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## High-Strength Magnetic Hydrogels with Photoweldability Made by Stepwise Assembly of Magnetic-Nanoparticle-Integrated Aramid Nanofiber Composites

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**ABSTRACT:** Hydrogels capable of transforming in response to a magnetic field hold great promise for applications in soft actuators and biomedical robots. However, achieving high mechanical strength and good manufacturability in magnetic hydrogels remains challenging. Here, inspired by natural loadbearing soft tissues, a class of composite magnetic hydrogels is developed with tissue-mimetic mechanical properties and photothermal welding/healing capability. In these hydrogels, a hybrid network involving aramid nanofibers, Fe<sub>3</sub>O<sub>4</sub> nanoparticles, and poly(vinyl alcohol) is accomplished by a stepwise assembly of the functional components. The engineered interactions between nanoscale constituents enable facile materials processing and confer a combination of excellent



mechanical properties, magnetism, water content, and porosity. Furthermore, the photothermal property of  $Fe_3O_4$  nanoparticles organized around the nanofiber network allows near-infrared welding of the hydrogels, providing a versatile means to fabricate heterogeneous structures with custom designs. Complex modes of magnetic actuation are made possible with the manufactured heterogeneous hydrogel structures, suggesting opportunities for further applications in implantable soft robots, drug delivery systems, human-machine interactions, and other technologies.

**KEYWORDS**: biomimetic materials, magnetic hydrogels, NIR-welding, nanofibers, porous, soft actuators

#### INTRODUCTION

The burgeoning field of soft robotics calls for advanced materials that can be actuated in response to external stimuli.<sup>1–7</sup> Among the variety of soft active materials, magnetic hydrogels are particularly useful for biomedical applications.<sup>3,8-10</sup> They are water-rich, biocompatible, and tailorable in mechanical properties. With the assistance of highmagnetization nanoparticles, they can be remotely manipulated through a biocompatible low-intensity magnetic field, allowing for safe operation in living organisms. Magnetic hydrogels could be implemented in advanced tools for drug delivery, tissue engineering platforms,<sup>12</sup> human-machine interactions,<sup>2</sup> and many other technologies. However, existing magnetic hydrogels exhibit limited mechanical strength and manufacturability, which restrict their practical applications.<sup>1,13</sup> Indeed, the required inorganic magnetic components often disrupt the structural properties of hydrogels that highly rely on the

porous network of polymers.<sup>14</sup> On the other hand, the complex synthesis schemes for mechanically strong hydrogels might not be compatible with functional design of the composites or fabrication of hybrid device structures.<sup>15–17</sup>

In biological soft tissues, such as skin, blood vessels, ligaments, etc., an excellent combination of mechanical properties, functionality, and water retention was accomplished with the hybrid nanofiber networks involving collagen and other soft biopolymers.<sup>3,8,18,19</sup> They provide a robust structural basis for various physiological functionalization, allowing for

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Figure 1. Tough and stretchable magnetic hydrogels stemming from hyperconnective ANF composites. (a) Schematics showing the formation process of the hierarchical structures. The surface ligand plays an essential role in stabilizing the magnetic nanoparticle against strong dipolar interactions. (b) TEM image showing  $Fe_3O_4$ -NH<sub>2</sub> nanoparticles, where the aggregation was induced by the drop-casting method using DI water. Scale bar: 200 nm. (c, d) SEM images showing the morphology of the (c) ANF-PVA hydrogel and (d) ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA composite hydrogel after critical point drying, where the nanoparticles are embedded on the surface of the nanofibers. Scale bars: 1  $\mu$ m. (e) Photograph showing as-fabricated ANF-PVA (left) and ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA (right) composite hydrogels. (f) Photograph showing the NIR welded hydrogel sample. (g) Photograph showing programed magnetic response of the sample. Each magnet used has dimensions of 40 × 5 mm, with a center magnetic field strength of 40 mT and edge magnetic field strength of 160 mT. Scale bars: 10 mm.

the formation of heterogeneous organs that operate under dynamic mechanical deformation. Emulating these features of natural soft tissues would enable a versatile materials platform for the construction of reliable functional structures for advanced biointegrated systems. However, it remains difficult due to the limited synthetic building blocks.

