

Research Article

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1 Biomimetic Mineralization of Recombinamer-Based Hydrogels 2 toward Controlled Morphologies and High Mineral Density

- ³ Yuping Li,*^{,†} Xi Chen,[†] Alex Fok,[†] Jose Carlos Rodriguez-Cabello,[‡] and Conrado Aparicio*,[†]
- 4 †Minnesota Dental Research Center for Biomaterials and Biomechanics, University of Minnesota, Minnesota, Minnesota 55455,
- 5 United States

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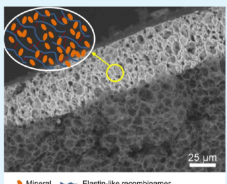
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- 6 [‡]GIR Bioforge, University of Valladolid, Valladolid 47002, Spain
- 7 Supporting Information

ABSTRACT: The use of insoluble organic matrices as a structural template for the bottom-up fabrication of organic—inorganic nanocomposites is a powerful way to build a variety of advanced materials with defined and controlled morphologies and superior mechanical properties. Calcium phosphate mineralization in polymeric hydrogels is receiving significant attention in terms of obtaining biomimetic hierarchical structures with unique mechanical properties and understanding the mechanisms of the biomineralization process. However, integration of organic matrices with hydroxyapatite nanocrystals, different in morphology and composition, has not been well-achieved yet at nanoscale. In this study, we synthesized thermoresponsive hydrogels, composed of elastin-like recombinamers (ELRs), to template mineralization of hydroxyapatite nanocrystals using a biomimetic polymer-induced liquid-precursor (PILP) mineralization process. Different from conventional mineralization where minerals were deposited on the surface of organic matrices, they were infiltrated into the



Mineral Elastin-like recombinamen

frameworks of ELR matrices, preserving their microporous structure. After 14 days of mineralization, an average of 78 μ m mineralization depth was achieved. Mineral density up to 1.9 g/cm³ was found after 28 days of mineralization, which is comparable to natural bone and dentin. In the dry state, the elastic modulus and hardness of the mineralized hydrogels were 20.3 \pm 1.7 and 0.93 \pm 0.07 GPa, respectively. After hydration, they were reduced to 4.50 \pm 0.55 and 0.10 \pm 0.03 GPa, respectively. These values were lower but still on the same order of magnitude as those of natural hard tissues. The results indicated that inorganic—organic hybrid biomaterials with controlled morphologies can be achieved using organic templates of ELRs. Notably, the chemical and physical properties of ELRs can be tuned, which might help elucidate the mechanisms by which living organisms regulate the mineralization process.

30 KEYWORDS: mineralization, elastin-like recombinamers, hydrogel, bone, dentine

1. INTRODUCTION

31 Hard tissues are organic-inorganic nanocomposites that 32 exhibit remarkable mechanical properties with hierarchical 33 structures. 1,2 In bone, a small amount of acidic noncollagenous 34 proteins (NCPs) and the self-assembled collagen fibrils play 35 critical roles in mineralization.^{3,4} Many of the NCPs are highly 36 negatively charged, abundant with carboxylate groups from 37 aspartic and glutamic acid residues or phosphate groups from 38 phosphoserine. They stabilize the initial amorphous calcium 39 phosphate (ACP) precursor phase, facilitate the precursor ion 40 infiltration into collagen fibrils, and mediate mineral phase 41 transformation. 5-8 Several studies have indicated that collagen 42 fibrils provide a structural template where the interstitial spaces 43 in the fibrils serve as confined compartments for mineral 44 deposition, $^{4,9-12}$ i.e., the ACP nanoclusters infiltrate into the 45 collagen fibrils and crystallize into oriented hydroxyapatite 46 (HA) nanocrystals with their [001] direction parallel to the 47 long axes of the collagen fibrils.³

