Robust Self-Healing Hydrogels Assisted by Cross-Linked Nanofiber Networks

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Given increasing environmental and energy issues, mimicking nature to confer synthetic materials with self-healing property to expand their lifespan is highly desirable. Just like human skin recovers itself upon damage with the aid of nutrient-laden blood vascularization, designing smart materials with microvascular network to accelerate self-healing is workable but continues to be a challenge. Here we report a new strategy to prepare robust self-healing hydrogels assisted by a healing layer composed of electrospun cross-linked nanofiber networks containing redox agents. The hydrogels process high healing rate ranging from seconds to days and great mechanical strengths with storage modulus up to 0.1 MPa. More interestingly, when the healing layer is embedded into the crack of the hydrogel, accelerated self-healing is observed and the healing efficiency is about 80%. The healing layer encourages molecular diffusion as well as further cross-linking in the crack region of the hydrogel, responsible for enhanced healing efficiency.

Self-healing is a key biological feature that prolongs the lifespan of living organisms. On human skin, for instance, a cut is able to heal spontaneously, where blood vascularization enables the delivery of nutrients for this recovery. Given increasing environmental and energy issues, mimicking nature to confer synthetic materials with self-healing property to expand their lifetime is highly desirable. Specifically, artificial microvascular systems have been created by some elaborate techniques such as soft lithography or direct-write assembly. And Sottos’ group incorporated the microvascular networks into epoxy coating to realize self-healing. Meanwhile, three-dimensionally cross-linked hydrogels with self-healing ability have aroused increasing interest owing to their potentials in diverse biomedical applications. In this system, self-healing usually occurs through the introduction of reversible covalent bonds, or non-covalent supramolecular interactions such as hydrogen bonds, π–π interactions, host-guest interactions or hydrophobic interactions. Despite great progress in this field, new facile strategies to endow hydrogels with combination of self-healing and mechanical toughness are still highly needed.

Herein, we developed cross-linked nanofiber networks filled with redox agents to mimic nutrient-laden blood vascularization to speed up the healing process of hydrogels with intrinsic self-healing capability. The hydrogels, prepared by using biomedical materials poly (1-vinyl-2-pyrrolidinone) (PVP) modified with O-carboxymethyl chitosan (O-CMC) and acrylamide (AAm), can bear external forces owing to their great mechanical strength and efficiently and autonomously heal themselves following damage, representing the rare examples of synthetic materials with combination of intrinsic self-healing capability and mechanical toughness. Moreover, inspired by healing process in human skin where damage can heal spontaneously with the aid of nutrient-laden blood vascularization, we designed a healing layer (a bilayer film) composed of electrospun cross-linked nanofiber networks to mimic blood vascularization, with oxidant and reductant loaded in separated layer to serve as nutrients (healing agents). The healing layer with cross-linked nanofiber networks could release and deliver redox agents efficiently through capillary force after embedded into the cracks of hydrogels, initiating further cross-linking reaction to rebond two hydrogel interfaces into an integrated one. Then, halved healing time without compromising mechanical properties of the hydrogels was realized. Importantly, the healing layer has the same composition as the bulk hydrogels, which can avoid chemical contamination and offer good affinity and hence endow high healing efficiency to the healing materials. This new facile strategy that employing electrospun cross-linked nanofiber networks containing redox agents mimic nutrient-laden blood vascularization to enhance self-healing performance of materials avoids the complex fabrication process required for self-healing microvascular systems, might contributing a technically simple and easy handling way to fabricate self-healing materials with useful applications.
amine and persulfide (Supplementary Fig. S1), initiating cross-linking properties. PVP has good biocompatibility and low toxicity, but poor mechanical strength limiting its potential application. Here we chose the cross-linking monomer AAm as the copolymerization monomer to address this issue. To fabricate Gel-1 bulk hydrogels, the ammonium persulfate (APS)/N,N,N',N'-tetramethylethylenediamine (TEMED) couple were used as the initiator. The redox reaction of APS can give two types of primary radical species from MED) couple were used as the initiator. The redox reaction of APS can give two types of primary radical species from the cross-linking monomer AAm as the copolymerization monomer to address this issue. To fabricate Gel-1 bulk hydrogels, the ammonium persulfate (APS)/N,N,N',N'-tetramethylethylenediamine (TE-MED) couple were used as the initiator. The redox reaction of APS and TEMED can give two types of primary radical species from amine and persulfide (Supplementary Fig. S1), initiating cross-linking reaction and profiting the formation of hydrogel networks25–27. To enhance healing performance of hydrogels, an electrospinning technique, a cost-effective approach for facile generation of nano-/micro-fibrous polymers28–32, was employed to prepare a cross-linked nanofiber bilayer film filled with both oxidant and reductant as a healing layer. As shown in Fig. 1b, an aqueous solution of Gel-1 pre-reaction product containing oxidant APS was electrosprun into nanofibers on a copper wire mesh collector, followed by the same implement for a Gel-1 pre-reaction product solution containing reductant TEMED, affording a bilayer film composed of a layer of Gel-1 nanofibers loaded with oxidant APS and the other layer of Gel-1 nanofibers loaded with reductant TEMED. Then, the Gel-1 bulk hydrogel was cut into two separated blocks, followed by embedding of the healing layer, to obtain a sandwich-structural hydrogel (Fig. 1c). Without any external intervention, these pieces merged into an integrated one much more quickly than in the situation without the healing layer. The morphologies and properties of Gel-1 hydrogels were thoroughly investigated. Scanning electron microscopy (SEM) measurements reveal cross-linked porous structure of Gel-1 bulk hydrogels (Fig.s 2a,b). As the ratio of [AAm]/[NVP] increases, the pore size of the hydrogel decreases, and so does its swelling capacity (Supplementary Fig. S6). Fig. 2c shows the storage modulus (G’) and loss modulus (G”) of the hydrogels as the functions of frequency at a fixed strain of 0.01%. The G’ values show a substantial elastic response and are larger than the G” values over the entire range of frequencies. Both moduli increase after introduction of small amount of cross-linker BIS (0.02 wt%). It is noted that the G’ value can be up to 0.1 MPa, comparable to the largest value reported for self-healing hydrogels37, suggesting that the hydrogels have high mechanical strengths. Strain sweeps shown in Fig. 2d reveal that the G’ value decreases and is slightly smaller than the G” value above the critical strain region of 0.42%, indicating a collapse of the gel state. Furthermore, the hydrogel can be stretched from 7.0 cm to 23.0 cm, and then shrunk back to 7.1 cm without any visible damage, indicating that it can be recovered rapidly (Figs. 2e–g). The results demonstrate that Gel-1 hydrogels have great mechanical strength and recover capability.

