Research Article

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Design and preparation of 3D printing intelligent poly N,N-dimethylacrylamide hydrogel actuators

https://doi.org/10.1515/epoly-2020-0033
received February 21, 2020; accepted April 23, 2020

Abstract: The intelligent poly N,N-dimethylacrylamide hydrogel material system with high mechanical strength and the 3D printable property was prepared via in situ free radical polymerization under vacuum successfully. With the increase in nanofibrillated cellulose (NFC) content, stress and strain of hydrogels increased gradually. As the effective reinforcement, NFC enhanced the crosslinking density, which realized the controllable regulation of rheology behaviors including viscosity, storage modulus, and loss modulus of hydrogels. Combined with the swelling rate and the existence of the gel–sol transition point, a hydrogel with 10 mg/mL NFC was treated as the 3D printing ink of hydrogel actuators. Variation of printing parameters significantly affected self-driven deformations. The hydrogel actuators with 90°/0° and 45°/135° configurations owned bending and spiral deformations, respectively. Actuators with a larger length–width ratio owned a lower pitch value. The precise anisotropic swelling property of the printed bilayer structure was the self-driven deformation mechanism of hydrogel actuators, which provided material candidates for the preparation of soft robots and actuators.

Keywords: hydrogel, 3D printing, mechanical strength, printing parameter, intelligent deformation

1 Introduction

As a kind of typical intelligent soft materials, intelligent hydrogels owned the reversible swelling and deswelling properties under the stimulations of temperature (1,2), pH (3), light (4–6), magnetic field (7,8), and electric field (9,10) to realize shape and volume variation. The efficient and repeatable deformation properties expanded the potential application of intelligent hydrogels in fields of soft actuator (11–14), soft robot (15), artificial muscle (16), and so on.

To realize the functional deformation properties, many kinds of methods were adopted to prepare intelligent hydrogels (17,18). In situ free radical polymerization was the main method to realize the polymerization of intelligent hydrogels. Based on the photopolymerization, a series of temperature-driven intelligent hydrogels were prepared (19,20). In the deformation structures, the bilayer structure was the simplest structure to realize the anisotropic bending property. Via the photothermal conversion function of graphene oxide, the near-infrared laser-driven bilayer intelligent hydrogels were prepared by in situ polymerization (21), exhibiting the multiple deformation patterns. The layered structure constructed by in situ free-radical polymerization and molding provided the effective anisotropic deformation mechanism for intelligent hydrogels. But, the disadvantage of relatively simple structure patterns restricted the potential application of intelligent hydrogels. The 3D printing technology owned the advantages of complex structure construction and accurate sample dimension, which provided effective solution for the disadvantage of molding. Combined with the ultraviolet light polymerization, the 3D printed hydrogel actuators imitated the deformation patterns of tendrils and flowers to realize the anisotropic swelling deformations (22,23). Based on changing printing parameters,
the bilayer intelligent hydrogel actuators were prepared via 3D printing. The relative complex printed structure realized deformations based on swelling and deswelling anisotropy. The deformations of 3D printed intelligent hydrogel actuators were determined by the hydrogel viscosity, printing parameters, and structure design (24). The relative low viscosity enhanced the printable property and restricted the printed structure fixation property of hydrogels. The relative high viscosity enhanced the printed structure fixation property and restricted the printable property of hydrogels. The controllable regulation of rheological behavior built the preparation base of hydrogel actuators. The structure design and printing parameters determined the final deformation patterns. Therefore, optimizing the printing properties was the key point for the 3D printing of intelligent hydrogels (23).

