces the formation of thicker fibrils with an elongated morphology. This tendency was observed in both SBC and SC but more evident in the former. As was also the case in the present study, Hayashi et al. [15] examined fibril formation in calf skin collagen for Pi concentrations of 50–300 mM and found that the fibril formation rate decreased with an increase in the Pi concentration. Further, Williams et al. [41] reported that Pi concentrations lower than 30 mM had negligible effect on the fibril formation rate in rat tendon collagen. In contrast, the formation rate decreased for Pi concentrations higher than 30 mM. However, few studies have directly described the relationship between fibril morphology and Pi concentration.
Different types of thick fibril-based structures were formed in SBC and SC at various Pi concentrations. Fusiform fibril structures grew in SBC at Pi concentrations of 30 mM and 60 mM. In contrast, thick bundles of fibrils formed in SC at a Pi concentration of 30 mM. These results indicated that the processes of fibril formation were different in the type I collagens obtained from the swim bladder and skin of the sturgeon. In a previous study, we found that the hydroxyproline (Hyp) content of SBC is higher than that of SC [27]. Since the hydroxyl groups of Hyp form hydrogen bonds to maintain the structure and stability of collagen, the difference in the Hyp contents of SBC and SC may be one of the reasons for different structures forming in the two types of collagens. This difference in the Hyp contents of SBC and SC may be attributable to a difference in the composition of the \( \alpha \)-chains in the collagen molecules, as is characteristic in fish. The molecules of type I collagen from mammals form a triple helical heterotrimer with two identical \( \alpha_1(I) \) chains and one \( \alpha_2(I) \) chain and can be expressed as \([\alpha_1(I)]_2\alpha_2(I)\). Although minor collagen molecules, including those of type V, are usually found with type I collagen, the composition of the type-I \( \alpha \) chain in different tissues is the same. In the molecules of the type I collagen found in fish, however, another type of \( \alpha \)-chain, named \( \alpha_3(I) \), is present [42]. Therefore, the type I collagen molecules in fish are a mixture of \([\alpha_1(I)]_2\alpha_2(I)\) and \(\alpha_1(I)\alpha_2(I)\alpha_3(I)\), with the ratio of the two types of molecules being specific to each tissue. Zhang [43] reported that the mRNA expression levels
of the α1(I) and α2(I) chains in the swim bladder and skin of the Amur sturgeon are different. Thus, a difference in the ratios of α1(I), α2(I), and α3(I) of SBC and SC could be the primary reason for the difference in the Hyp contents.

The diameter ranges of the fibrils formed during the *in-vitro* fibrillogenesis experiments in SBC and SC were 50–1600 nm and 50–350 nm, respectively. The thicknesses of collagen fibrils *in vivo* are 0.5–3 µm [44, 45]. This suggests that, at the right Pi concentration, sturgeon collagen can form fibrils with diameters similar to those of the fibrils formed *in vivo*. Hence, sturgeon collagen can potentially be used as a biomedical material. In addition, the T_m value of the fibrils increased with the Pi concentration. In one of our previous studies, we found that the concentration of NaCl showed no effect on the thermal stability of the fibrils [46]. This indicates that the effects of phosphate ions on collagen fibrils are different from those of NaCl. The precise mechanism responsible for this increase in T_m remains to be clarified. However, it is likely that the structural differences between the fibrils are responsible for the observed T_m values.

The present study also showed that higher Pi concentrations inhibit the number of fibrils formed over a period of 1 h (Fig. 2). This effect was more pronounced in SC than in SBC, with no SC fibrils forming at a concentration of 120 mM. It is possible that, as the Pi concentration increased, more collagen molecules undergo linear aggregation to form large
subfibrillar units, owing to a decrease in the degree of lateral aggregation. The subfibrillar units were probably too narrow to affect the turbidity and could not be collected by centrifugation. Currently, performing in-situ observations of the developing subfibrillar units remains a challenge because a technique with a molecular-level resolution is required. The development of such techniques will improve our understanding of the mechanisms by which the Pi concentration affects fibril morphology.