Recently emerged nanocomposites based on aramid nanofibers (ANFs) indicate a route for the engineering of magnetic hydrogels that could address these challenges.<sup>8,20-23</sup> ANFs self-assemble into 3D networks that exhibit microstructural properties similar to those of natural collagen fibers, providing a biomimetic framework for the construction of various soft composites.<sup>8</sup> The polar groups on ANFs allow interactions with other organic materials, facilitating the formation of functional hydrogels that retain the high structural robustness arising from the nanofiber skeleton.<sup>20,24</sup> Nevertheless, incorporation of magnetic components in the ANF network is still difficult. For instance, in situ synthesis of magnetic nanostructures highly relies on surface chemistry, wherein the functional groups determine nanoparticle nucleation and growth. Due to the chemical stability of *para*-aramid (poly(*p*phenylene terephthalamide)) (PPTA) chains, chemical modification of ANFs requires complex and cumbersome procedures that might not be compatible with magnetic components.<sup>22</sup> Physical deposition methods,<sup>23,25,26</sup> including vacuum-assisted filtration and matrix infiltration, require diffusion of functional building blocks in aqueous or vapor phase. In the presence of the significant interactions between magnetic particles, these processes may cause severe aggregation or nonuniform distribution of the particles, leading to compromised functionality or structural failure by stress concentration. Therefore, developing strategies for reliable incorporation of magnetic components into ANF-based composite hydrogels becomes essential.

Herein, we report a facile fabrication strategy for ANF-based magnetic hydrogels with high mechanical strength and excellent manufacturability. Magnetite  $Fe_3O_4$  nanoparticles functionalized with alkyl chains terminated with amine  $(-NH_2)$  groups exhibit good stability in suspension while enabling desired interactions with ANFs. Stepwise assembly of ANFs,  $Fe_3O_4$  nanoparticles, and poly(vinyl alcohol) (PVA) via solution-based mixing processes leads to a magnetically responsive nanofiber network with excellent structural robustness and uniformity. In addition, the photothermal property of



Figure 2. Mechanical properties of an ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA magnetic composite hydrogel. (a) Photographs showing a sample of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-1 magnetic hydrogel with 0% (left) and 66.6% (right) tensile strains. (b) Photographs showing a sample of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-1 magnetic hydrogel without (left) and with (right) a compressive load of 20 N (2 kg). (c, d) Uniaxial (c) tensile and (d) compressive stress-strain curves for ANF-PVA and ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA magnetic hydrogels with four concentrations. (e) Tensile and compressive moduli of various samples. (f) SEM images showing the morphology of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogel samples with increasing content of Fe<sub>3</sub>O<sub>4</sub> nanoparticles (from ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-1 to ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-4). Scale bars: 1  $\mu$ m.

Fe<sub>3</sub>O<sub>4</sub> nanoparticles and the reconfigurable hydrogen bonding within the nanofiber hydrogels afford fast and reliable welding with near-infrared (NIR) radiation.<sup>27-30</sup> Alternatively, a functional moiety, such as nanoparticles or organic dyes<sup>3</sup> dispersed in the composites, might enable selective welding via laser irradiation. However, the use of noble nanoparticles as functional components dramatically increases the cost, whereas the organic dye molecules cannot sustain the high temperature required to soften the polymer composites. The use of Fe<sub>3</sub>O<sub>4</sub> nanoparticle in the composite hydrogels is particularly beneficial since it enables photothermal effects without additional components. NIR-laser-based photothermal welding enables production of Janus-type hydrogel structures or other complex and heterogeneous patterns, which allows for various modes of magnetic actuation with custom designs. The nanofibrous magnetic hydrogels also exhibit good biocompatibility, indicating potential applications in biointegrated device systems.

#### **RESULTS AND DISCUSSION**

**Synthesis of the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA Hydrogels.** To fabricate composite hydrogels with desirable properties in

bulk, a stepwise assembly strategy is employed (Figure 1a). Preparation of the composite hydrogel starts with intermixing the colloidal suspensions of  $Fe_3O_4$ - $NH_2$  nanoparticles and ANFs in dimethyl sulfoxide (DMSO). The hydroxyl and amine groups on the magnetite nanoparticles allow spontaneous packing with the ANFs via hydrogen bonding. The DMSO suspension of ANF- $Fe_3O_4$  is then mixed with a DMSO solution of PVA, which will further enhance stretchability and toughness of the nanofiber networks.<sup>8</sup> The ternary mixture appears as a viscous and moldable fluid (Figure S1), since DMSO is a strong hydrogen-bonding acceptor and prevents unwanted gelation at this stage. After the mixing steps, a solid hydrogel can be obtained when the mixture is immersed in water, wherein DMSO is exchanged by water, a weaker hydrogen bond acceptor.<sup>8</sup>

