Photo-Cross-Linkable Human Albumin Colloidal Gels Facilitate In Vivo Vascular Integration for Regenerative Medicine

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ABSTRACT: Biodegradable cellular and acellular scaffolds have great potential to regenerate damaged tissues or organs by creating a proper extracellular matrix (ECM) capable of recruiting endogenous cells to support cellular ingrowth. However, since hydrogel-based scaffolds normally degrade through surface erosion, cell migration and ingrowth into scaffolds might be inhibited early in the implantation. This could result in insufficient de novo tissue formation in the injured area. To address these challenges, continuous and microsized strand-like networks could be incorporated into scaffolds to guide and recruit endogenous cells in rapid manner. Fabrication of such microarchitectures in scaffolds is often a laborious and time-consuming process and could compromise the structural integrity of the scaffold or impact cell viability. Here, we have developed a fast single-step approach to fabricate colloidal hydrogels, which are made up of randomly packed human serum albumin-based photo-cross-linkable microparticles with continuous internal networks of microscale voids. The human serum albumin conjugated with methacrylic groups were assembled to microsized aggregates for achieving unique porous structures inside the colloidal gels. The albumin hydrogels showed tunable mechanical properties such as elastic modulus, porosity, and biodegradability, providing a suitable ECM for various cells such as cardiomyoblasts and endothelial cells. In addition, the encapsulated cells within the hydrogel showed improved cell retention upon implantation, achieving rapid recruitment of hematopoietic cells and, ultimately, enhancing the tissue regeneration capacity of implanted scaffolds.

INTRODUCTION

Opening a new path to the effective delivery of biological factors (e.g., stem cells, growth factors, drugs, small interfering RNA (siRNA), and microRNA) to injured tissue sites, engineered biodegradable scaffolds are capable of creating suitable extracellular matrices (ECMs) for implanted cells, thus facilitating and enhancing tissue regeneration. Implantable and biodegradable scaffolds developed to date have utilized numerous natural and synthetic biomaterials; polyester urethane-based porous synthetic scaffolds loaded with siRNA used for promoting in vivo angiogenesis and fibrin-based scaffolds for enhanced cell survival after transplantation, reduction of infarct expansion, and development of neovascularure in the ischemic myocardium are some examples. A major challenge in successful long-term integration of scaffolds within the host tissue is enabling the scaffold to rapidly interface with the existing host microenvironment and cellular architecture while delivering numerous biological factors (e.g., growth factors and drug molecules) to target cells. While several studies have focused on fabricating biomaterials with prevascularized structures or delivering growth factors that induce the growth of endothelial cells at the implant site, the fabrication process is often not straightforward, thus limiting the usability of the scaffold. Moreover, delivering growth factors to the injured site has been limited by the insufficient half-life and instability of these biomolecules, thus impacting the long-term effectiveness of implanted scaffolds.

Among various biomolecules, albumin, one of the most abundant water-soluble proteins found in blood plasma (40–50 mg/mL, 65–70 kDa) due to its negatively charged surface, possesses an ability to bind to various active biological substances (e.g., vitamins, hormones, fatty acids, ions, and drugs) and can transport these materials to the required tissues by enhancing the in vivo half-life of the bound active substances. In addition, albumin is capable of binding to inflammation-inducing substances and free radicals, taking part...
Albumin also takes part in congenital endothelial stabilization and acts as an essential factor for the growth and function of various cells (e.g., endothelial cells, fibroblasts, and smooth muscle cells). Consequently, fabricating albumin-based biodegradable scaffolds can provide unique advantages to tissue regeneration upon implantation since various growth factors and active substances can be incorporated into the implantable scaffold with improved half-lives and increased concentrations. Furthermore, albumin can be self-assembled into aggregates that can pack into a 3D cubic lattice with abundant cavities, i.e., porous channels, in specific conditions. This unique self-assembly property and the biological nature of albumin can be useful for creating continuous and microsized internal networks inside scaffolds. Accordingly, albumin-based hydrogels have recently emerged as promising scaffolds for bone, cardiac regeneration, and wound healing. The fabrication method typically involves thermal or pH-induced albumin gelation or chemically cross-linking with other conjugates. These methods, however, are time-consuming, provide suboptimal cell attachment, and could cause immunogenic reactions upon implantation. Moreover, recent studies have mostly focused on employing bovine serum albumin (BSA), which does not accurately recapture the human serum albumin (HSA) protein sequence and implications for tissue regeneration in humans.