Although extensive research has been performed on collagen-49 based mineralization, it should be noted that collagen matrices can only be obtained from natural sources.^{4,48-50} There are 50 concerns with disease transmission and immunogenic response. 51 The organization of collagen molecules is also very sensitive to 52 the processing conditions.⁵¹ Only collagen fibrils with banding 53 patterns can be mineralized in a way that the minerals infiltrate 54 within them.⁵² Other natural macromolecules and synthetic 55 polymers, such as chitosan, polycaprolactone, polylactide, and 56 Pluronic, have been used as organic matrices for apatite 57 mineralization. 40,53-55 However, mineralization of these 3D 58 bulk scaffolds using traditional crystallization processes, such as 59 incubation in simulated body fluid (SBF), often resulted in the 60 heterogeneous nucleation of HA on the surface of substrates or 61 loosely trapped within porous scaffolds, 13,16,17 compromising 62 their structural integrity as distinct organic and inorganic phases 63 are formed. Such a scenario is also different from that of mature 64 bone, which is composed of two continuous organic and 65

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66 inorganic phases that determine its superior mechanical 67 properties. ¹⁸

In biologically controlled mineralization, the living organism 69 controls the mineralization process directly or indirectly to 70 produce biominerals with selected size, morphology, and 71 structure. 19,20 The elaborate biomineral morphologies may 72 result from the complexity of the template; i.e., the transient 73 amorphous clusters accrete into the insoluble organic template 74 and "mold" into the 3D structure of the matrix. 21-23 It is 75 believed that the development of organic-inorganic nano-76 composites on a large scale with complex 3D morphologies can 77 only be achieved via a templating process, where the shape of 78 the template defines the form of the final material.²⁰ The fluidic 79 character of the amorphous precursor phases has many 80 advantages for generating complex morphologies. 23 It has 81 been shown that amorphous calcium carbonates can adapt to 82 the shape of a polymeric matrix that mimicked the structure of 83 the organic matrix in living organisms. 22,24,25 Calcite crystals 84 can be molded into complex 2D and 3D structures thru the 85 polymer-induced liquid-precursor (PILP) process, which has 86 proven to be the underpinning mechanism of shape control in 87 biomineralization.²² Combination of the process with various 88 substrates, templates, or compartments may build a variety of 89 biomimetic structures useful for hard tissue engineering.

Elastin is an extracellular matrix protein that is known for providing elasticity to tissues and organs, such as blood vessels, elastic ligaments, lung, and skin. Elastin-like recombinamers (ELRs) are biosynthetic recombinant polypeptides based on a repeating pentapeptide sequence derived from tropoelastin, (VPG-Xaa-G), where Xaa is a guest amino acid (excluding proline). One of the unique properties of these recombinamers is their inverse transition temperature (T_t) that allows them to transit between a soluble form at temperatures below T_t and an insoluble aggregation at temperatures above T_t . Although mineralization of ELRs has been investigated by directly mixing them with minerals or incubating them in conventional mineralization solutions, the role of ELRs in calcium phosphate mineralization has not been well explored.

Here, we demonstrated that ELR-based hydrogels can be mineralized through a biomimetic mineralization process by which the minerals infiltrate and deposit within their matrix framework. Thus, the original microstructure of the hydrogel controlled their final morphologies after mineralization and resulted in high mineral content. Because the ability to engineer peptide sequences derived from elastin allows the precise control of the physicochemical and structural characteristics of the recombinamers, i.e., mechanical stability, elasticity, in inherent bioactivity, and self-assembly properties, in particular their their biomineralization and use in hard-tissue regeneration.

2. MATERIALS AND METHODS

2.1. Materials. ELRs HSS₃ and REDV were synthesized according to the published procedures, and their physicochemical properties are listed in Table 1. Sequences of the two ELRs are as follows: HSS₃: [((VPGIG)₂VPGKG(VPGIG)₂)DDDEEKFLRRIGRFG-120 ((VPGIG)₂VPGKG(VPGIG)₂)]₃(VPAVG)₂₀[((VPGIG)₂VPGKG-121 (VPGIG)₂)DDDEEKFLRRIGRFG((VPGIG)₂VPGKG(VPGIG)₂)]₃; 22 REDV: [(VPGIG)₂(VPGKG)(VPGIG)₂EEIQIGHIPREDVDYHLYP-123 (VPGIG)₂(VPGKG)(VPGIG)₂(VGVAPG)₃]₁₀.