To match the diameter of the nanofibers, we synthesized  $Fe_3O_4$  nanoparticles with diameters of ~25 nm.<sup>24</sup> The nanoparticles are surface functionalized with alkyl chains terminated with amine and hydroxyl groups (Figure 1b). The magnetite  $Fe_3O_4$  lattices were further verified by X-ray diffraction (Figure S2), vibrating-sample magnetometer (Figure S3a), and transmission electron microscopy (TEM)



Figure 3. Photothermally induced welding of an ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA magnetic composite hydrogel. (a) Infrared images showing the temperature change of the hydrogel from 28.93 to 470.25 °C in 5 s under the illumination of a NIR laser. (b) Photographs showing the separated hydrogel thin films (left) and thick stripes (right) can be welded together by the NIR laser. (c) Stress-strain curves for the welded hydrogels. (d, e) Numerical simulation of (d) the spatial distribution of temperature on an array of Fe<sub>3</sub>O<sub>4</sub> nanoparticles and (e) surrounding environment under NIR laser irradiation. (f) Corresponding length-temperature plot showing temperature distribution on an array of Fe<sub>3</sub>O<sub>4</sub> nanoparticles under NIR laser, where the center part of the array shows the highest temperature due to the coupling effect. (g) SEM images at different welding stages showing that the interfaces are experiencing merging processes and finally fused together. Scale bars: 20  $\mu$ m. (h) Photographs showing an LED powered through a conductive hydrogel sample before and after the cutting-healing process. Scale bars: 2 cm.

ring diffraction pattern (Figure S4), which is consistent with theoretical calculations based on the reported lattice parameters (Table S1). A superparamagnetic response with a maximum of 74 emu/g was achieved for  $Fe_3O_4$  nanoparticles. Besides controlling the size of nanoparticles using hydro-thermal synthesis, there are other strategies to manipulate the morphology and composition of the magnetic nanoparticles, leading to high saturation magnetization for operation under a weak magnetic field.<sup>9</sup> For instance, thermal decomposition methods, introducing shape anisotropy, or adding zinc or cobalt elements<sup>32–34</sup> may be considered depending on the

functional requirement of the material system. During synthesis of  $Fe_3O_4$  nanoparticles, 1,6-hexanediamine was used as the modulator, which helps to control the size of the particles and produces sufficient alkyl chains terminated with amine groups on their surfaces. In deionized (DI) water, the surface alkyl chains closely pack on the surface of  $Fe_3O_4$ -NH<sub>2</sub> nanoparticles due to their poor solubility in water. The dissociation of amine and hydroxyl groups is thus limited, producing weak electrostatic repulsion between nanoparticles. The nanoparticles aggregate due to dominated dipolar and van der Waals attractions and settle to the bottom of glass vials within seconds (Figure S5), which contrasts with highly charged Fe<sub>3</sub>O<sub>4</sub>-COOH particles (Figure S6). However, in an organic solvent such as DMSO, the alkyl chains are highly dissolved and solvated. The steric repulsion of Fe<sub>3</sub>O<sub>4</sub>-NH<sub>2</sub> nanoparticles counteracts the dipolar attraction or van der Waals interactions, forming a stable colloidal suspension (Figures 1a and S6b).

Scanning electron microscopy (SEM) shows the microscale morphology of the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels. The Fe<sub>3</sub>O<sub>4</sub> nanoparticles attached to the nanofibers without significant aggregation or disruption of the 3D interconnected network (Figure 1c,d). It is conceivable that the nanofiber network provides an excellent template to guide the assembly of Fe<sub>3</sub>O<sub>4</sub> nanoparticles and acts as a robust structural framework to bear mechanical loads. We fabricated hydrogels with four different concentrations of loaded Fe<sub>3</sub>O<sub>4</sub> nanoparticles (7.3, 13.7, 24.1, and 32.2 wt %) denoted as ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-1 to ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-4. The magnetic response can be adjusted through controlling the composition of the hydrogel without major influence on the superparamagnetic properties of nanoparticles (Figure S3b,c). In addition, due to the inherent magnetic and photothermal properties of Fe<sub>3</sub>O<sub>4</sub> nanoparticles, the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA and ANF-PVA hydrogels can be further processed with NIR welding to form hybrid structures with designed magnetic responses (Figure 1e-g).