The self-healing behavior of Gel-1 hydrogels is shown in Fig. 3. When the bulk hydrogel is cut into separated pieces by a razor blade and then juxtaposed in contact, they will merge into an integrated one after a certain time without any external intervention. The effect of various preparation parameters on the healing rate of self-healing hydrogels was investigated, as shown in Fig.s 3a–c and Supplementary Table S1. As the ratio of [AAm]/[NVP] or the content of cross-linker BIS increases, the healing rate of the hydrogels decreases (Figs. 3a,b). As shown in Fig. 3a, the hydrogels prepared from [AAm]/[NVP] = 1:1, 2:1, and 3:1 mol/mol, respectively, show healing time of 1 day (d), 2d, and 3d, respectively. These results are consistent with the previous report that the chemical hydrogels with high cross-link density lack the ability to self-heal37. In addition, chitosan is a bioadhesive molecule that it can strongly bind surrounding tissue through hydrogen bonding. In our case, the increase in O-CMC concentration leads to the increase in the healing rate of the hydrogels (Fig. 3c). This indicates that the presence of chitosan increases the healing rate, which is in good agreement with the results in the literature34. Specifically, when the hydrogel prepared from NVP = 0.04 mol, BIS = 0.02 wt%, and O-CMC/monomer = 2.5% (w/w) was cut by a blade, it spontaneously healed within 15 s (Fig. 3d–f; Supplementary Movie S1). The width of the wound upon fresh cut was 119.2 μm (Fig. 3d), and then became 105.2 μm after 5 s (Fig. 3e). With increasing time, the wound disappeared fast, and the damaged areas self-healed completely within 15 s (Fig. 3f).

Significantly, accelerated self-healing was observed after the healing layer composed of electrosprun cross-linked nanofiber networks with redox agents was embedded into the damaged area of hydrogels (Fig.s 1c and 3a–c). For example, as the healing layer was introduced.