Sodium phosphate dibasic and N-hydroxysuccinimide (NHS) were purchased from Fischer Scientific (Pittsburgh, PA, USA). Poly(L-126 aspartic acid) (polyAsp) sodium salts were purchased from Alamanda Polymers (Huntsville, AL, USA). 1-Ethyl-3-[3-(dimethylamino)-

Table 1. Physicochemical Properties of the Elastin-Like Recombinamers a

ELR	$\begin{array}{c} \text{molecular} \\ \text{weight } M_{\text{w}} \\ \text{(kDa)} \end{array}$	isoelectric point	net charge at pH 7.0	ratio of hydrophilic residues/total number of residues (%)
HSS_3	44.9	10.3	8.0	13
REDV	80.1	5.3	-18.1	10

"Peptide calculator: http://www.bachem.com (accessed on October 19, 2015).

propyl] carbodiimide hydrochloride (EDC) and all other chemicals 128 were purchased from Sigma-Aldrich (St. Louis, MO, USA). 129

2.2. Preparation of Cross-Linked ELR Hydrogels and 130 Mineralization. ELR molecules were cross-linked with EDC and 131 NHS. Briefly, 7 mg of ELRs were dissolved in 1 mL of 2-(N- 132 morpholino) ethanesulfonic acid (MES, 50 mM, pH 6.8) mixed with 133 50 mM EDC and 25 mM NHS and stored at 4 °C overnight for cross- 134 linking. They were then centrifuged at 37 °C, and the precipitate was 135 immersed in a solution containing 0.1 M Na₂HPO₄ and 2 M NaCl for 136 2 h to hydrolyze any remaining activated carboxyl groups of peptides 137 and EDC. After rinsing with distilled water, the cross-linked hydrogels 138 were obtained. Mineralization of the cross-linked ELR hydrogel was 139 conducted via the PILP process.³⁴ Mineralization solution was 140 prepared by mixing equal volumes 9 mM CaCl₂·2H₂O and 4.2 mM 141 K₂HPO₄ in Tris-buffered saline (pH 7.4 at 37 °C). polyAsp sodium 142 salt (M_w : 27 000 Da, 50 μ g/mL) was dissolved in the CaCl₂ solution 143 described above before mixing. Mineralization (7-28 days) was 144 performed because it has been generally used in biomimetic 145 mineralization systems.^{35,36} The PILP solution was refreshed every 3 146 days during the mineralization.

2.3. Characterization of the Mineralized Hydrogels. Differ- 148 ential scanning calorimetry (DSC, TA Instruments Q1000, USA) was 149 used to analyze the thermal response of HSS₃. A 10 μ L aliquot of a 50 150 mg/mL HSS $_3$ aqueous solution was placed in a 20 μ L aluminum pan 151 and sealed. An equal volume of distilled water was placed in a 152 reference pan. They were heated from 0 to 40 °C at a constant rate of 153 5 °C per min to obtain isothermal curves. The morphologies of the 154 hydrogels before and after mineralization were analyzed by FE-SEM 155 (JEOL 6500, Tokyo, Japan) at 5 kV and FE-TEM (FEI Tecnai G2 156 F30, Hillsboro, OR, USA) at 300 kV. For SEM, the samples were 157 lyophilized and sputter-coated with 5 nm of Pt. Energy dispersive X- 158 ray spectroscopy (EDS) analysis was performed during SEM analysis 159 at 15 kV. TEM samples were prepared by embedding the lyophilized 160 sample in epoxy resin, cut with an ultramicrotome (Leica Reichert 161 UltraCut S) at room temperature, and collected on copper grids. 162 Selected-area electron diffraction (SAED) was performed during TEM 163 analysis to identify the crystallinity of the minerals. The crystal 164 structure of the minerals was also characterized using a micro- 165 diffractometric system with a 2D area detector (Bruker AXS, 166 Germany) at 45 kV and 40 mA. The incident angle was 15°, and 167 the detector position was fixed at 30°, which covered the angular range 168 from 15 to 45° in 2θ . Data collection time was 1000s, and the results 169 were analyzed using the JADE8 software (Materials Data Inc., JADE, 170 Livermore, CA, USA). Microcomputed tomography (μ -CT, HMX-XT 171 225, X-tek system, United Kingdom) was used to determine 3D 172 structure and mineral density of the hydrogel after mineralization using 173 the following operational parameters: 90 kV, 90 µA, 720 projections, 174 and 4 frames per projection. The volumetric reconstruction of the 175 microradiographs was performed using CT Pro 3D (Nikon Metrology, 176 Brighton, MI, USA); then, the reconstructed volume was analyzed 177 using VG Studio Max (Version 2.1.3, 64 bit, Volume Graphics, 178 Charlotte, NC, USA). An HSS₃ hydrogel after 28 days of 179 mineralization (approximately $0.4 \times 1.5 \times 1.5 \text{ cm}^3$) as well as a 180 bovine dentin and cortical bone block were used for this analysis. 181 Mineral density was quantified by comparing the attenuation 182 coefficient of the mineralized hydrogel with CT-based calibration 183 phantoms of HA.