Mechanical Properties of the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA Hydrogels. Due to the hybrid nanofiber network, <sup>20,35</sup> ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels are mechanically strong and stretchable (Figure 2a,b). To quantitatively analyze their mechanical properties, we performed tensile and compression tests on pure ANF-PVA hydrogels and ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels with various concentrations of nanoparticles (Table S2). The magnetic composite hydrogels exhibit high strength (1.32-2.4 MPa) and stretchability (53.3-66.6%) similar to those of pure ANF-PVA hydrogels (Figure 2c-e). However, the Fe<sub>3</sub>O<sub>4</sub> nanoparticles functionalized with amine or hydroxyl groups might have a slight interference with the hydrogen bonding between ANFs and PVA, which plays a significant role in the mechanical properties of the hydrogels. With lower nanoparticle concentrations (7.3 or 13.7 wt % of dried samples), the fracture strains of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels are similar to that of the pure ANF-PVA hydrogel (Figure 2c). The energy dissipation characteristics of these ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels are also similar to those of pure ANF-PVA (Figure S7).8 Samples with higher nanoparticle concentrations (24.1 or 32.2 wt %) exhibit decreases in tensile strength and fracture strain (Figure 2e). From SEM examination, all of the samples retain the 3D nanofiber network originating from the ANF framework (Figure 2f). Further optimization of the magnetic responses and mechanical properties of the composite hydrogels can be achieved by tuning the composition based on specific device applications.

The compressive behaviors of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels are less dependent on the loading of nanoparticles (Figure 2d,e). Samples loaded with Fe<sub>3</sub>O<sub>4</sub> nanoparticles retain similar compressive responses to the pure ANF-PVA hydrogels (Figure 2d). Different from the tensile responses that mostly arise from the nanofiber network, the compression resistance of the hydrogels is also related to the trapping of interstitial water flow that is less dependent on nanoparticle loading.<sup>8</sup> Overall, the mechanical robustness of the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels is sufficient for device applications even with a high loading of Fe<sub>3</sub>O<sub>4</sub> nanoparticles.

Near-Infrared Laser-Induced Welding. Welding and fusing of functional hydrogels are important for the fabrication of hybrid devices or repairing of damaged structures. Existing processing methods for soft materials, including thermal annealing, solvent treatment, or chemical cross-linking, lack the capability of spatially selective welding.<sup>36,37</sup> In addition, given the good solvent stability and low thermal conductivity of ANFs combined with the water-rich feature of the matrix, these traditional methods are incapable of welding ANF-based composite hydrogels. On the other hand, incorporation of photothermally active components may enable an approach to selective welding of ANF-based hydrogels. Typical photothermally active components are exemplified by organic dyes and gold nanoparticles,<sup>38,39</sup> but their incorporation into the ANF-PVA-Fe<sub>3</sub>O<sub>4</sub> system may complicate the manufacturing process, increase the cost, or impact the mechanical properties. Interestingly, the photothermal properties of Fe<sub>3</sub>O<sub>4</sub> nanoparticles embedded in the PVA-rich composite hydrogels may enable a convenient method for selective welding through NIR laser irradiation.<sup>40,41</sup> Under the radiation of a NIR laser with a maximum power of 0.5 W, the temperature of the hydrogel ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-1 (7.3 wt % Fe<sub>3</sub>O<sub>4</sub> nanoparticle loading) increased to 470 °C in 5 s as recorded by an infrared thermal imaging system (Figure 3a). This temperature rise, which was also demonstrated in other samples (Figure S8), can be utilized to weld separate hydrogel strips and achieve high healing efficiency (Figure 3b,c). The rapid temperature elevation can be ascribed to the high density of Fe<sub>3</sub>O<sub>4</sub> nanoparticles on the surface of nanofibers, which can be further confirmed by numerical simulation on an array of nanoparticles (Figure 3df). The center of the array displays a higher temperature than the end part under the illumination of the NIR laser, due to coupling of Fe<sub>3</sub>O<sub>4</sub> nanoparticles. Moreover, the internal heating source dispersed within the hydrogel matrix is beneficial for the high temperature rise since the aramidbased materials have a low thermal conductivity.<sup>22,42</sup> Though significantly enhanced photothermal properties can be achieved by magnetic particle clusters,<sup>28-30</sup> when designing nanoparticle-hydrogel systems, a balance between enhanced photothermal properties and overall homogeneity should be handled carefully. Twofold factors are considered: first, the aggregation of nanoparticles is irregular and uncontrollable due to their strong interparticle interaction; second, such aggregation would induce structural failure or nonuniform heating, resulting in reduced efficacy and even defects due to local overheating.