We further attempted to use the SBC and SC fibrils as cellular scaffolds for tissue engineering, since collagen fibrils are the primary scaffolds in vivo. Therefore, we coated SBC and SC fibrils on cell culture wells and examined the responses of fibroblastic L929 cells in terms of their morphology and proliferation. The L929 cells seeded on the control wells had irregular polygonal shapes and proliferated rapidly in a tilted manner on the bottom surfaces of the wells. In contrast, the cells seeded on the collagen fibrils had higher cell height (larger cell thickness), and many of them extended two long cellular processes in the opposite direction. Notably, the cells attached to the SBC-60 and -120 fibrils spread the cellular processes along the fibrils.

This type of cell morphology can be attributed to the specific signal transduction from the fibrils, which passed through the integrins and transformed the cellular receptors to type I collagen fibrils [47]. Generally, the signaling pathways downstream of the integrins
regulate different aspects of cell behavior, such as cell morphology, orientation, proliferation, and differentiation [48–50]. According to Friedrichs et al. [44], cells expressing integrins are polarized strongly along collagen fibrils. In contrast, cells lacking integrins attach to the substrates but do not exhibit polarization.

We were also able to determine the appropriate fibril diameter range for cell polarization. Because the cells in some human tissues, such as bone and tendon, are arranged in parallel arrays [51], being able to regulate the cell direction is essential for tissue engineering. Thus, the SBC-60 and -120 fibrils have high potential for use as scaffolds for the engineering of tissues consisting of aligned cell arrays.

The cell proliferation rate on the SBC and SC fibrils was lower compared to that on the control (noncoated) surface. On the SBC fibrils, effects of fibril diameter on the proliferation activity were apparent: as the fibril diameter increased, the cell proliferation activity decreased, with the activity being lowest at a Pi concentration of 60 mM. Thus, the cells probably receive different signals for proliferation from fibrils having different diameters.

Artavanis-Tsakonas [52] found that signal transmission between cells also affects cell proliferation. Thus, it is possible that the high degree of cell alignment on the SBC-60 and -120 fibrils causes the transmission of different signals, resulting in a decrease in proliferation.

Mouse MC3T3E1, ATDC5, 3T3-L1, and C2C12 cells also exhibit reduced proliferation on
SBC fibrils, especially when the fibril diameter is high [53]. While the exact reason for this phenomenon is yet unknown, it suggests that the inhibitory activity of SBC fibrils is not specific to L929 cells. This study has a limitation in that the effects of SBC fibrils on cellular differentiation were not determined. Since there is a general trade-off relationship between cell proliferation and differentiation, SBC fibrils may also affect the latter. Future studies on the differentiation of cells on SBC fibrils will further clarify the potential of SBC fibrils as cellular scaffolds. In addition, the present method is not applicable to the standard three-dimensional cell culture system containing cells in the collagen gel. Since the standard method seeds the cells in a mixture of collagen and the cell culture medium and, thereafter, induces gelation, the high Pi concentrations used for the regulation of collagen fibrillogenesis in this study may have had some adverse effects on the cells. Future studies on the application of the present method to the three-dimensional scaffold and cell culturing are essential.

5. Conclusions

In this study, the regulation ability of the fibril morphology of sturgeon collagens was clarified to demonstrate their potential as scaffold materials for tissue engineering. It was found that the Pi concentration has a determining effect on the morphology as well as the rate and degree of fibrillogenesis and the thermal stability of the formed fibrils. At low Pi
concentrations, thin fibrils with a low thermal stability were formed. However, as the Pi concentration increased, the fibril formation rate decreased, and thick fibrils with higher thermal stability were formed. Moreover, the morphology of the fibrils determined the morphology and proliferation of the fibroblasts grown on them. We could successfully coat SBC and SC fibrils of different morphologies on the bottoms of cell culture wells and found that, while thin fibrils are better for cell extension and proliferation, thick fibrils limit cell extension and proliferation but regulate cell orientation. Given that there have been no previous studies on the effects of Pi concentration on the characteristics of fish collagen fibrils, this study is the first to highlight the possibility of Pi-induced regulation of the morphology of sturgeon collagen fibrils from a tissue engineering viewpoint. In particular, SBC is suitable for use as a scaffold material, because SBC fibrils are rich in sturgeon by-products [27], quicker to assemble [27, 29], retain high Tm (higher than 37 °C ), are affected by the Pi concentration to a greater degree, and exhibit a more diverse morphology as compared to SC fibrils. Future studies on the differentiation of cells on SBC fibrils will further clarify the potential of SBC fibrils as cellular scaffolds.