Here, we introduce a human albumin-based colloidal hydrogel, namely, packed photo-cross-linkable human serum albumin methacryloyl (AlMA) microparticles, to improve cell retention and survivability within thick scaffolds. This unique self-assembly property and the biological nature of albumin can be useful for creating continuous and microsized internal networks inside scaffolds. Accordingly, albumin-based hydrogels have recently emerged as promising scaffolds for bone, cardiac regeneration, and wound healing. The fabrication method typically involves thermal or pH-induced albumin gelation or chemically cross-linking with other conjugates. These methods, however, are time-consuming, provide suboptimal cell attachment, and could cause immunogenic reactions upon implantation. Moreover, recent studies have mostly focused on employing bovine serum albumin (BSA), which does not accurately recapture the human serum albumin (HSA) protein sequence and implications for tissue regeneration in humans.

Here, we introduce a human albumin-based colloidal hydrogel, namely, packed photo-cross-linkable human serum albumin methacryloyl (AlMA) microparticles, to improve cell retention and survivability within thick scaffolds. Specifically, albumin from human serum can be conjugated with methacrylic functional groups to form AlMA, which can then self-assemble into hydrogels upon UV and visible light cross-linking.
exposure. Furthermore, the physical and mechanical properties of AlMA hydrogels could be tuned by the degree of methacrylation (DM) and the concentration of AlMA. Consequently, the formed AlMA hydrogel with continuous and microscale internal networks can serve as a guide for infiltrated cells to achieve rapid recruitment and allow for the ingrowth of circulating cells as a “regenerative pod,” unlike any hydrogel with randomly formed porous structures. In the future, the straightforward and highly tunable fabrication process of AlMA biomaterials could be leveraged with recent additive manufacturing technologies, namely, in situ bioprinting, to rapidly create nanocarrier- and growth factor-conjugated scaffolds with longer half-lives, geometries customized to the defect size, and an enhanced functionality in inducing angiogenesis and tissue regeneration.

RESULTS AND DISCUSSION

Fabrication of the Photo-Cross-Linkable AlMA Aggregates. AlMA was synthesized through the nucleophilic substitution reaction, in which electron-rich groups such as thiol (−SH), hydroxyl (−OH), and amino groups (−NH₂) on the side chain of amino acids donate electron pairs to the carbonyl carbon of MA to create acyl derivatives. Among other moieties, the amine groups in lysine predominate the overall reaction due to their steric accessibility within the albumin structure. Therefore, methacryloyl groups were covalently conjugated to free amine groups in recombinant human serum albumin (Figure 1a). It is important to note that maintaining subneutral-pH and low-temperature conditions (<4 °C) is essential to prevent protein denaturation and degradation. ¹H NMR spectra verified the conjugated methacryloyl groups to the AlMA. Due to the strong water-attracting property of albumin, the baseline of the spectrum was not smooth enough to reliably quantify the molecules through integrating the area under the curve. Methacryloyl modification of albumin was therefore confirmed after manual adjustment of the baseline and phase correction. As shown in Figure 1b, we observed two newly generated peaks at 5.4 and 5.7 ppm (red arrows) in the AlMA spectrum, which were created by the protons in the methacrylate vinyl groups. Simultaneously, a reduction in methylene of the lysine was observed at the 2.9 ppm signal (blue arrow), indicating successful conjugation.

Next, we set out to tune the mechanical and physical properties of the AlMA hydrogel by tuning the concentration of MA from 0.05 to 2.0% (v/v), thus achieving various degrees of methacrylation as shown in Figure 1c. As a result, we were able to achieve a wide range of DM (~35 to 82%) in proportion to the concentration of MA, providing a potential to tune the stiffness of AlMA hydrogels. However, a DM of more than 82.1 ± 3.0% (equivalent to 2.0% v/v MA) was not achievable due to the denaturation of the albumin molecules in the presence of high-concentration MA (data not presented), which could be pertinent to the denaturing effect of the adsorption of methacryloyl groups onto the protein structure via strong hydrophobic interactions. To sum up, the degree of methacrylation of AlMA used for the experiments were either 0.6% MA (~61.5% DM) or 2% MA (~83.1% DM), and two different AlMA polymer concentrations (10 or 20% w/v) within each degree of methacrylation were used (Table 1).

Next, we optimized albumin concentration and the incubation conditions to guide the self-assembly of AlMA aggregates at a neutral pH similar to that of biological environments. Albumin is a natural colloidal particle that maintains the colloidal osmotic pressure of blood. The formation of albumin aggregates was observed at high albumin concentrations (>0.05%) at 25 °C for 1 h (data not shown). Furthermore, the conjugated hydrophobic methacryloyl groups on the albumin molecules may take part in altering the amphiphilic nature of albumin, which could affect the formation of the self-assembled aggregates and their sizes. We then demonstrated the effects of the DM on the self-assembled AlMA aggregates. Specifically, AlMA aggregates formed under different DM conditions ranged from ~12 to 2000 nm in size (Figure 1d). The size of the aggregates formed at high (2.0%) and medium (0.6%) DMs were increased (Figure 1e) in comparison to low-DM (0.05%) and pristine albumin aggregates due to the increased hydrophobicity of AlMA molecules upon increased methacryloyl substitutions. The increased hydrophobicity might have induced strong hydrophobic interactions among AlMA molecules in an aqueous environment, which could have resulted in the creation of more large aggregates with increased diameters. The formation of AlMA aggregates at high DM was visually confirmed by cryo-transmission electron microscopy (cryo-TEM), applying the negative staining with uranyl acetate solution (Figure 1f). At high concentration, we confirmed continuous clustered networks (Figure 1f) and individual aggregates were observed at a low concentration of aggregates (Figure 1g), which is in agreement with the previously reported TEM results.