Investigations on the microscale morphology revealed the healing/welding process (Figure 3g). Under the NIR illumination, the elevated temperature enhances the fluidity of the fibrous nanocomposites, providing driving forces that bypass the kinetic traps and reduce the free interfacial energy. The nanofibers connect and merge as they are softened by elevated temperature. The water-rich protective PVA layer mitigates the high transient temperature that can damage the fibers and provides a time window for operation. Further indicated by the change in external appearance of as-welded samples, the welding process results in densification of the networks. For thin film samples with 80  $\mu$ m thickness, the welded regions become more transparent, while the same regions for thick stripes (250  $\mu$ m in thickness) exhibit obvious shrinkage (Figure 3b). As a comparison, ANF-Fe<sub>3</sub>O<sub>4</sub> aerogels are easily burned by a NIR laser, even under low laser intensity (Figure S9). By increasing the illumination time, two hydrogels



Figure 4. Schematics of various recycling processes of aramid materials, including those based on chemical stripping, mechanical fibrillation, and NIR-aided welding.

are fused together, of which the interface becomes smooth but condensed.

The mechanical properties of healed hydrogels are dependent on the thickness of samples, as indicated by tensile tests (Figure S10a,b). Besides the homogeneous welding of the same type of hydrogels, our method can also realize "macroscopic assembly" by welding the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA and ANF-PVA hydrogel, forming heterojunctions, as indicated by their distinct colors (Figures 1e-g and S10c). Tensile stressstrain curves indicate 65.33% fracture strain is retained by the healed hydrogel sample, achieving a high healing efficiency of 98%, when comparing to the 66.60% fracture strain of the original hydrogel sample (Figure 3c). The heterojunction can resist up to 42.73% strain and 0.8385 MPa stress before fracture. Nonuniform nanoparticle distribution may cause insufficient heating at the interface, resulting in reduced healing efficiency. Compared with healed stripe samples that retain 94.8% tensile modulus and 98.09% fracture strain, thin film samples exhibit a significantly decreased healing efficiency in modulus (79.4%) and strain (69.94%) (Figure S11). Since the healing process is protected by PVA and interstitial water, under the same healing time, thinner films may experience more damage due to lower water content. To optimize the healing efficiency, finding the balance between water content, nanoparticle concentration, and welding power would benefit future investigations.

In addition, the hydrogels can be employed as soft ionic conductors with infiltrated sodium citrate. This conductive hydrogel is also healable using NIR-induced welding, and electrical conductivity of the hydrogel is maintained even under bending and twisting (Figure 3h). In addition to Fe<sub>3</sub>O<sub>4</sub> nanoparticles, we synthesized gold (Au) nanoparticles with a diameter of ~25 nm (Figure S12) to replace Fe<sub>3</sub>O<sub>4</sub> nanoparticles for the fabrication of an ANF-Au-PVA composite hydrogel (Figure S13), as gold nanoparticles are biocompatible<sup>43</sup> and exhibit good electrical conductivity.<sup>44</sup> The resistance change against strain exhibits a linear and monotonic increase when stretched up to 70%, the rupture strain of the ANF-Au-PVA hydrogel. With a gauge factor of 1.979, calculated by the resistance change divided by the strain, this linear trend shows a feasible electrical property as a strain sensor.

This welding and healing approach for aramid-based materials benefits the improvement of device fabrication and materials recycling (Figure 4). First, the traditional method to

recycle the aramid materials faces severe challenges, including high energy/solvent cost, harsh degradation conditions, and, specifically for aramid composites, an expensive purification process. Our method provides a reliable idea to recycle ANFs forming desirable shapes with high energy efficiency and without solvent-aided physical and/or chemical treatment.<sup>45</sup> Second, the ANFs have long been used as building blocks constructing highly connective networks as reinforcing components. It is not an easy task to dial in functionalities to such materials in a clickable manner or fabricate hybrid structures. In addition, the welding can be repeated multiple times without significant degradation of mechanical strength (Figure S14). Our strategy of heterogeneous welding provides a customizable, modular approach that integrates various components with distinct functionalities.

