

Infiltration of Silica Inside Fibrillar Collagen**

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Diatom frustules (the hard and porous cell wall of diatoms) are created under the control of biomolecules (silaffins, silacidins, and long-chain polyamines) at close to physiologic conditions.^[1–4] The mechanism of biosilica formation was traditionally based on the ability of zwitterionic water-soluble proteins to create macromolecular assemblies for silica polymerization.^[5–7] Recent discoveries of water-insoluble collagen matrices within certain sponge biosilica spicules,^[8] chitin-based scaffolds in sponge and diatom biosilica formations,^[9,10] as well as cingulins within diatom girdle bands,^[4] revive the use of insoluble biomimetic organic templates for morphogenesis of nonporous silica structures. The use of fibrillar collagen as templates for biosilica synthesis was unsuccessful in the past as only extrafibrillar silica deposition was observed.^[11,12] Intrafibrillar mineralization of collagen has important implications from a biophysical perspective.^[13] Herein, we report a collagen biosilicification scheme based on

the fusion of stabilized polysilicic acid into a fluidic precursor phase upon their infiltration into polyamine-enriched collagen. The latter serves as a template and catalyst for polymerization of the precursor phase into silica that faithfully reproduces the collagen tertiary architecture. Our findings provide a new concept in biosilica materials synthesis, which does not require phosphate supplements.

Type I collagen has been widely used for hybrid biomaterials synthesis because of its biocompatibility.^[14,15] Nevertheless, the lack of mechanical resistance of highly porous, nonmineralized collagen matrices limits their application as stress-bearing scaffolds for bone repair.^[5] The potential stimulating effect of silicic acid on osteogenesis^[16,17] provides the incentive to produce silicified collagen matrices for bone regeneration applications. Previous studies have explored methods to produce silica–collagen hybrid materials by simultaneous self-assembly of collagen and silica. However, the kinetics of collagen self-assembly and silica polymerization are not fully compatible.^[18] Other studies that utilized fibrillar collagen as templates produced silica particles in close vicinity of the collagen fibrils but not within the fibrils, which limits the strength of the biohybrid materials.^[12] Silica–collagen hybrids may also be potentially used for preparation of nonbiological functional materials;^[19,20] however, the fundamental prerequisite is that collagen fibrils should be fully infiltrated with a high intrafibrillar silica content.

During diatom cell wall synthesis, silicic acid is stabilized and prevented from spontaneous polymerization into silica.^[21] Silicic acid gels rapidly in vitro at pH 5.5 without stabilization (see the Supporting Information S2, Figure S2 A and S3, Figure S3 A). Thus, we used choline chloride to stabilize silicic acid hydrolyzed from tetraethyl orthosilicate.^[22] Based on the assessment of gelling times, 72 mM choline chloride is required to stabilize 1.5 % silicic acid (pH 5.5) for 72 h before gelling (see the Supporting Information S2, Figure S2 A). This method provided sufficient time for the choline-stabilized silicic acid (Ch–SA) to infiltrate type I collagen sponges. The mean particle diameter of Ch–SA was approximately 9.7 nm based on dynamic light scattering (DLS) measurements (see the Supporting Information S2, Figure S2 B and S2 C).

A series of control experiments was initially performed using unstained sections prepared for transmission electron microscopy (TEM) to determine the optimal conditions for silicification of collagen sponges with Ch–SA (see the Supporting Information 3, Figure S3 A–E). By embracing the model of cingulin proteins as water-insoluble templates in the silicification of diatom girdle bands,^[4] successful intrafibrillar silicification of collagen was achieved in 4 days after immersing polyallylamine (PAH)-enriched collagen sponges