**Figure 1** Schematic preparation and enhanced self-healing procedure of Gel-1. (a) Synthetic scheme of Gel-1. (b) Fabrication of Gel-1 healing layer formed of nanofiber networks. (c) Enhanced self-healing procedure upon embedding a healing layer into the damage area of Gel-1 bulk hydrogels.
into the cracks of the hydrogels prepared from [AAm]/[NVP] ratios of 3:1 and 1:2 (mol/mol), respectively, the structural integrity of the corresponding hydrogels could be restored after 2 d, 1 d, and 0.5 d, respectively (Fig. 3a), noting that the embedment of the healing layer reduces half of the healing time for the hydrogels. This finding also accords with the other runs (Fig.s 3b,c, and Supplementary Table S1), indicating that the healing layer plays a vital role in enhancing healing rate. Moreover, as the healed cracks recut, the self-healing process occurred once again, suggesting this self-healing process could be repeatable.

To investigate the healing efficiency of the hydrogels upon embedding with the healing layers, we constructed a 7.0-cm-long column by juxtaposing hydrogel blocks (transparent), healing layer (not seen), and hydrogel blocks doped with rhodamine B (red) in sequence and alternately (Fig. 3g). Even though the blocks were not cut freshly, the healing layer easily stuck them together. The resulting column is very stable, even not damaged after bended by hands (Fig. 3h). Additionally, we connected together two 2.5 cm-long blocks inserted with a healing layer in the same way (Fig. 3i). As shown in Fig. 3j, the joint between the two blocks in the merged sample was strong enough to sustain vigorous stretching. The tensile tests conducted on the original hydrogel gave an elongation ratio at break of 773 ± 38% at 136.9 ± 3 kPa, while those on the healing layer-embedded healed hydrogel gained an elongation ratio at break of 626 ± 32% at 111.5 ± 2 kPa (Supplementary Table S2). The elongation ratio of the healed hydrogel is similar to the original one and the healing efficiency of this hydrogel is about 80%. The results suggest that the introduction of the healing layer into the damaged area of hydrogels can highly accelerate self-healing without compromising mechanical properties of the hydrogels.

**Discussion**

What is the origin of the self-healing nature of Gel-1 hydrogels and the enhanced self-healing performance of hydrogels by embedding the healing layer? We propose that self-healing nature of Gel-1 hydrogels could be attributed to hydrogen bonding across two hydrogel pieces. As seen in Supplementary Fig. S7a, two Gel-1 hydrogel pieces can be merged into an integrated one through hydrogen bonding between amide, hydroxyl, and carboxylic functional groups. To testify our proposal, we immersed the self-healed Gel-1 hydrogels into the cracks of the hydrogels prepared from [AAm]/[NVP] = 3:1, 2:1, and 1:1 mol/mol, respectively, the structural integrity of the corresponding hydrogels could be restored after 2 d, 1 d, and 0.5 d, respectively (Fig. 3a), noting that the embedment of the healing layer reduces half of the healing time for the hydrogels. This finding also accords with the other runs (Fig.s 3b,c, and Supplementary Table S1), indicating that the healing layer plays a vital role in enhancing healing rate. Moreover, as the healed cracks recut, the self-healing process occurred once again, suggesting this self-healing process could be repeatable.

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into an aqueous solution of 30% urea to disrupt the hydrogen bonds. As expected, the two blocks of the self-healed Gel-1 hydrogels were separated after immersed in the urea solution (Supplementary Fig. S7b).

We also explored the mechanism for enhanced self-healing performance of hydrogels by embedding the healing layer. Field emission scanning electron microscopy (FESEM) measurements show that the original Gel-1 pre-reaction product nanofibers have an average diameter of about 100 nm (Fig. 4a, and Supplementary Fig.s S7, S8). Whereas, the nanofibers in the healing layer of bilayer film loaded redox agents display cross-linked network structure with increased diameter of 300 nm (Fig. 4b), which are further cross-linked and swollen with diameter of ca. 500 nm after embedded into the hydrogels (Fig. 4c). These features suggest the occurrence of cross-linking reaction initiated by APS and TEMED filled in the nanofibers. This result implies the presence of redox agents in the nano-network healing layer and their implement in cross-linking reaction to further form hydrogel networks. To further confirm this point, we prepared a sample coating with a healing layer composed of a nanofibrous bilayer film at the outside of the hydrogel and that without healing layer for comparison. Additionally, in order to explain the role of nano-networks, we fabricated a bilayer film without nanofiber structure using spin-coating method, with the same composition as the bilayer film with nanofiber structure, as a negative control. After immersed in Gel-1 pre-reaction product aqueous solutions containing 0.05 wt% rhodamine B for 1 d, the sample coating with the nanofibrous bilayer film absorbed the most amount of the Gel-1 pre-reaction product solution, to form the most amount of new hydrogels around the original one (Supplementary Fig. S10). This experiment indicates the nano-network healing layer promotes the release of redox agents which initiate cross-linking reaction to further form more hydrogel networks.