Soft Programmable Actuators. Due to the inherent magnetic nature of  $Fe_3O_4$  nanoparticles (Figure S3a), the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels can serve as programmable soft magnetic actuators (Figure S3b,c). Existing soft magnetic hydrogels usually achieve actuation patterns based on anisotropic assembly of magnetic nanoparticles with predesigned alignments.<sup>1</sup> However, the sophisticated assembly process may impose restrictions on device fabrication or mechanical performance. For ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels, due to their photothermal weldability, they can be integrated with nonmagnetic ANF-PVA hydrogels and achieve modulated, controllable actuation patterns through hybrid structures (Figure 5a). Actuation experiments were performed using nine combined cylindrical magnets with a total center magnetic field strength of 390 mT perpendicular to the operation plane under ambient conditions. The pure ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA stripe exhibits a bending motion under a magnetic field applied at one end (Figure 5b). To further demonstrate the magnetic actuation of the hydrogel samples, we tested the anisotropic responses of four hydrogel samples with heterojunctions under a magnetic field applied at the central position (Figure 5c-e, Videos S1 and S2). Through different welding patterns, the actuators show different transformations, demonstrating the possibility of programmable and customizable control. Our working principles also allow for the construction of more complex shapes and patterns. For example, a flower-shaped hydrogel is formed by ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA petals and an ANF-PVA core, potentially serving as a reversible robotic hand for grasping objects (Figure 5f). Compared with other magnetic



Figure 5. Magnetic actuation and biocompatibility of photothermally welded ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels. (a) Photographs showing the welding process of a flower-like heterostructure fabricated from ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA and ANF-PVA hydrogels. (b–f) Photographs showing reversible actuation of hydrogel structures with (b) pure ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA and heterostructures with (c) one, (d) two, or (e) three pieces of magnetic hydrogel patches, as well as (f) a complex flower-like structure. The actuation was induced by 9 magnets with a total magnetic field strength of 390 mT applied perpendicular to the plane. Each cylindrical magnet has a dimension of 20 × 5 mm. Scale bars: 4 mm. (g) Fluorescent images (top) and cell viability histogram (bottom) of fibroblasts cultured on an ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA magnetic composite hydrogel, measured on day 1, day 3, and day 5 after cell seeding. Scale bars: 100  $\mu$ m.

hydrogels based on self-healing strategies utilizing precursor loading, metal-coordination, hydrogen bonds,<sup>46–48</sup> etc., which are limited by sophisticated chemical processes and insufficient mechanical performance,<sup>46,49</sup> our methods exhibit a high healing efficiency, mechanical strength, and manufacturability that does not require complex chemical modifications.

The chemical stability of our hydrogels further limits hazardous waste production during polymer reconfiguration and shape morphing via photothermal effects, mechanical forces, and magnetic fields, ensuring a high degree of safety. This high stability of the hydrogels yields excellent biocompatibility even suitable for primary human cells such as chondrocytes. Though generally considered biologically safe,<sup>3</sup> studies have indicated that  $Fe_3O_4$  nanoparticles' concentration, sizes, and interaction with cells<sup>50,51</sup> influence the cytotoxicity. To further investigate the cytotoxicity and biocompatibility of the composite hydrogel, the fibroblast cell line NIH/3T3, a major cell line in connective tissues, was cultured on the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-1, ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-2, and ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-3 hydrogel samples without surface modification. From day 1 to day 5, no obvious difference is observed in cell viability (Figures 5g and S15). Though potential biocompatibility concerns may relate to high temperature rise, by controlling irradiation time and laser intensity, our magnetic hydrogel may also be utilized to

perform photothermal therapy at a temperature level around 50  $^{\circ}\text{C}.^{52,53}$ 

#### CONCLUSIONS

In conclusion, we have introduced a facile and scalable approach to the fabrication of mechanically robust and magnetically active hydrogels with potential applications in soft robots and remotely controllable biomedical actuators. Using NIR welding methods, we can achieve custom-designed heterogeneous structures with the possibility of complex modes of deformation. Our recent work has demonstrated the ability to integrate various microelectronic sensors and electroactive polymers onto ANF-PVA-based materials.<sup>21</sup> In combination with the soft electronics, the multifunctional hydrogel platform may achieve more sophisticated sensing and actuation capabilities in a closed-loop manner. The hybrid device platform could enable diverse applications in implantable surgical tools, human-robot interactions, controlled release of drugs, photothermal therapies, and many other technologies.