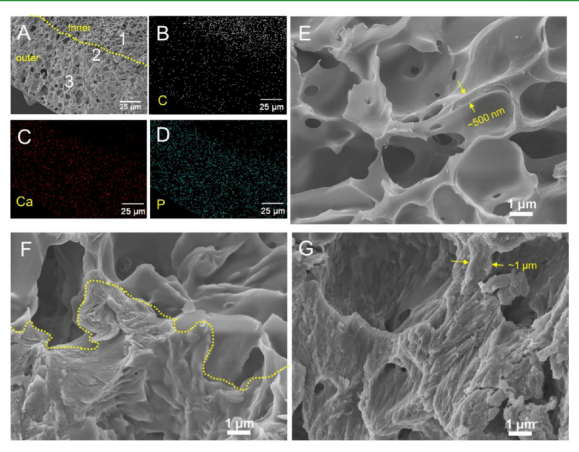


Figure 1. (A–D) Cross-sectional SEM image and elemental mapping of the cross-linked HSS₃ hydrogel after 14 days of mineralization in the PILP solution. (E–G) Cross-sectional SEM images from the regions of 1, 2, and 3 marked in A, respectively. The dashed lines indicate the interface between the outer mineralized and the inner nonmineralized regions.

185 2.4. Mechanical Testing. Mechanical properties of the ELR 186 hydrogel after 28 days of mineralization were examined by nanoindentation using a MTS nanoindenter XP, equipped with a Berkovich tip at room temperature. Testworks 4 software, which 189 incorporates the Oliver-Pharr data analysis method³⁷ was used to 190 analyze their elastic modulus and hardness. Ultramicrotome sectioning was used to reveal the indentation surfaces of the mineralized region of 192 the hydrogel that was embedded in an epoxy resin. Nanoindentation was also performed on a cortical bone from bovine tibia. Preparation of bone sample was performed as we previously described.3bovine cortical bone was embeded into PMMA, sectioned with a low-196 speed diamond saw (Isomet, Buehler, Lake Bluff, IL, USA), ground with sand papers (SiC, 600, 1000, and 1200), and polished with 198 alumina suspensions (5, 1, and 0.1 μ m) to obain a smooth surface suitable for nanoindentation. A total of 10 indents were performed on 199 each sample. Mean elastic modulus and hardness were obtained from 200 representative properties-displacement curves at a depth of 2000 μ m. 2.5. Statistical Analysis. Analysis of the statistically significant 203 differences on mechanical properties among groups was performed 204 with one-way ANOVA tables (SPSS v.19, IBM). The level of statistical 205 significance was set at p < 0.05.

3. RESULTS AND DISCUSSION

3.1. Thermoresponsive Behavior of HSS₃. HSS₃ used in this study is an ABA amphiphilic triblock ELR. Block A is compose of [[(VPGIG)₂]₂(VPGKG)] (VPGIG)₂]₂DDDEEKFLRRIGRFG[(VPGIG)₂(VPGKG)] (VPGIG)₂]₃. Block B is (VPAVG)₂₀. This block polymer displays the aforementioned reverse thermal response. DSC analysis revealed its endotherm ranged from 22 to 37 °C, which was resolved in two peaks corresponding to the individual

transitions of the hydrophobic moieties of the two different $_{214}$ blocks (Figure S1). Although the hydrophobic moieties of both $_{215}$ blocks have similar polarity, the influence of the nearby polar $_{216}$ amino acids might shift the $T_{\rm t}$ of the block A to a value higher $_{217}$ than that of the block B. $_{38}$