Development and Characterization of AlMA Hydrogels. We first studied the characteristics of hydrogels formed upon photo-cross-linking AlMA colloidal particles in the presence and absence of cells (Figure 2a). Upon UV exposure, the methacryloyl groups in the self-assembled structures were cross-linked, resulting in a randomly packed colloidal hydrogel with continuous and microporous networks. Prepolymer solutions fabricated at high DMs or high AlMA polymer concentrations exhibited significantly decreased transmittance and increased turbidity at 360–480 nm UV wavelengths (Figure 2b,c). AlMA hydrogels fabricated at high DMs showed higher opacity than medium-DM AlMA hydrogels for both low and high AlMA concentrations (Figure 2d). This may be due to increased size of aggregates at higher degrees of MA (Figure 1d). Following UV exposure optimization, large-sized AlMA hydrogels (>5 mm) could be fabricated for structurally robust and easy to handle implantable units for surgery (Figure 2de). Interestingly, when the AlMA hydrogel was observed under a brightfield microscope, a unique internal pore structure different from GelMA hydrogel was confirmed. The internal

| Table 1. Sample Names, % Volume of Methacrylic Anhydride Used, Degree of Methacrylation, and AlMA Polymer Concentrations Used for the Experiments |
|---|---|---|---|
| Sample name | % volume methacrylic anhydride (MA) | degree of methacrylation (%) | AlMA polymer conc. (%) |
| 0.6/10 | 0.6 | 61.5 | 10 |
| 0.6/20 | 0.6 | 61.5 | 20 |
| 2/10 | 2 | 83.1 | 10 |
| 2/20 | 2 | 83.1 | 20 |
patterning of the hydrogel occurred according to the direction of the UV for cross-linking (Figure 2f,g). This internal patterning phenomenon occurred regardless of the presence or absence of cells (Figure 2h and Figure S1).
To study the mechanical tunability of AlMA hydrogels, we varied the DM (0.6 to 2.0%) and observed an increase in the elastic modulus up to $\sim 5$ kPa at high DMs and $\sim 2$ kPa at low DMs (Figure 2i). The elastic modulus of the 10% w/v AlMA hydrogels was comparable to 4 mg/mL collagen hydrogels (control 1) (Figure S2), which is commonly used for tissue engineered scaffolds (e.g., bone, cartilage, skin, vascular grafts, etc.). Increasing the concentration of AlMA, however, showed little effect on the elastic modulus possibly due to the reduction of transmittance followed by the inhibition of PI free radicals and, eventually, a reduction in cross-linking density of the hydrogels (Figure 2b). A controllable degradation profile in an engineered polymer is a crucial factor for balanced tissue formation and the sustained release of bioactive compounds. Albumin has been proposed as an injectable delivery vehicle for various drugs and can be degraded by collagenase and other digestive proteases such as papain or trypsin. To study the degradation behavior of AlMA hydrogels, we selected collagenase type 2 as a representative tissue dissociation enzyme (TDE). We observed that degradation behavior and hydrogel swelling were strongly affected by the UV cross-linking density (Figure 2j,k). Most AlMA hydrogels showed equilibrium swelling behavior within 24 h but afterward exhibited high swelling ratios (5−15%) due to high micro-porosity and absorption of water. Similarly, increasing AlMA polymer concentration and DM resulted in a significant reduction in swelling behavior due to high UV cross-linking density and the reduction of porosity (Figure S3). AlMA hydrogels with low DM (0.1 and 0.2%) exhibited rapid degradation behavior up to $\sim 90$% degradation within 1 h of incubation (Figure 2k), while medium-DM hydrogels (0.4−0.6%) degraded to $\sim 50$% after 72 h of incubation and high-DM hydrogels (2.0%) exhibited 80% remaining mass over the same period. Consequently, high-DM hydrogels with improved elastic moduli and reduced swelling behavior showed relatively slower degradation and were therefore selected for the following analyses.