The mechanical strength was the deformation base of intelligent hydrogel actuators. Many kinds of intelligent hydrogels have been used for the preparation of hydrogel actuators. As a typical kind of hydrogel material with high mechanical strength, the poly N,N-dimethylacrylamide (PDMAA) hydrogel was widely used in the preparation of hydrogel actuators. Moreover, the physical crosslinking PDMAA hydrogel was used in 3D printing. As a kind of effective reinforcement, nanofibrillated cellulose (NFC) was adopted to enhance the mechanical strength of intelligent hydrogels. Therefore, changing the rheological behavior to satisfy 3D printing and improving the mechanical strength of PDMAA hydrogel further were the key points for the preparation of 3D printing intelligent hydrogel actuators.

In this study, a series of novel NFC-reinforced PDMAA hydrogels were prepared via physical crosslinking. Combining the design of printing parameters, the intelligent hydrogel actuators realized 3D printing by constructing the relationship between NFC content and rheological behaviors.

2 Experimental

2.1 Materials

The N,N-dimethylacrylamide (DMAA, C₅H₉NO; Aladdin, Shanghai, China), nanosized synthetic hectorite clay (Laponite XLG, Mg₃.₃₄Li₀.₆₆Si₆O₂₀(OH)₄; Rockwood, Ltd, Germany), potassium peroxydisulfate (KPS, K₂S₂O₈; Shanghai Aibi Chemical Reagent Co., Ltd, Shanghai, China), and N,N,N‘,N‘-tetramethylethylenediamine (TEMED; Tianjin Weiyi Chemical Technology Co., Ltd, Tianjin, China, 98%) were treated as a monomer, a crosslinking agent, an initiator, and a catalyst, respectively. NFC (Guilin Qihong Technology Co., Ltd, Guilin, China, 1,342 nm) was treated as reinforcement to regulate the rheological property of hydrogels. Pure water was obtained by deionization and filtration with a Millipore purification apparatus (resistivity ≥18.2 MΩ cm).

2.2 Synthesis of hydrogel actuators via 3D printing

To investigate whether the addition of NFC affected the synthesis of intelligent hydrogels or not, in situ free-radical polymerization was adopted to prepare PDMAA hydrogels. About 3 mg of methyl blue was added into pure water for dye of hydrogels. Also, the XLG was added into pure water with methyl blue. Then, the mixture was stirred for 1 h and ultrasonically radiated for 30 min. The NFC with various contents was added and stirred for 1 h under the ice–water bath environment. Then, the DMAA was added and stirred for 1 h. After the addition of KPS and TEMED and stirring for 5 min, the prepared hydrogel reaction solution was put into a rubber mold with a dimension of 70 × 20 × 2 mm (length × width × thickness). The mole ratio of monomer, initiator, and catalyst was kept at 100:0.370:0.638. After 24 h of polymerization in 25°C, the prepared intelligent hydrogels were mold unloading. Based on the NFC contents of 0, 10, 11, 12, 13, and 14 mg/mL, the intelligent hydrogels were defined as NFC0, NFC10, NFC11, NFC12, NFC13, and NFC14, respectively. The compositions of intelligent hydrogels with different NFC contents are listed in Table 1.

The molding preparation provided material reference for 3D printing. To investigate the deformation of 3D printing intelligent hydrogel actuators, the printing structure models were designed. Before 3D printing of intelligent hydrogels, the STL date was produced by SolidWorks. Slic3r software was used for slicing. The laboratory-made 3D printer with stainless steel needle of 20G (inner diameter of 0.6 mm) was controlled by the open source software of Prontface. The printed intelligent hydrogels were put into a vacuum dryer at 25°C for 24 h to realize complete polymerization.

2.3 Characteristics

2.3.1 Microstructure

After removing water in hydrogels in a completely swollen state via a freeze-drying oven (LGJ-10C; Beijing
Four Ring Scientific Instrument Factory Co., Ltd, Beijing, China), the microstructure of intelligent hydrogel was observed by the scanning electron microscope (Model Evo18 Carl Zeiss, Oberkochen, Germany).