3.2. Fabrication and Mineralization of Cross-Linked 219 HSS₃ Hydrogels. When the HSS₃ molecules were cross-linked 220 with carbodiimides, they formed microporous hydrogels 221 (Figure S2). These hydrogels were transparent at 4 °C. They 222 became opaque rapidly when incubated at 37 $^{\circ}\text{C}$, indicating the $_{223}$ occurrence of phase transition. Mineralization via the PILP 224 process using 50 μ g/mL polyaspartate (M_w : 27 000 Da) as 225 process-directing agent was conducted, and the PILP solution 226 was replaced every 3 days to keep it clear. After 14 days of 227 mineralization, mineralization depth of 78 \pm 4 μ m was achieved 228 on the outer region of the porous HSS₃ hydrogel showing 229 strong signals of calcium and phosphate, whereas a large 230 amount of carbon along with sparse signals of calcium and 231 phosphate were found in its inner side (Figure 1). The 232 fl thickness of the hydrogel framework at the nonmineralized 233 region was around 500 nm (Figure 1E), and it was 234 approximately 1 μ m at the mineralized region (Figure 1G). 235 In contrast to the conventional mineralization of hydrogels, 236 where HA was precipitated on the surface of the organic 237 matrices, 16,39 the minerals were specifically deposited within the 238 framework of the HSS3 hydrogel, preserving its microporous 239 structure. Besides, the texture of the mineralized region showed 240 striking similarity to that of the cortical bone at the nanoscale 241 (Figure S3).

Figure 2. TEM images of the cross-linked HSS₃ hydrogel after 14 days of mineralization. (A) Representative TEM image and (B) corresponding SAED pattern of the hydrogel at the mineralized region. The minerals were homogeneously distributed in the framework of the hydrogel, and they were randomly oriented HA nanocrystals. (C) Hydroxyapatite nanocrystals were needlelike. Platelike fragments (pointed to by arrows) without lattice fringes were presumably from the organic hydrogel. (D) HR-TEM lattice image of HA nanocrystals showing 0.8 and 0.34 nm planar spacing, which correspond to the HA (100) and (002) planes, respectively.

TEM images revealed that nanocrystals were homogeneously deposited in the hydrogel framework at the mineralized region (Figure 2A). SAED produced a multiple-ring-shaped diffraction pattern for HA, including a ring for the (002) and (210) planes and one for the combined (211), (112), and (300) planes (Figure 2B), indicating that the crystals were randomly oriented in the hydrogel framework. Such crystal orientation is different from the one found in mineralized collagen fibrils where the 251 nanocrystals were oriented parallel to the longitudinal axes of 252 the collagen fibrils. Because the polymer chains in the hydrogel 253 framework were randomly distributed, it may not contain oriented confinements to guide the crystal orientation that 255 found in the collagen system. 40,41 The crystals were needlelike, 256 approximately 10 nm wide and 50-150 nm long (Figure 2C). 257 They had 0.8 and 0.34 nm planar spacings as assessed with the 258 HR-TEM lattice image (Figure 2D), which correspond to the 259 (100) and (002) planes of HA, respectively. X-ray diffraction 260 spectra further confirmed that the minerals were HA

nanocrystals (Figure 3). Peaks for the (002), (210), (211), 261 f3 (202), and (310) planes of HA were broad and matched well 262 with the diffraction pattern of bovine cortical bone.

3.3. The Role of Elastin-Like Recombinamers on 264 Mineralization. Unlike in classical crystal nucleation, where 265 nucleation from a supersaturated solution occurs when a 266 sufficiently large density fluctuation of the solution overcomes a 267 free energy barrier between the crystalline and solution phases, 268 calcium phosphate biomineralization proceeds via an amor- 269 phous precursor phase. ^{8,42} The PILP process is a biomimetic 270 mineralization system in which anionic polyaspartic acids mimic 271 the role of acidic proteins in biominerals, forming a liquidlike 272 ACP precursor complex. ⁴ Because of its fluidlike characteristics, 273 the PILP process does not require specific interactions with 274 specific crystallographic planes in contrast to the classical 275 crystallization. ²² These liquid precursors are well-suited to 276 achieve the controlled morphogenesis using templates because 277 they can easily adapt to any shape before crystallization. ²⁰ They 278

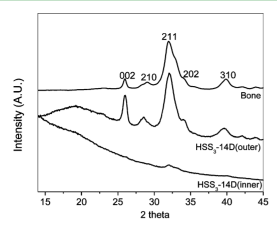


Figure 3. XRD spectra of the cross-linked HSS₃ hydrogel after 14 days of mineralization in the inner nonmineralized and outer mineralized regions as well as bovine cortical bone.