Another interesting physical property of albumin is that it strongly binds with various biological substances such as growth factors, hormones, and fatty acids. Wettability has been shown to determine adsorption attributes such as kinetics, quantities, deformation, and reversibility. After measuring the contact angle of water, one can estimate the hydrophilicity, wettability, adhesiveness, and surface free energy of the AlMA hydrogels (Figure 2i). For the comparative analysis, two common hydrogels used in tissue engineering,
10% w/v PEGDA and GelMA hydrogels, were studied (Figure S4). The contact angles of AlMA hydrogels increased in proportion to the increment of the concentration of AlMA colloidal solution (Figure 2m). The 20% w/v AlMA hydrogel showed the highest angle of 124.0 ± 26.6° among the three polymers (65.5 ± 0.5° for GelMA hydrogel and 24.1 ± 0.1° for PEGDA hydrogel) as shown in Figure S4. The surface hydrophilicity can be estimated to be higher in the order of PEGDA > GelMA > low-concentration AlMA > high-concentration AlMA. According to our data, wettability and solid-surface free energy were also the lowest for the AlMA hydrogel compared to the PEGDA and GelMA hydrogels. We then observed the dominant interfacial forces from protein adsorption attributes under the influence of surface charge and wettability (Figure 2n and Figure S4c). Bovine serum albumin (BSA) and fetal bovine serum (FBS) were similarly adsorbed on all hydrogels without any significant difference. The GelMA (10% w/v) and AlMA hydrogels with two different concentrations (10 and 20% w/v) showed good adsorption compared with PEG hydrogels. To improve the adsorbed number of proteins on the hydrogel, both hydrophilic and hydrophobic properties that can induce various types of physical interactions, such as electrostatic interactions, hydrogen bonding, and hydrophobic intersections between proteins and the surfaces of hydrogels, are required, compared with hydrogels that can only possess highly hydrophilic or hydrophobic properties.

**In Vitro Characterization of the AlMA Hydrogels.** Albumin plays an essential role in transporting conjugated drugs to cells through caveolae-mediated endocytosis, particularly in specialized membrane architectures such as endothelial cells, fibroblasts, and smooth muscle cells.16,34 Aside from its central role as a functional carrier and ligand-binder in both in vivo and in vitro cellular environments, albumin has proven to be instrumental in alleviating reactive oxygen species (ROS)-induced oxidative tissue damage by binding to free radicals as well as providing protection against hydrodynamic stress induced by in vitro vessels (e.g., bioreactors).16 To ensure that AlMA hydrogels provide favorable ECM for cell proliferation and spreading, we next examined cellular behavior in hydrogels seeded with a representative subset of cell types whose successful culture is dependent on albumin-rich microenvironments such as those found in native blood vessels: human epithelial cells (hEPCs), rat cardiomyoblasts (H9C2), and NIH-3T3 fibroblasts. Unlike collagen or GelMA hydrogels, albumin does not contain cell-
binding sites such as Arg-Gly-Asp (RGD) peptides. However, AlMA hydrogels can adsorb and maintain abundant biological molecules, which are excised in the cell culture media or secreted from cells and could improve cellular viability and proliferation. According to our study, 10–20 wt % AlMA hydrogels at high DMs (2.0%) exhibited high levels of seeded cell viability (>~95%) (Figure 3a,b). Low-DM (0.6%) hydrogels, on the other hand, exhibited slightly lower viability (~80%), possibly due to their insufficient stiffness for supporting cell attachment and growth.35 We observed that increasing the DM had a major impact in increasing the elongated cell area on the surface of the AlMA hydrogels (Figure 3c,d). In the case of H9C2 cells, cell spreading in high-DM hydrogels was so significant on day 3 that the formation of interconnected cytoskeleton networks was observed throughout the surface (Figure 3d). Increasing the concentration of AlMA from 10 to 20% w/v led to a reduction in cell adherence and spreading by ~20 wt % in high-DM hydrogels and ~5% in low-DM hydrogels (Figure 3c). High-DM % w/v AlMA hydrogels exhibited comparable cellular spreading behavior to 10% w/v GelMA hydrogels (Figure 3c), both of which demonstrated an elastic modulus of ~5 kPa (Figure S2). With the help of photolithography, AlMA hydrogels can also be micropatterned with various shapes of masks for having a potential to fabricate complex ECM architectures (Figure 3e). NIH-3T3 cells seeded on the micropatterned AlMA hydrogels exhibited great elongation and interconnected cellular networks (Figure 3f and Figure S6). We observed a ~30% increase in cell area by increasing the concentration of AlMA from 10 to 20% w/v (Figure 3g), and this increase was consistent in 70 h of culture.