Diffusion has been reported to be the most critical stage of crack healing process, and we found that the introduction of the healing layer with cross-linked network into the hydrogels facilitates molecular diffusion. As shown in Fig.s 4d,e, two-block hydrogels with and without the healing layer embedded into the merged interface were prepared, respectively, where one hydrogel block in each kind of...
healed hydrogels was doped with fluorescent dye marker rhodamine B. Upon UV excitation (\(\lambda = 302\) nm), the hydrogel blocks with rhodamine B show bright orange PL emission (580 nm), whereas the hydrogel blocks without rhodamine B exhibit blue PL emission (380 nm) (Figs. 4d,e and Supplementary Fig. S11). After the healed hydrogels without healing layer stored for 7 d, rhodamine B diffused from the position of 3 cm (the boundary of the two blocks) to the position of 3.2 cm. Whereas, for the healed hydrogels embedded with healing layer, rhodamine B diffused to the position of 3.8 cm. These reveal the molecular diffusion in the sample embedding with the healing layer is much faster than in the sample without healing layer. These diffusion features can also be confirmed by fluorescence spectra, where the sample embedding with the healing layer shows relatively lower fluorescent intensity at the position of 2.8 cm and higher fluorescent intensity at the position of 3.2 cm in comparison with those without healing layer, respectively (Fig. 4f). The above results indicate the use of electrospun cross-linked nanofiber networks with redox agents as healing layer to embed between two hydrogel blocks can accelerate the molecular diffusion from one hydrogel block to the other based on capillary force, through which redox agents can release efficiently and disperse across the interfaces of hydrogel pieces to initiate further cross-linking and rebond two hydrogel interfaces into an integrated one, and hence accelerated self-healing is achieved.

In conclusion, we synthesized O-CMC-modified poly(NVP-co-AAm) self-healing hydrogels with high mechanical strength. By adjusting the preparation parameters in this system, the self-healing processes with healing rate ranging from seconds to days were available, which is meaningful for the application of self-healing hydrogels. Abundant hydrogen bonding in this cross-linked system might be responsible for the intrinsic self-healing capability of hydrogels. Moreover, we developed a new facile method to accelerate the healing rate of hydrogels based on the introduction of electrospun cross-linked nanofiber networks containing redox initiators as the healing layer, mimicking nutrient-laden blood vascularization enabling spontaneous recovery of a cut in human skin. This approach shows the unique advance that cross-linked nanofibers profit the molecular diffusion along the boundary of the damaged blocks and the released redox agents initiate further cross-linking across the rupture, and thus reduce half of the healing time. Also importantly, the healing layer has the same composition as the bulk hydrogels, avoiding chemical contamination to the healing materials. With healing efficiency up to 80%, the mechanical strength of the healed hydrogels is similar to the original one. This work opens promising avenues to achieve the potential development of more appropriate self-healing hydrogels with electrospun cross-linked nanofiber networks, and thus might be adapted for more practical applications, for instance, in controlled drug delivery materials, regenerative medicines and wound dressings.

**Methods**

**Materials.** Acrylamide (AAm; 97%), 1-vinyl-2-pyrrolidinone (NVP; 99%), N,N'-methylenebisacrylamide (BIS; 98%), ethanol absolute (99.5%), and N,N,N',N'-tetramethylethylenediamine (TEMED; 99%) were purchased from Sigma-Aldrich and used as received without any further purification. O-carboxymethyl chitosan (O-CMC) was obtained from Zhejiang AoXing Biotechnology Co. Ltd. (China) with a substitution degree of 0.8, \(M_n\) of 9 000, and used without any further treatment. Ammonium persulfate (APS; 98%) was ordered from Sigma-Aldrich and purified by recrystallization in distilled water before use. High-purity water with the resistivity of greater than 18 M\(\Omega\) cm\(^{-1}\) was used in the experiments.

**Preparation of Gel-1 bulk hydrogels.** The appropriate amounts of AAm, NVP, O-CMC, and APS were mixed in a certain amount of water to form a homogeneous solution in a container. A typical composition was water = 60 wt%, \([\text{AAm}]/[\text{NVP}] = 3 : 1\) (mol/mol), O-CMC/monomer = 2.5% (w/w), and APS = 0.04 wt%. After the
reagents TEMED ([APS]/[TEMED] = 1:4 mol/mol) was added and mixed homogeneously, the solution was sealed in the container and placed at ambient temperature without stirring until the hydrogel was formed. After the completion of reaction, the samples were purified by immersing in deionized water for 7 days to remove all water-soluble materials. Then the samples were dried in a vacuum oven at 60°C until the weight of the specimen was constant.