279 can even infiltrate into small cavities of the insoluble organic 280 matrix as found in the bone and dentin; i.e., the ACP 281 nanoclusters deposit into the gap zones and nanochannels of 282 the type-I collagen fibrils.^{3,4,12}

It has been demonstrated that elastin-like poly(VPGVG) and 1284 its analogs poly[f_v (VPGVG), f_x (VPGKG)] (0.1 $\leq f_x \leq$ 0.2, f_v + 1285 f_x = 1) undergo hierarchical self-assembly. At the nanoscale, 1286 those ELRs self-assembled into 5 nm wide twisted filaments 1287 consisting of several β spirals. These filaments were aligned in

parallel into fibrils several hundred nanometers in diameter. 288 The HSS₃ molecules in aqueous solution can also self-assemble 289 into anisotropic fibrils around 150 nm in diameter at 37 °C 290 (Figure S4). Although the HSS3 chains in the hydrogel were 291 cross-linked, the ordered structures, such as β spiral and twisted 292 filament may still form at a temperature above T_t because the 293 hydrogel exhibited opalescence, an indication of microphase 294 separation (microdomains in tens of nanometers) between 295 hydrophobic and hydrophilic moieties. 31,44 Therefore, nano- 296 pores and/or nanochannels from the hydrophilic moieties may 297 form within the hydrogel framework, serving as compartments 298 for mineral deposition. It is possible that the amorphous 299 precursor nanoclusters infiltrate into the nanocompartments of 300 the ELR hydrogel and then coalesce, solidify, and crystallize to 301 form close-packed crystals. Further investigations on the 302 mechanisms of mineralization in the ELR hydrogels are 303 underway.

The peptide sequence of SN_A15 (DDDEEKFLRRIGRFG), 305 derived from statherin, may promote the uptake of the liquid 306 ACP precursor into the hydrogel. Statherin is an acidic 307 phosphopeptide with a high degree of structural and charge 308 asymmetry. SN_A15 has a high binding affinity and crystal 309 growth inhibition to calcium phosphate minerals. Previously, 310 we have demonstrated that statherin-derived ELRs have the 311 ability to induce and control the growth of minerals on 312 biofunctionalized surfaces. However, the sequence of SN_A15 313 may not be critical for mineral infiltration. As shown in Figure 314 64

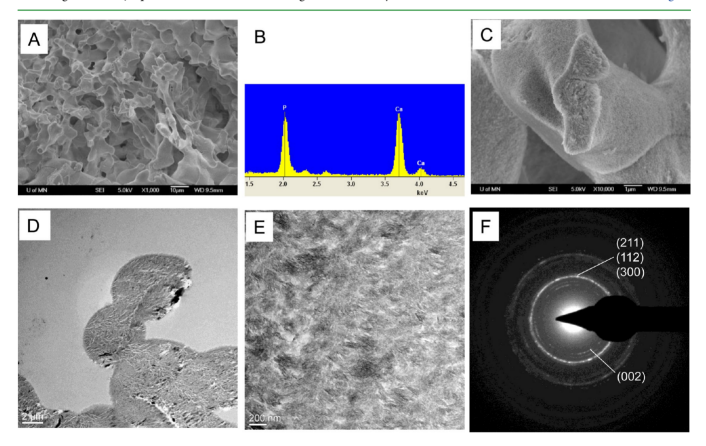


Figure 4. SEM and TEM analysis of the REDV hydrogel after 14 days of mineralization by the PILP process. (A) SEM image and (B) its corresponding EDS of the mineralized REDV, showing that calcium phosphate minerals were deposited into the REDV hydrogel framework. (C) Fractured surface of the REDV hydrogel after mineralization. (D) TEM image of the mineralized REDV hydrogel showing densely packed and homogeneously distributed minerals in the polymeric matrix. (E and F) TEM image and the corresponding SAED pattern, respectively, revealed that the randomly oritented minerals were needlelike HA nanocrystals.