Next, we set out to encapsulate the representative cells (i.e., NIH-3T3, H9C2, and hEPC) into micropatterned AlMA hydrogels to form 3D constructs. Moving from a 2D culture to 3D constructs, photo-cross-linkable hydrogels are expected to exhibit a relative loss in cellular viability due to the mechanical stresses induced by the encapsulation process as well as increased UV exposure duration and PI concentration, which leads to the generation of free radicals.36 We encapsulated NIH-3T3 and H9C2 cells in high-DM 20% w/v AlMA hydrogels and observed a uniform distribution with high viability in micropatterned AlMA hydrogels (Figure 4a,b). To optimize UV exposure for maintaining cell viability, we constructed the hydrogels at two UV exposure durations (40 and 50 s) and measured the number of viable cells over the course of 96 h (Figure 4c,d). At higher UV exposure time, which yielded higher mechanical stiffness (Figure S3), cell-laden AlMA hydrogels showed a significant increase in the number of embedded cells, which suggested that the hydrogel stiffness was well suited to support cell proliferation. hEPC-laden AlMA hydrogels also demonstrated >90% cellular viability over the course of 24 h (Figure 4e and Figure S7). Moreover, encapsulated hMSCs and C2C12 showed an increase in metabolic activity rates up to day 5 (Figure 4f,g).

**AlMA Hydrogels Exhibit Successful Host Invasion and Degradation In Vivo.** Confirming the viability using various cell types encapsulated in AlMA hydrogels, we next explored the implantability of AlMA hydrogels. In a model of circulating cell invasion, AlMA and collagen (control) hydrogels of identical geometrical properties (cylindrical diameter, 8 mm; thickness, 1 mm) were implanted in the rat dorsal skin subcutaneously. After 14 days of in vivo culture, the AlMA implants exhibited a noticeable amount of host infiltration (Figure S5a). Analysis of explants harvested on day 3 revealed infiltration of circulating cells within the internal pores of AlMA hydrogel and was covered by a thin layer of fibrous capsule at the hydrogel interface (Figure S5a and Figure S8). On day 7, circulating cell infiltration intensity was considerably greater, and AlMA hydrogels were divided into fragments surrounded by collagen fibers as shown with
Masson’s trichrome staining (Figure 5a and Figure S9). Additionally, revealing a high number of CD31+ vessels, AIMA hydrogel implants demonstrated neovascularization in the peri-implant area (Figure Sb,c) in comparison with the collagen or sham + conditions (Figure S10). Higher proliferation of percentage of CD45+ cells was significantly greater in the AIMA group compared to the control group. Higher percentage of CD45+ cells was significantly greater in the AIMA group compared to the control group. Ligation of CD31 to the leukocyte surface is associated with the activation of functional leukocyte integrin, and leukocyte transmigration throughout the endothelium suggests an affinity for CD31 interaction. In other words, hydrogel based on albumin, a major protein in body serum, can rapidly collect circulating cells and has high binding ability. Further staining with CD45 after 14 days revealed that the number of leukocytes increased throughout the tissue comprising the peri-implant area (Figure 5d). The quantification revealed that the percentage of CD45+ cells was significantly greater in the collagen scaffold (Figure 5e).

CONCLUSIONS

In this study, we successfully designed and characterized human serum albumin-based photo-cross-linkable hydrogels (AIMA) that can be used for fabricating tissue-engineered implantable cell-laden constructs. In addition to their micro-porosity, the proposed AIMA hydrogels showed tunable porosity, the proposed AIMA hydrogels showed tunable biomechanical properties of target ECMs. Moreover, the hydrogel demonstrated a capability to support implant survival with increased survivability in vivo and served as a guide for host infiltration and rapid recruitment of hematopoietic cells. Overall, the results of the proposed AIMA hydrogels suggest a new biomaterial capable of tunable stiffness and degradation, therapeutic delivery to injured sites, and cellular ingrowth for regenerative medicine applications.

Exploiting albumin’s inherently valuable physiological functions such as binding to key biological compounds (e.g., vitamins, hormones, fatty acids, ions, and drugs), nanocarrier- and growth factor-conjugated scaffolds can be fabricated with enhanced half-life, making AIMA potentially suitable for wound closures or wound dressing applications. The high affinity toward numerous biomolecules makes AIMA a great candidate for the delivery of wound healing factors to the injured site. Furthermore, since there are albumin-based products that have already been approved by the FDA, obtaining approval for the clinical use of AIMA is projected to be relatively easier than other biomaterial-based scaffolds. In addition to the hemostatic and wound healing aspect, AIMA would also perform well as a photo-cross-linkable bioprintink for a wide range of 3D bioprinting applications that have emerged in the past decade. In this regard, rapid fabrication methods targeting large-scale scaffolds with intricate detail yet robust microarchitectures can benefit from using AIMA-based additive bioprinting techniques. Finally, tunability and ease of fabrication of AIMA hydrogels can be applied to the fabrication of geometries customized to the defect size to enable enhanced integration, angiogenesis, and tissue regeneration.

MATERIALS AND METHODS

Materials. Albumin from human serum (A1653, ≥96%; lyophilized powder), methacrylic anhydride (MA), and 3-(trimethoxysilyl) propyl methacrylate (TMSPMA) were obtained from Sigma-Aldrich (MO, USA). Dialysis membrane (Standard Regenerated Cellulose Membrane, Spectra/Por) was purchased from Spectrum Labs (CA, USA). Ultraviolet (UV) light curing system (Omniscure S2000) was purchased from EXFO Photonic Solutions Inc. (Ontario, Canada), and the photomasks with printed patterns used for hydrogel patterning were custom-made from CADart (Washington, USA).