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(treatment with 6.67×10^{-4} M PAH for 4 h) in 1.5 % Ch-SA. Unlike hydrated nonsilicified collagen sponges that collapsed after drying, the silicified sponges retained their original shapes after dehydration (see the Supporting Information S3, Figure S3F). This collagen silicification scheme does not involve the use of phosphate ions to induce phase separation of polycyclic aromatic hydrocarbons (PAHs) into macromolecular assemblies.^[23] Despite identification of extrafibrillar silica particles by scanning electron microscopy (SEM), the silicified collagen sponges remained highly porous (see the Supporting Information S3). The well-packed manifestation of the silicified collagen fibrils was attributed to their extensive intrafibrillar silicification. Under TEM, electron-dense minerals were identified within the fibrils that reproduced the cross banding and microfibrillar architecture of fibrillar collagen (Figure 1). The presence of silicon within the collagen fibrils was confirmed using scanning transmission electron microscopy energy-dispersed X-ray analysis (STEM-EDX; Figure 2). Examination of the initial stage of silicification at 24 h identified a continuous electron-dense phase that was formed by fusion of Ch-SA nanoparticles after their infiltration into the PAH-enriched collagen fibrils (see the Supporting Information S4). Micro-computed tomography of the silicified collagen sponges showed that silicification occurred throughout the entire sponge (see the Supporting Information S5).

Silicified PAH-enriched collagen sponges were characterized using different analytical techniques. Fourier transform infrared spectroscopy was used to identify peaks associated with hydrated silica (see the Supporting Information S6). As

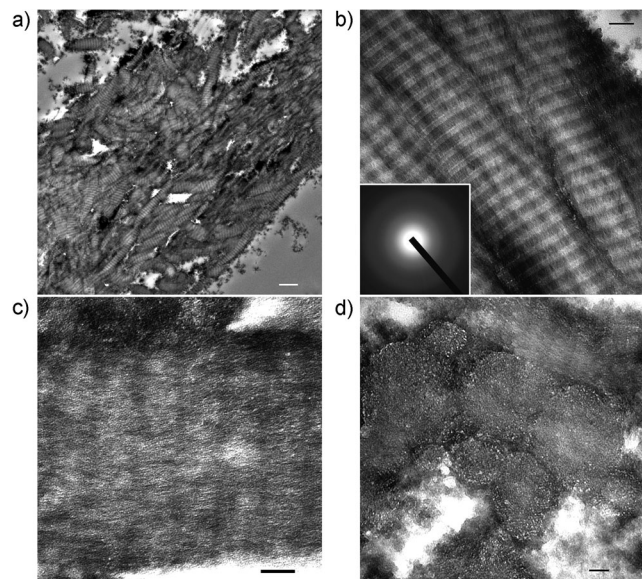


Figure 1. Unstained TEM images of silicified PAH-enriched collagen. a) A leaflet from a silicified collagen sponge showing electron-dense minerals inside collagen fibrils (bar = 1 mm). b) Intrafibrillar electron-dense minerals replicate the cross-banding and microfibrillar architecture of fibrillar collagen (bar = 100 nm). Inset: selected area electron diffraction reveals the amorphous nature of the infiltrated minerals. c) A heavily silicified collagen fibril showing continuous intrafibrillar mineral strands (bar = 50 nm). d) Cross-section showing mineral infiltration from the surface to the center of the fibrils (bar = 50 nm).

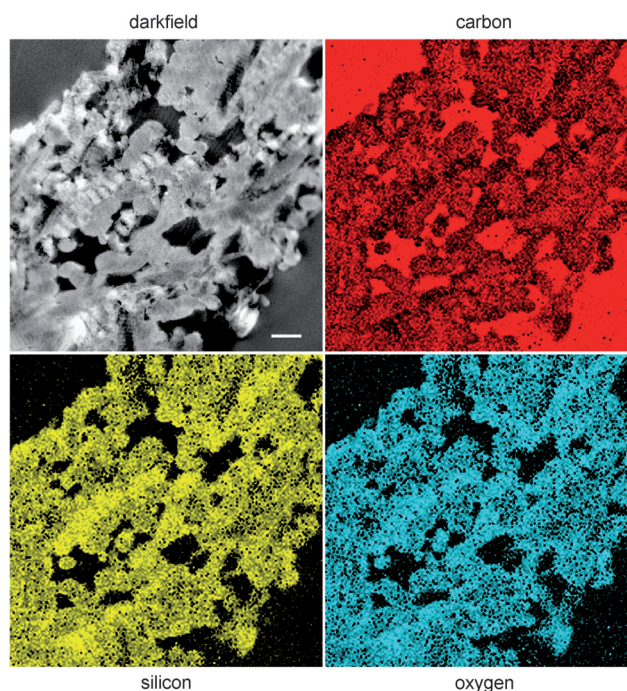


Figure 2. STEM-EDX of an unstained leaflet within a silicified PAH-enriched collagen sponge. Darkfield image shows mineral deposition within the collagen fibrils (bar = 500 nm). The latter contain predominantly silicon and oxygen atoms.