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References


Figure captions

Fig. 1. Effects of phosphate ion concentration on in-vitro formation of fibrils of (A) swim bladder and (B) skin collagens of Bester sturgeon.

Fig. 2. Effects of phosphate ion concentration on the degree of fibril formation of (A) swim bladder and (B) skin collagens of Bester sturgeon. Columns and bars represent mean and standard error for three experiments. Differences between groups with different letters are significant (Tukey–Kramer post-hoc test, p < 0.01).

Fig. 3. Scanning electron micrographs of swim bladder and skin fibrils formed at 21 °C for 1 h. Scale bars: 5 μm. Black and white arrows indicate fusiform and bundle structures in SBC and SC, respectively.

Fig. 4. Diameter distributions of swim bladder and skin collagen fibrils formed at 21 °C for 1 h.

Fig. 5. DSC curves of (A) swim bladder and (A) skin collagen fibrils formed at 21 °C for 1 h. Black arrowheads indicate maximum transition temperatures.
Fig. 6. Scanning electron micrographs of bottom surfaces of cell culture wells coated with swim bladder and skin collagen fibrils. Scale bars: 5 μm. Black arrows indicate fusiform structures in SBC-coated wells.

Fig. 7. Phase-contrast microscopy images of L929 cells cultured for 6 and 24 h. Scale bars: 75 μm.

Fig. 8. Scanning electron micrograph images of L929 cells cultured for 2 days. White arrows show the direction of collagen fibril alignment. Scale bars: 10 μm.

Fig. 9. Proliferation of L929 cells cultured for 5 days. Columns and bars represent mean and standard error for six measurements. Differences between groups with different letters are significant (Tukey–Kramer post-hoc test, p < 0.01).
Fig. 1

(A) Turbidity (320nm) over time (sec) for different concentrations (15 mM, 30 mM, 60 mM, 120 mM).

(B) Turbidity (320nm) over time (sec) for different concentrations (15 mM, 30 mM, 60 mM, 120 mM).
Fig. 2

A

B
Fig. 3

SBC 15 mM PB  
SC 15 mM PB

SBC 30 mM PB  
SC 30 mM PB

SBC 60 mM PB  
SC 60 mM PB

SBC 120 mM PB  

Fig. 4

(A) SBC 15 mM □ SBC 30 mM □ SBC 60 mM □ SBC 120 mM

(B) SC 15 mM □ SC 30 mM □ SC 60 mM
Fig. 7

6 hours

SBC 15 mM PB  SBC 30 mM PB  SBC 60 mM PB  SBC 120 mM PB

SC 15 mM PB  SC 30 mM PB  SC 60 mM PB  Control

24 hours

SBC 15 mM PB  SBC 30 mM PB  SBC 60 mM PB  SBC 120 mM PB

SC 15 mM PB  SC 30 mM PB  SC 60 mM PB  Control

SBC  15 mM PB
SBC  30 mM PB
SBC  60 mM PB
SBC  120 mM PB

SC  15 mM PB
SC  30 mM PB
SC  60 mM PB
Control

SBC  15 mM PB
SBC  30 mM PB
SBC  60 mM PB
SBC  120 mM PB

SC  15 mM PB
SC  30 mM PB
SC  60 mM PB
Control
Fig. 9