Preparation of Human Serum Albumin Methacryloyl (AIMA). Human serum albumin methacryloyl (AIMA) was synthesized by substituting amine groups mainly in lysine residues with methacrylic groups as previously described with several modifications. Briefly, lyophilized human serum albumin was dissolved in distilled water (pH 7) in 5% w/v concentration at 4 °C and until albumin is completely dissolved. Methacrylic anhydride (MA) was added to the albumin solution at a rate of 200 μL/min to reach the specified target concentration (0.05, 0.2, 0.6, 0.8, 1, and 2% v/v) under constant stirring, and the reaction time was set to 2 h (4 °C) to minimize the precipitation. Then, additional distilled water was added, after which the resulting mixture was dialyzed (pH 6.5–7) using a 12–14 kDa cutoff dialysis membrane for three days. Maintenance of pH is crucial to preserve isoelectric point-induced albumin precipitation at around pH 4.7. The solution was finally lyophilized and stored at −80 °C before the experiments.

Characterization of AIMA. 2,4,6-Trinitrobenzenesulfonic Acid Assay (TNB5A). The degree of methacrylation (DM) of AIMA was calculated using 2,4,6-trinitrobenzenesulfonic acid assay (TNB5A) to quantify unreacted free amine groups and the percentage of the number of reacted amine groups divided by the number of free amine groups before the chemical modification was calculated. Glycine was used to generate a standard curve. The hydrogen map of amine groups substituted with methacrylic moieties was profiled by using 1H nuclear magnetic resonance (1H NMR) spectroscopy.

1H NMR. Chemically modified HSA was lyophilized and dissolved (10 mg/mL) in deuterium oxide (Sigma-Aldrich) and stored at 4 °C until NMR data acquisition. 1H NMR spectra were acquired at 25 °C using a Bruker 500 MHz spectrometer with a spectral width of 10,000 ppm, 128 scans, 4 dummy scans, and a total acquisition time of 1.64 s. Solvent presaturation was employed to minimize the impact of water on the spectrum. Phase and baseline corrections were manually applied to obtain purely absorptive peaks. The double peaks at 5.4 and 5.7 ppm were used as an indicator of incorporated hydrogens attached to the double bond of methacrylic anhydride.

Assessment of AIMA Aggregates. To induce self-assembled aggregates, AIMA (0, 0.05, 0.6, and 2% DM; 2 mg/mL) was dissolved in PBS and allowed to stir in the dark at room temperature. The hydrodynamic diameter and polydispersity index (Pdi) of the resulting self-assembled AIMA aggregates were measured by dynamic light scattering (DLS) using a nanoparticle analyzer (Malvern Zetacizer Nano ZS90, Malvern Instruments, Malvern, UK).

Transmission electron microscopy (TEM) analysis was conducted to study the aggregation of 20% AIMA. AIMA
macromolecule solution was applied to a charged carbon–formvar-coated grid and stained (1% uranyl acetate, 1 min). The samples were then examined with the TEM (Hitachi H7600, Tokyo, Japan) at 80 kV. High-DM AIMA solution specimens with a concentration range of 0.1 ng/mL–10 mg/mL were subject to TEM measurements.

Transmittance Study. The light transmitted through the different prepolymer solutions with different concentrations of MA and AIMA was measured using a UV–Vis spectrophotometer (OPTIZEN POP, Mecasys, Daejeon, Korea). For the measurement, the prepolymer solution was scanned (acquisition mode) from 200 to 900 nm at 5.0 nm intervals using quartz cuvettes with 10 mm path lengths. PBS was used as the blank. The collected absorbance data was used to calculate %T.

Preparation of Ultraviolet (UV)-Cross-Linked Hydrogels. Lyophilized AIMA was dissolved using phosphate-buffered saline (PBS) with 0.5% w/v 2-hydroxy-1-(4-(hydroxymethyl) phenyl)-2-methyl-1-propanone (Irgacure 2959, CIBA Chemicals, Basel, Switzerland) to generate free radicals to initiate the photo-cross-linking reaction. Microscope slides (Marienfeld, Germany) were acylated with TMSPMA to chemically anchor the hydrogels on the surface.41 After being fully dissolved, the prepolymer solution was dropped onto the TMSPMA-coated glass slide between two cover glasses separated by a 1 μm spacer and exposed to UV light (8.3 mw/cm², 360–380 nm) for 150 s. Immediately after the formation of the hydrogel, each sample was placed in PBS at 37 °C for 24 or 48 h. At each timepoint, the weight was recorded after removing excess PBS, and the samples were then lyophilized and weighed one more time. Finally, the mass swelling ratio was expressed as the ratio of swollen hydrogel mass to the mass of the dry polymer. The number of tested samples was 5 per group.