Synthesis of Gel-1 pre-reaction products. Gel-1 pre-reaction products were prepared by aqueous free radical copolymerization of AAm and NVP with O-CMC using APS as an initiator in a four-necked round flask with a mechanical stirrer, a thermometer, a reflux condenser, and a nitrogen gas-introducing tube. A typical synthesis of Gel-1 pre-reaction products is as follows: A mixture of 5.63 g AAm, 2.93 g NVP, and 0.22 g O-CMC was dissolved in 40 g purified water in a 100 ml round flask, and the stirring speed was set at 250 rpm. The flask was immersed in an oil bath set at 70°C. The reaction mixture was purged with nitrogen for 20 min. Once the reaction temperature had been reached, polymerization was initiated by 10 g aqueous solution containing 0.04 g APS. The polymerization reaction was allowed to continue for 2 h. The resulting viscous solution was then allowed to cool and was subsequently precipitated in an excess amount of ethanol and separated by centrifugation. The precipitate was washed with ethanol several times to remove the unreacted monomers, and then the precipitated Gel-1 pre-reaction products were dried in a vacuum oven at 60°C overnight.

Fabrication of electrospun cross-linked nanofiber networks. The electrospinning apparatus used in this work consists of a high-voltage power, a syringe, an 18 gauge stainless steel needle and a grounded stainless steel plane collector (wrapped with a copper wire mesh) mounted on an insulating stand at a distance of 15 cm from the needle. The electrospinning solution was maintained at a feed rate of 0.5 mL/h under a constant potential of 15 kV. The ambient temperature was 25°C and the humidity was 55%. The cross-linked nanofiber networks contained two layers of nanofibers. The layer 1 was electrospun from a aqueous solution of Gel-1 pre-reaction product (12 wt%) containing oxidant APS (0.1 wt%) on the copper wire mesh for 2.5 h, and then the layer 2 was electrospun from a Gel-1 pre-reaction product aqueous solution (12 wt%) containing reductant TEMED (0.2 wt%) over layer 1 for 2.5 h, affording a two-layered film composed of a layer of Gel-1 nanofibers loaded with oxidant APS and the other layer with Gel-1 nanofibers loaded with reduced TEMED. Finally, the obtained cross-linked nanofiber networks were peeled off from the copper wire mesh.

Characterizations. To further identify the chemical structures of the samples, fourier transform infrared spectra (FT-IR) were recorded on a Nicolet 6700 spectrometer with KBr pellets in the 400–4000 cm⁻¹ region. 1H high-resolution magnetic-angle spinning (HR-MRS) NMR spectra were collected at 400 MHz on a Bruker Avance 400D instrument. Molecular weight distributions were analyzed by Gel permeation chromatography (GPC, Waters Instruments) with an Ultrahydrogel column. The morphology of the electrospun hydrogel nanofibers was observed by field emission scanning electron microscopy (FESEM) with a Hitachi S4800 instrument at 20 kV. Photographic images of the samples were captured by digital camera (SX210 IS, Canon) or a stereomicroscope (SZM45). The structures of the bulk hydrogels obtained were examined using a QUANTA 200 SEM (Philips-FEI, Holland) operated at 20.0 kV. The samples used for SEM measurement were immersing in distilled water after synthesis to remove the residual unreacted monomers. During this period, the distilled water was changed daily until the samples reached the maximum swelling ratios. The samples were then freeze-dried for 12 h. Dried samples used for SEM measurement were cut to expose their inner structure. The swelling properties of the as-prepared bulk hydrogels were measured based on gravimetric analysis. The rheological experiments of the bulk hydrogels were performed on a rheometer (Avesta) at 25°C under the following conditions: crosshead speed = 15 mm/min, sample length between jaws = 48 mm. The tensile strength and percentage elongation at break were recorded. For reproducibility, at least six samples were measured for each hydrogel sample and the results were averaged.

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Author contributions
S.C. conceived the project and designed the experiments. Y.F., Z.-H.Z. and H.S. performed the experiments and analyzed the data. Y.F., C.-F.W. and S.C. contributed to the interpretation of the results and wrote the paper.

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