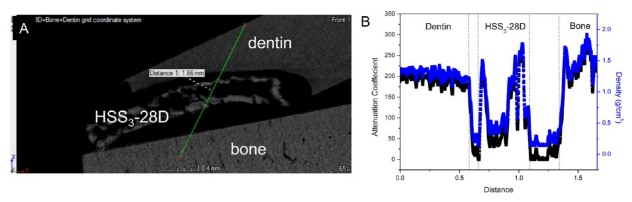


Figure 5. (A) Micro-CT image of dentin, bone, and the HSS₃ hydrogel after 28 days of mineralization (HSS₃-28D). (B) Distribution of the attenuation coefficient and mineral density of the region marked with a green line in A.

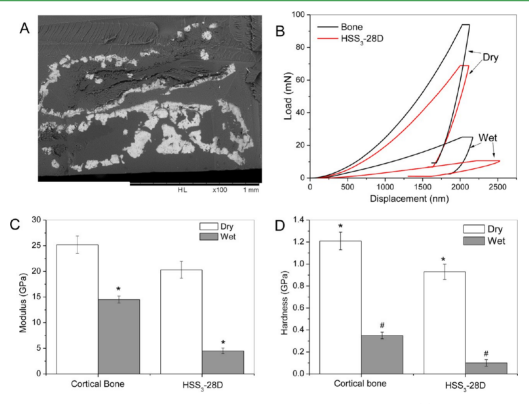


Figure 6. (A) SEM image of the microtome section of HSS₃-28D used for nanoindentation. Representative (B) nanoindentation load—displacement curves, (C) elastic modulus, and (D) hardness of the bovine cortical bone and the HSS₃-28D in the dry and wet states.

315 4, the REDV hydrogel contains no SNA15 and was also 316 successfully mineralized. After 14 days of mineralization, the 317 framework of the cross-linked REDV hydrogel was still discernible (Figure 4A), and the corresponding EDS analysis confirmed the presence of calcium and phosphate (Figure 4B). The fractured surface of the mineralized REDV showed a 320 homogeneously granular morphology, indicating that the 321 322 minerals have deposited within the framework and were wellintegrated with the organic REDV hydrogel (Figure 4C). TEM (Figure 4D,E) and SAED (Figure 4F) images demonstrated that needlelike HA nanocrystals were randomly distributed in 326 the REDV hydrogel, similar to those found in the mineralized 327 HSS₃ hydrogel (Figure 2). Previous studies suggested that electrostatic interactions between amorphous ACP clusters and 329 collagen-mediated infiltration of the mineral into the collagen 330 fibrillar matrix, i.e., polyaspartic acid molecules formed 331 negatively charged complexes with ACP that interacted with

a positively charged region of collagen located at the C-terminal 332 region of the gap zone. Our results demonstrated that ACP 333 nanoclusters can infiltrate into negatively charged REDV 334 matrices (Table 1), suggesting that the electrostatic interactions 335 may not be the major driving force for the mineralization in 336 these organic matrix frameworks.

3.3. Mineral Density and Nanoindentation. Micro-CT ₃₃₈ analysis of the HSS₃ hydrogel after 28 days of mineralization ₃₃₉ demonstrated that a mineral density of up to 1.90 g·cm⁻³ was ₃₄₀ achieved, which is higher than that of dentin but lower than ₃₄₁ that of bovine cortical bone (Figure 5A,B).

Mechanical properties of the hydrogel at the mineralized 343 region were measured by nanoindentation and compared to 344 those of bovine cortical bone. As shown in Figure 6A, the 345 66 thickness of the mineralized region ranged from 40 – 200 μ m. In 346 the dry state, the load–displacement curve (Figure 6B) was 347 remarkably similar to the one obtained from bovine cortical 348

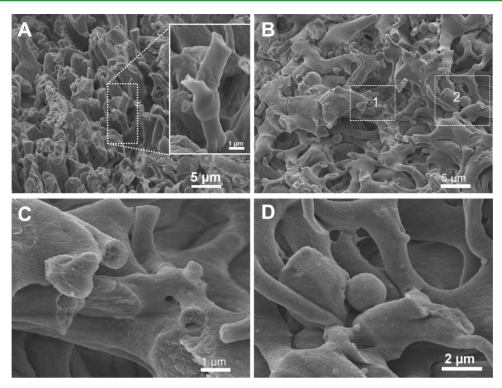


Figure 7. SEM images of the cross-linked HSS_3 hydrogel after 7 days of mineralization. (A) Fractured framework of the hydrogel. Insert: a hydrogel strut that displayed mineralized top and unmineralized bottom. (B) Complex interwoven network of struts and microspheres with granular surfaces. (C and D) Higher-magnification SEM images of the fractured surface of the hydrogel framework from areas 1 and 2 marked in B, which are indicative of the full integration between the minerals and the HSS_3 matrices.