Enzymatic Degradation Profile. To determine the stability of hydrogel matrices under physiological conditions, polymerized hydrogels (55 μL, 20% w/v, and 150 s) were placed into a free-standing cylinder and incubated in 1 mL of 2.5 U/mL collagenase type II (Worthington Biochemical Corp., NJ, USA) at 37 °C with gentle shaking (< 150 rpm).43 The degradation profiles of the hydrogels were expressed as the percentage of remaining mass (%), which was calculated by dividing the initial mass (mg) of gels at 0 h by the remaining mass (mg) of gels at various timepoints based on dry weight recorded after freeze-drying. The hydrogels used in the analysis had varying DMs (27, 36, 51, and 58% DM; each corresponding to 0.1, 0.2, 0.4, and 0.6% MA used for the modification). Three replicates were made at nine different timepoints (0, 1, 3, 6, 10, 24, 35, 48, and 72 h after the incubation).

Contact Angle Measurement. To determine the surface hydrophilicity of the hydrogels, the surface glycerol contact angles of three different types of hydrogels were measured (room temperature) following the method described in the ASTM D5946 (“Standard test method for corona-treated polymer films using water contact angle measurements”) by using the Phoenix 300 Touch instrument (SEO). Samples were prepared with 10 and 20% w/v concentrations with 8 mm diameters and 1 mm thicknesses and directly cross-linked on the glass slide in a free-standing form. As a reference control, the same concentrations of polyethylene glycol diacrylate (PEGDA; MW 1000, Polysciences, Inc., USA) and GelMA hydrogels that are commonly used as biopolymers were chosen. The contact angle was measured by dropping 0.003–0.005 mL of glycerol solution onto the hydrogel matrices with a 27-gauge size needle and 3 mL volume syringe. Measurements were performed three times for each type of polymer and averaged with data variation expressed as standard deviation (SD) and coefficient of variation (CV). The range of measurement was 10–180° with an accuracy of 0.1°.

Protein Adsorption. Total protein content was calculated by performing a Micro BCA assay (Thermo Fisher Scientific, MA, USA). To measure the protein adsorption on hydrogel surface, test samples were prepared by pipetting (10 μL) prepolymer solution (0.5% PI) onto Culturewell chambered coverslips of 1 mm thickness (8 mm diameter) and exposed to UV light (8.3 mw/cm², 360–380 nm) for 150 s. To characterize the Young’s modulus E, the AFM indentation technique was applied. Here, the hydrogel sample was firstly immersed in 1× PBS in a petri dish mounted onto the AFM setup. Then, the AFM cantilever (DNP, Bruker, Camarillo, CA, USA) was positioned onto the sample with the assistance of an inverted microscope underneath the petri dish. The cantilever tip was then indented into the sample for the characterization. For each sample, six well-scattered points were selected randomly. Upon completion of the nanoindentation test, the corresponding force–depth curves were analyzed. Since the indentation depth in the micrometer scale is far larger than the radius of the AFM tip radius in the nanoscale, the theoretical model for conical indentation was properly applied to analyze the indentation results.42

Swelling Properties. For the measurement of swelling properties, the test specimens were fabricated by pipetting the prepolymer solution between two glass slides separated by a 1 mm spacer and exposed to UV light (8.3 mw/cm², 360–380 nm) for 150 s. Immediately after the formation of the hydrogel, each sample was placed in PBS at 37 °C for 24 or 48 h. At each timepoint, the weight was recorded after removing excess PBS, and the samples were then lyophilized and weighed one more time. Finally, the mass swelling ratio was expressed as the ratio of swollen hydrogel mass to the mass of the dry polymer. The number of tested samples was 5 per group.

In Vitro Studies. Cell Culture. For the cell culture of the H9C2 rat heart myoblast and the NIH-3T3 cell line (purchased from the Korean Cell Line Bank, Seoul, Korea),
Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS) and 1× penicillin/streptomycin (Welgene, Daegu, Korea) were used. Both cell lines were grown at 5% CO₂ and 37 °C conditions. Cells were passaged at least twice per week at subconfluence in a 1:3 ratio, and the media was changed once every 2 days. Human endothelial progenitor cells (hEPCs) were kindly provided by Prof. S.M. Kwon (Pusan University, Pusan, Korea).44,45 The hEPCs were cultured on gelatin-coated (1%) dishes in EC basal medium 2 (EBM-2, Lonza, Walkersville, MD, USA) supplemented with 5% FBS, EGM-2 growth factor mixture, and 1× penicillin/streptomycin (5% CO₂ at 37 °C).