both electron diffraction (Figure 1 b, inset) and powder X-ray diffraction of the silicified sponges showed that the infiltrated mineral phase was amorphous, we sintered the silicified sponges in atmospheric air to 1000 °C and observed different crystalline silica phases (predominantly tridymite and cristobolite) after sintering (see the Supporting Information S7). This result confirms that the mineral phase prior to sintering is amorphous silica. Using ^{29}Si CP-MAS NMR, three different Q species were delineated from the NMR spectrum that provide information on the connectivity of the silica network: Q4 (bulk siloxane), Q3 (single silanol), and Q2 (geminal silanol; Figure 3 a).^[24] The Q4/Q3/Q2 intensity ratio (1:0.76:0.19) of the biosilicified collagen matrix is comparable to the ratio identified from the diatom *Coscinodiscus granii* (1:0.71:0.05),^[23] indicating that they have a similar degree of silica condensation. When ^{13}C CP-MAS NMR was performed on the silicified collagen sponges, we observed an amino acid profile that is characteristic of collagen (Figure 3 b).^[24] High-resolution thermogravimetric analysis showed that the mineral content in the silicified PAH-enriched collagen sponges was 57.2 % (Figure 4 a). Although this mineral content is lower than that of mineralized bone (ca. 60–70 wt %),^[25] one has to take into account the lower molecular weight of silica ($M_r = 60.1$) versus that of hydroxyapatite ($M_r = 502.3$). Changes in derivative of weight loss with temperature further revealed weight loss rate profiles (Figure 4 b) characteristic of collagen^[26] and hydrated silica.^[27]

The mechanical properties of silicified collagen sponges were investigated by examining their compressive stress-strain responses (Figure 5). The tangent modulus of the silicified sponges based on responses from 0–5 % strain is