349 bone. Its elastic modulus was 20.3 ± 1.7 GPa, comparable to 350 natural cortical bone, whereas its hardness (0.93 \pm 0.07 GPa) 351 was significantly lower than that of bone (Figure 6C,D). When the specimen was hydrated, the mechanical properties of both the mineralized hydrogel and cortical bone were significantly 354 reduced. The elastic modulus and hardness of mineralized 355 hydrogel were decreased to 4.50 \pm 0.55 and 0.10 \pm 0.03 GPa, respectively. In the wet state, the mechanical properties of the mineralized region of the hydrogel were significantly lower, around one-third of those measured for bovine cortical bone. It has been reported that the elastic modulus of demineralized dentin lesions can be restored from 0.2 GPa to near 10 GPa in wet state after 14 days of mineralization using the PILP process.⁵⁶ Our previous study also demonstrated that the 363 hardness and elastic modulus of the nanoporous intrafibrillarly 364 mineralized collagen films were 0.7 and 9.1 GPa, respectively, in 365 the dry state, whereas in the wet state they were 9 and 177 366 MPa, respectively.³⁴ In contrast, the mechanical properties of the mineralized ELRs studied here were over 1 order of magnitude higher than those of the nanoporous intrafibrillarly mineralized collagen films and on the same order of magnitude 370 as those of the restored dentin lesions.

3.4. ELR-Template-Directed Control of Hybrid Mor372 phologies. The use of insoluble organic matrices as a
373 morphological template for the bottom-up fabrication of
374 organic—inorganic nanocomposites is a powerful way to build
375 a variety of advanced hybrid biomaterials. In contrast to metals
376 and ceramics, polymers are much more easily fabricated into
377 diverse shapes. By controlling mineral deposition in the organic
378 matrices, predictable morphology of the nanocomposites can
379 be obtained. In our study, many struts were observed on a HSS₃
380 hydrogel after mineralization (Figure 7A). In some cases, they

showed a granular/rough surface (top part in inset of Figure 381 7A) or a smooth surface (bottom part, inset of Figure 7A), 382 revealing the occurrence of ongoing mineralization. Interwoven 383 struts and microspheres with fully granular surfaces were also 384 found (Figure 7B–D). These complex morphologies and 385 shapes with curved surfaces suggested that the original 386 microstructure of the hydrogel dictated the final morphology 387 of the nanocomposites.

CONCLUSIONS

We have demonstrated that hydrogels of the elastin-like 390 recombinamers templated calcium phosphate mineralization 391 where the minerals selectively deposited within the hydrogel 392 frameworks. By using a bottom-up method, a new class of 393 hybrid nanocomposites with controlled morphologies was 394 developed. These composites possessed microporous structures 395 and mineral density comparable to that of natural bone. Their 396 mechanical properties were on the same order of magnitude as 397 those measured from bovine cortical bone. The use of the ELR 398 hydrogels opens the possibility to study in vitro model systems 399 that reproduce biomimetic processes. By designing the 400 sequence of the ELRs and controlling the morphologies of 401 ELRs matrices at different dimensional levels, diverse hybrid 402 nanocomposites with optimized mechanical and biological 403 properties can be constructed and can be suited for the 404 treatment of bone defects using regenerative medicine 405 approaches.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the 409 ACS Publications website at DOI: 10.1021/acsami.5b07628. 410

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DSC analysis on phase transition of HSS₃ in an aqueous solution, SEM visualization of the porous HSS₃ hydrogels, SEM comparative images of the mineralized scaffolds and cortical bone, and the self-assembled fibrillar structures of HSS₃. (PDF)

416 AUTHOR INFORMATION

417 Corresponding Authors

- 418 *E-mail: apari003@umn.edu.
- 419 *E-mail: lixx1191@umn.edu. Tel.: +1-612-625-4467. Fax: +1-
- 420 612-626-1484.

421 Notes

422 The authors declare no competing financial interest.

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