#### **METHODS**

Synthesis of Fe<sub>3</sub>O<sub>4</sub> Nanoparticles with Amine Groups on the Surface. To match the diameter of the aramid nanofibers, the Fe<sub>3</sub>O<sub>4</sub> nanoparticles with 20-30 nm diameter were synthesized via the reported strategy.<sup>54</sup> Briefly, 0.6 g of FeCl<sub>3</sub> as a ferro source was dissolved with 6.5 g of 1,6-hexanediamine and 2.0 g of anhydrous sodium acetate in 30 mL of glycol at 50 °C by vigorously stirring and sonification. The fully dissolved, transparent solution was transferred into a Teflon-lined autoclave and put into an oven at 198 °C for 6 h. The magnetite nanoparticles were then washed by ethanol and DI water via a magnet collection and dispersion process (2 to 3 times) to remove the residue reactant and finally dispersed in DI water. Due to the strong attractions between magnetic particles, the particles tend to form clusters and even aggregate. The particles are then washed by DMSO (three times) through a magnet-induced precipitation and redispersion process, forming a stable suspension prior to hydrogel fabrication. TEM (FEI Tecnai G2 20 scanning TEM) is employed to characterize the nanoparticles. X-ray diffraction measurements were performed using a Cu K $\alpha$  source ( $\lambda$  = 1.541 Å, Rigaku SmartLab) in the  $2\theta$  range of  $15-95^{\circ}$  using a scanning rate of  $5^{\circ}$  min<sup>-1</sup>. To prepare the samples, the as-synthesized Fe<sub>3</sub>O<sub>4</sub> nanoparticles were drop-casted onto the sample holder and dried at 60 °C under ambient condition before measurement. The magnetic characterization was performed on Fe<sub>3</sub>O<sub>4</sub> nanoparticles, ANF-Fe<sub>3</sub>O<sub>4</sub> composites, and ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels under a magnetic field from -2 to 2 T (Lakeshore VSM 7307).

Fabrication of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA Hydrogel. An ANF suspension (2 wt %) in DMSO was prepared using the reported method and mixed with the  $Fe_3O_4$  DMSO suspension.<sup>35</sup> The mixing ratio can be varied by changing the concentration of nanoparticles, while the volume of the suspension keeps constant. The mixture was stirred vigorously for 2 min to ensure adequate assembly with the Fe<sub>3</sub>O<sub>4</sub> particles through hydrogen bonding. Then a 10 wt % PVA DMSO solution with a volume equal to the ANF suspension was added, followed by another 2 min of stirring. This process is highly water sensitive and should be performed under an inert environment such as nitrogen gas (in a glovebox). Any water absorption will induce the microscale ANF gelation and lead to microphase separation, which is detrimental for their mechanical performance. The mixture was either coated as a film or cast in customized molds. The entire stirring and molding processes were performed on a hot plate (80 °C, with a magnet) to maintain good fluidity. The shaped mixture was then immersed in DI water to generate the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogel. Fabrication procedures for ANF-PVA and ANF-Au-PVA hydrogels are the same, where ANF-PVA hydrogels do not require addition of nanoparticles, and ANF-Au-PVA hydrogels add Au particles instead of  $Fe_3O_4$ . For SEM (Hitachi S4800 FEG SEM) characterization, supercritical  $CO_2$  drying was employed to remove water from the hydrogel matrix without further damage.

**Mechanical Test of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA Hydrogel.** The mechanical tests were performed under ambient conditions using a universal testing machine (Zwick Roell, German). For tensile tests, thick (250  $\mu$ m) and thin (80  $\mu$ m) rectangular strips of 5 mm in width and 6 cm in length were prepared using a laser cutting machine and tested under a strain rate of 100%/min. For the compression test, cylindrical samples of 10 mm diameter and 3 mm thickness were prepared by pouring and degassing mixed solution into Teflon molds. The compression tests were performed under a strain rate of 100%/min. For healed samples, the tensile tests were performed under a strain rate of 100%/min.

**NIR-Induced Welding of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA Hydrogel.** Prior to NIR welding, the ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogel samples were fabricated in bulk by the doctor blade coating method. After the solvent exchange process, the hydrogel samples were laser cut into desired shapes. The NIR welding is realized by the photothermal effect of  $Fe_3O_4$  nanoparticles. Two pieces of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogel films with the same width were overlapped along their long axis, followed by the NIR laser illumination (785 nm, Shanghai Laser & Optics Century, IRM785RMA-300FC). The NIR power and welding duration rely on the thickness of films and the demand of mechanical strength at the joint, both enhancing the NIR power and extending the welding duration result in denser and stronger joints.