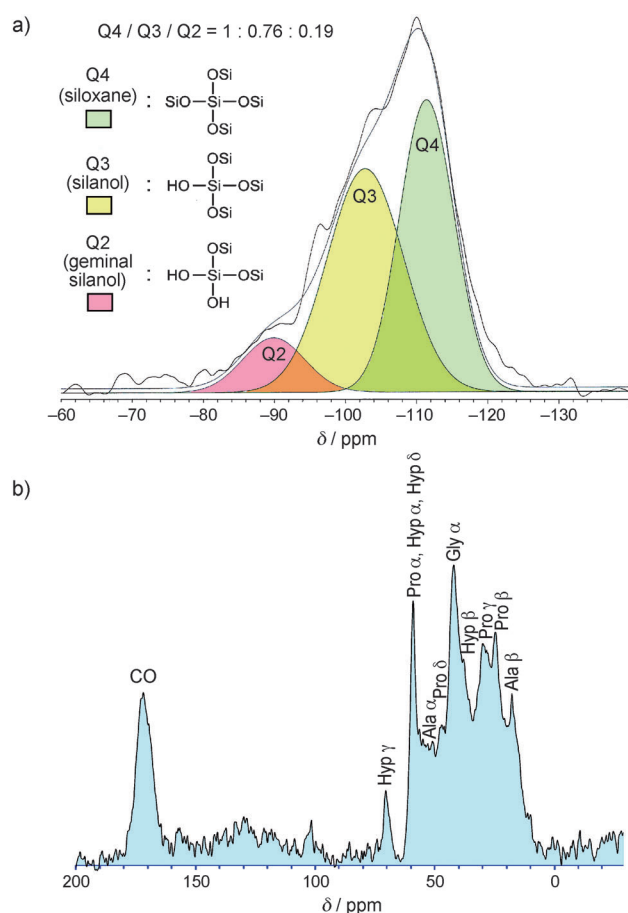


Figure 3. Solid-state NMR spectra of silicified PAH-enriched collagen sponges. a) ^{29}Si CP-MAS spectrum. The broad peak between -78 to -125 ppm was deconvoluted to reveal Q4 (siloxane) at approximately 110 ppm, Q3 (single silanol) at approximately 100 ppm, and Q2 (geminal silanol) at approximately 91 ppm. b) ^{13}C CP-MAS spectrum. Signals assigned to glycine (Gly), alanine (Ala), proline (Pro), and hydroxyproline (Hyp) correspond to those identified from fibrillar collagen.^[24]

599.8 ± 166.0 kPa, which is approximately 48000 times higher than that of nonsilicified sponges (0.013 ± 0.004 kPa). For comparison, the elastic modulus of human trabecular bone is highly dependent on density and ranges from approximately 10 MPa to over 10 GPa.^[28] The modulus of toughness of the silicified collagen sponges (165.3 ± 5.0 kPa) is approximately 1500 times higher than the value obtained for nonsilicified sponges (0.11 ± 0.002 kPa). The modulus of toughness of trabecular bone with a density $= 0.3 \text{ kg cm}^{-3}$ is 1.2 MPa.^[29] Although both the stiffness and toughness of the silicified collagen matrices are lower than human bone, one has to consider that fiber leaflets of a collagen sponge are much thinner ($< 5 \text{ }\mu\text{m}$; Figure 2 and the Supporting Information S3) than natural bone trabeculae (mean trabecular thickness 140 – $172 \text{ }\mu\text{m}$).^[30,31] Collagen fibrils within fiber leaflets of collagen sponges are also somewhat isolated and are not able to reinforce one another, as in bone. Moreover, collagen sponges lack the different levels of hierarchy that are seen in bone. As silicified collagen sponges demonstrated improved stress-strain relationships over the highly compliant nonsilicified sponge, they are potentially useful as porous scaffolds for

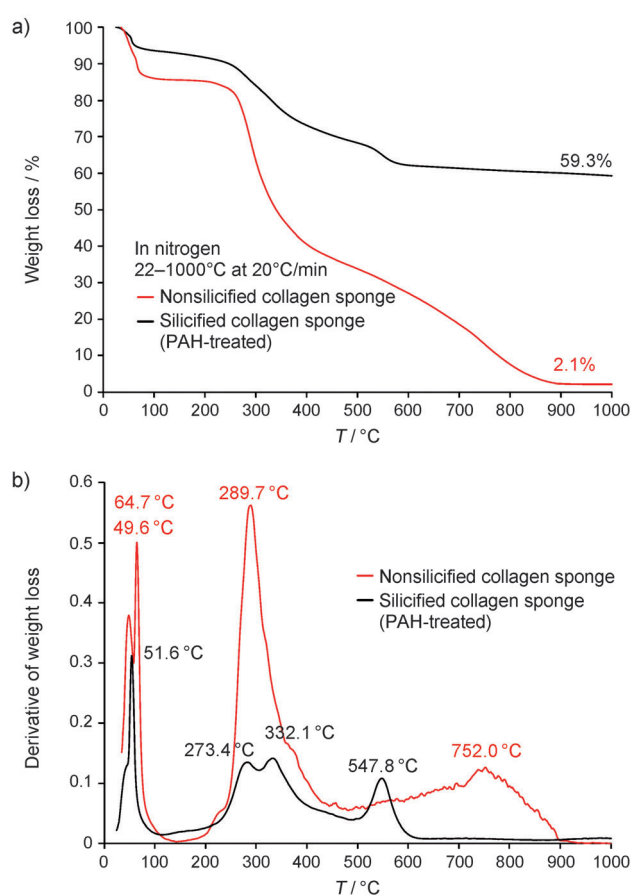


Figure 4. Thermogravimetric analysis of nonsilicified vs. silicified PAH-enriched collagen sponges. a) Plot of weight loss vs. temperature showing a $57.2 \text{ wt}\%$ increase in mineral content after silicification. b) Plot of the derivative of weight loss vs. temperature. For the nonsilicified sponge, the two peaks at 49.6°C and 64.7°C are caused by dissociation of water from the primary and secondary hydration compartments of type I collagen. The two additional peaks observed at high temperatures are attributed to decomposition (289.7°C) and combustion of the organic matrix (752°C).^[26] For the silicified sponge, three desorption processes can be observed: physisorbed water (51.6°C), hydrogen-bonded structural water (273.4°C), and water released by condensation of silanol to siloxane (547.8°C).^[27]

bone repair in regions with minimal to moderate load-bearing requirements.