**Cell Adhesion.** Five different types of biopolymers were used in the cell adhesion study for comparative analysis: GelMA, 10% AIMA (0.6 and 2% DM), and 20% AIMA (0.6 and 2% DM) hydrogels. Hydrogel sheets (8 mm diameter × 150 μm height) were fabricated onto TMSPMA-coated glass slides by using a UV intensity of 8.3 mW/cm² for 25 s. Polymerized hydrogels were washed with DPBS prior to incubation in culture media for 12 h. Cell suspensions containing 2.5 × 10⁵ cells/mL of cell lines (H9C2 or NIH-3T3) or primary cells (hEPC) were pipetted onto the hydrogel surface and incubated for 1 h prior to gentle washing with DPBS. Cells that adhered to hydrogel sheets were stained using a LIVE/DEAD Viability/Cytotoxicity Kit (Thermo Fisher Scientific, MA, USA) to evaluate cell viability after 24 h. On day 3, cells were fixed with 4% formaldehyde solution, and cytoskeleton structures were stained with Phalloidin-FITC (F-actin filament; Thermo Fisher Scientific, MA, USA) and DAPI (cell nuclei; Sigma-Aldrich, MO, USA).

**Cell Encapsulation.** For the 3D cell encapsulation study, cell lines (H9C2 or NIH-3T3) and primary cells (hEPC) were suspended in the 10 and 20% w/v AIMA prepolymer with different DMs at 2 × 10⁵ cells/mL. The cells containing prepolymer solution with 0.5% PI were exposed to a 8.3 mW/cm² UV light for 25 s on the TMSPMA-coated glass between 150 μm spacers. The cell-encapsulated hydrogels were thoroughly washed with DPBS and incubated for 4 days. Cell viability was measured by using a LIVE/DEAD Viability/Cytotoxicity Kit (Thermo Fisher Scientific, MA, USA).

**In Vivo Studies.** *Animal and Hydrogel Disk Implantation.* Male Sprague–Dawley rats (n = 30; 7 weeks old; 300–330 g) were purchased from Oriental Bio (Seongnam, Korea) and were maintained under a controlled environment in specific pathogen-free conditions. All experiments were carried out following protocols approved by the Institutional Animal Care and Use Committee of Konkuk University (approval no. KU 17025). Four hydrogel disks were subcutaneously implanted for each rat. Rats were anesthetized by continuous inhalation of 2% isoflurane gas. Incisions were made on the central dorsal area to reach the subcutaneous space. Then, subcutaneous pockets were created for the implantation of the hydrogel disks. After implantation, the skin incisions were closed using interrupted 3-0 silk sutures.

**Immunohistological Analysis.** The implanted hydrogels were harvested at days 3, 7, and 14 for each group and then fixed in Bouin’s solution (Sigma-Aldrich, MO, USA). The paraffin block samples were sectioned (3 μm) and permeabilized using 0.1% Triton X-100 (Sigma-Aldrich, MO, USA). Prepared sections were then stained with hematoxylin and eosin (H&E) to identify the nuclei gathering into the surrounding tissues and Masson’s trichrome staining for collagen and blood vessel distribution around the implanted hydrogel. Sections were further stained with CD31 (1:400, Novus, CO, USA) as a marker for blood vessels and were incubated overnight at 4 °C. CD45 (1:500, ab10558, Abcam, Tokyo, Japan) was used to examine the migration of macrophages into the transplanted hydrogels, and the sections were incubated overnight at 4 °C. Secondary Alexa Fluor 488-conjugated antibodies (Invitrogen, MA, USA) were diluted (1: 500) and reacted at room temperature for 2 h. The nucleus was stained by dilution to 1:10000 with DAPI (Sigma-Aldrich, MO, USA) with counterstain.

**Statistical Analysis.** All parametric data are indicated as mean ± standard deviation (SD). Unpaired Student’s t tests and one- or two-way ANOVA were performed to determine significant differences with the appropriate post-tests using GraphPad Prism5 (GraphPad, San Diego, USA). A P value of <0.05 was considered to indicate statistical significance.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.1c04292.

Supporting methods, cross-sectional SEM images, and elastic modulus of AIMA hydrogels; water contact angle images of 10% GelMA and 10% PEGDA hydrogels; F-actin and DAPI image of cultured H9C2 on the 10% GelMA hydrogel on day 3; cultured NIH-3T3 and H9C2 cell behavior on the 10 and 20% AIMA hydrogels; live/dead images of encapsulated hEPCs in the AIMA hydrogels; supporting in vivo characterizations (PDF)

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Author Contributions

Author Contributions

H.B. and S.R.S. designed the experiments; H.J.Y., S.Y.S., H.L., and Y.A.J. performed the experiments. H.B., S.R.S., H.J.Y., and Y.A.J. arranged the data and wrote the manuscript; H.B., S.R.S., H.J.Y., and Y.A.J. revised the manuscript. All authors significantly contributed to the study and to the interpretation of the data.

Notes

The authors declare no competing financial interest.

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