**Magnetic Actuation Experiments.** The actuation was induced by 9 cylindrical magnets with a total magnetic field strength of 390 mT applied perpendicular to the plane. Each cylindrical magnet has dimensions of  $20 \times 5$  mm. Here, the as-prepared patterned hydrogel samples were laid on a plane, and the magnet set was held upright and moved to approach the sample for actuation behavior.

**Cell Culture and Fluorescence Characterization.** The hydrogel samples were cut and washed with ethanol and phosphate-buffered saline (Gibco pH 7.4 basic  $(1\times)$ ) prior to cell culture. Dulbecco's modified Eagle medium (DMEM, Gibco, high glucose), fetal bovine serum (Gibco, qualified, Brazil), and penicillin-streptomycin (Gibco, 10,000 U/mL) were mixed as received at 89%, 10%, and 1% volume fraction, respectively. Mouse embryonic fibroblast NIH 3T3 cells were cultured. And the samples were characterized on day 1, 3, and 5 during culturing. To characterize the samples, a LIVE/DEAD cell imaging kit (488/570) (Invitrogen) was used as received, and the cells were treated for 15 min under ambient room conditions. The florescence was characterized using a Nikon Eclipse Ci-L.

**Numerical Simulation.** COMSOL Multiphysics (version 5.4a) was used to simulate the temperature on and around light-heated particles. Seven closely attached  $Fe_3O_4$  nanoparticles with a diameter of 25 nm were placed in a 1000 nm × 1000 nm × 1000 nm box filled with hydrogel as media. The model involves modules of heat transfer in solids.

- $\nabla \cdot \boldsymbol{q} = Q$
- $q = -k\nabla T$

where q is the heat flux vector, Q is heating source power, T is temperature, and k is thermal conductivity. The upper half surface of spherical particles was endowed with a boundary heat source, which mimics light heating.

 $-\mathbf{n} \cdot \mathbf{q} = Q_{\rm b}$ 

The *n* is the boundary normal vector. The boundary source power,  $Q_b$ , was set as 1.810 W m<sup>-2</sup>. The sides of the box were set to be at room temperature, 293.15 K. Due to the high water content, the hydrogel was regarded to have the same heat transfer properties as deionized water, possessing a thermal conductivity of 0.59 W m<sup>-1</sup> K<sup>-1</sup>, while the thermal conductivity of Fe<sub>3</sub>O<sub>4</sub> nanoparticles was set as 5.9 Wm<sup>-1</sup>K<sup>-1</sup>. After these settings, the model can simulate the temperature distribution on and around particles, where we found the central part of the nanoparticle array to be more heated.

#### ASSOCIATED CONTENT

#### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsnano.3c03156.

Synthesis of Au nanoparticles; photographs of processing of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels; electron diffraction pattern, X-ray diffraction pattern of Fe<sub>3</sub>O<sub>4</sub> nanoparticles; magnetic hysteresis curves of Fe<sub>3</sub>O<sub>4</sub> nanoparticles, ANF-Fe<sub>3</sub>O<sub>4</sub> nanocomposite, and ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels; photographs of nanoparticle suspension stability; table for material composition of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels; cyclic tensile and compression tests of magnetic hydrogels; near-infrared images of ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA hydrogels heated by a near-infrared laser; SEM images of burned ANF-Fe<sub>3</sub>O<sub>4</sub> aerogel; photographs of a stretching process applied on welded hydrogel samples; mechanical test comparison of tensile stress and strain at break between ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA-1 hydrogels and the same hydrogels experiencing a cut-welding process; photographs and sensitivity test of ANF-Au-PVA composite hydrogel as a strain sensor; mechanical tests on heterojunction hydrogels after secondary welding; fluorescence microscope images of fibroblasts cultured on ANF-Fe<sub>3</sub>O<sub>4</sub>-PVA magnetic composite hydrogels (PDF)

Video for actuation of hydrogel heterostructures with one piece of magnetic hydrogel patches (MP4)

Video for actuation of hydrogel heterostructures with two pieces of magnetic hydrogel patches (MP4)

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#### **Author Contributions**

Zuochen W. conceived the research. Zuochen W., H.Z., H.L., and B.Y. planned the experiments. L.W. and L.X. supervised the research. Zuochen W. and H.Z. prepared the samples. H.Z. carried out actuation and cell experiments. M.S. carried out mechanical tests. Zhisheng W. and Y.W. carried out numerical simulation and discussed nanoparticle synthesis. L.X., Zuochen W., and H.Z. cowrote the manuscript with assistance from the other authors. Zuochen W. and H.Z. contributed equally to this work.

#### Notes

The authors declare no competing financial interest.

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