To verify that intrafibrillar silicification of collagen is not a “staining artifact” (sections were examined unstained), we sintered silicified collagen sponges to completely remove the organic components (see the Supporting Information S8). The appearance of nonporous fiber cores indicates that collagen fibrils were heavily silicified with amorphous silica prior to sintering. To verify that silicification of reconstituted collagen sponges can be reproduced in natural collagen, we silicified PAH-enriched rat tail tendon collagen in 1.5% Ch-SA and obtained the same results (see the Supporting Information S9). Electron tomography and 3D visualization of silicified rat tail tendon collagen are included in the Supporting Information S9.

Our findings provide a new concept in biosilica materials synthesis that does not require phosphate supplements. Figure 6 is a schematic depicting the hypothetical events

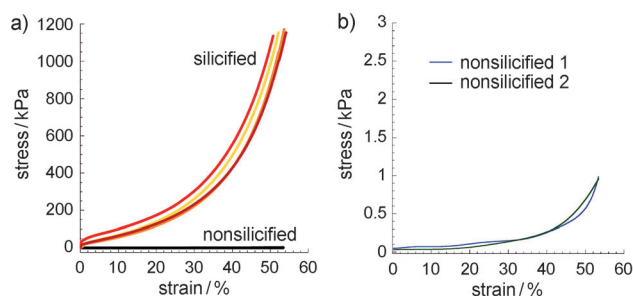


Figure 5. Compressive stress–strain responses of silicified PAH-enriched vs. nonsilicified, hydrated collagen sponges. a) Stress–strain responses were recorded only to the maximum capacity of the load cell and did not represent failure of the silicified sponges. b) For the nonsilicified sponges, the stress axis is scaled down for better visualization.

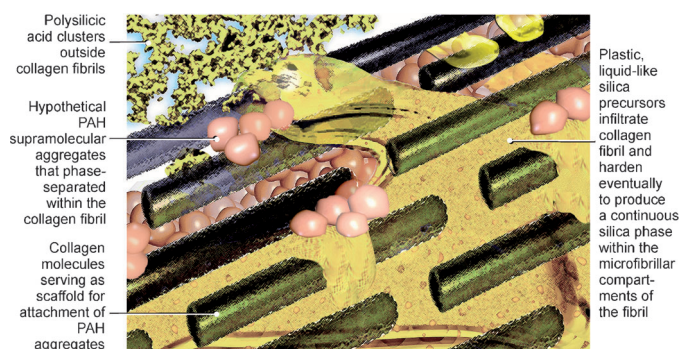


Figure 6. A schematic representation of a biosilicification approach based on the use of a polyamine-enriched collagen template.

that occur in this biosilicification scheme. It is possible that PAH treatment of collagen fibrils results in phase separation of the PAH into macromolecular assemblies within the microfibrillar milieu of the fibrils. As Ch–SA nanoparticles infiltrate the collagen fibril, they fuse to produce a liquidlike silica precursor phase resembling the polymer-induced liquid precursors in calcium carbonate and calcium phosphate mineralization of collagen,^[32,33] that eventually fills the gap zones and microfibrillar spaces. An alternative mechanism is that PAH covers the collagen sponge surface, with its positive charge attracting the negatively charged silica species to facilitate silicification of the PAH-coated sponges. Either mechanism results in the PAH-enriched collagen serving as a template for polymerization of the silica precursor phase. Complete infiltration of the water-filled spaces by the liquid-like silica precursor phase results in replication of the cross-banding patterns and microfibrillar architecture of the collagen fibril.

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