### Table 1: Compositions of intelligent hydrogels

| Sample | H₂O (mL) | DMAA (mL) | XLG (g) | NFC (mg) | KPS (mg) | TEMED (µL) | Methyl blue (mg) |
|--------|----------|-----------|---------|----------|----------|------------|-----------------|
| NFC0   | 17.5     | 1.97      | 0.65    | 0        | 20       | 30         | 3               |
| NFC10  | 17.5     | 1.97      | 0.65    | 200      | 20       | 30         | 3               |
| NFC11  | 17.5     | 1.97      | 0.65    | 220      | 20       | 30         | 3               |
| NFC-2  | 17.5     | 1.97      | 0.65    | 240      | 20       | 30         | 3               |
| NFC13  | 17.5     | 1.97      | 0.65    | 260      | 20       | 30         | 3               |
| NFC14  | 17.5     | 1.97      | 0.65    | 280      | 20       | 30         | 3               |

### 3 Results and discussion

#### 3.1 Microstructure

After freeze-drying, the microstructure of hydrogel was maintained integrally. Figure 1 shows the microstructure characteristics of PDMAA hydrogels with different NFC contents via in situ polymerization. The NFC-reinforced intelligent hydrogels exhibited relative harmonious pore sizes in specific sample. From Figure 1a–e, it can be found that NFC contents significantly affected the variation of micropore size. With an increase in NFC contents, the micropore size decreased from about 500 to 100 µm gradually. The reasons can be attributed to the increase in crosslinking density of microstructures, enhancing the compactness of hydrogels. Figure 1 exhibits that the addition of NFC into hydrogel materials maintained the efficient polymerization, proving the feasibility of material composition from point view of microstructure.

#### 3.2 FT-IR spectra and mechanical strength analysis

The FT-IR spectra of intelligent PDMAA hydrogels with various NFC contents are exhibited in Figure 2. The variation of NFC contents maintained the positions of typical bands of hydrogels. The band at 3,546 cm⁻¹ was an –OH stretching vibration peak. The bands at 3,070 and 1,380 cm⁻¹ were –CH₃ stretching vibration peaks. The band at 1,652 cm⁻¹ was the C=O stretching vibration peak. The band at 1,506 cm⁻¹ was a –C=N– stretching vibration peak. Figure 2 indicates that hydrophilic acylamino existed in the prepared PDMAA intelligent hydrogels.
From the point view of functional base, variation of NFC content maintained the original functional chemical bonds of DMAA, which proved the beneficial role of NFC.

Figure 3 shows the stress–strain profiles of intelligent PDMAA hydrogels with various NFC contents. Variation of NFC content significantly affected the mechanical strength characteristics of hydrogels. With the increase in NFC contents, stress and strain of PDMAA hydrogels with 10, 11, 12, 13, and 14 mg/mL NFC increased. The stress average values of intelligent PDMAA hydrogels were 1.85, 2.03, 2.24, 2.41, 2.53, and 2.72 MPa, respectively. The corresponding strain average values of intelligent PDMAA hydrogels were 784%, 795%, 813%, 873%, 915%, and 994%, respectively. The NFC14 intelligent PDMAA hydrogel owned the highest mechanical strength. Figure 1 indicates the increase in crosslinking density of intelligent hydrogels, which increased the stress and strain values. The material composition of PDMAA hydrogel with different NFC contents.
contents exhibited high mechanical property, which provided the mechanical strength base for the application of 3D printing hydrogel actuators.

3.3 Rheology characteristics and 3D printing structure design

To investigate whether the PDMAA hydrogel reaction solution with various NFC contents was appropriate for 3D printing or not, the rheology characteristics of apparent viscosity, storage modulus, and loss modulus were analyzed, as shown in Figure 4. From Figure 4a, it can be found that the hydrogel reaction solution with and without NFC owned shear thinning phenomenon, exhibiting the non-Newtonian fluid property. The addition of NFC significantly increased the viscosity of hydrogel reaction solution, which enhanced the printable property. With the increase in NFC content, the viscosity of hydrogel reaction solution increased, indicating the controllable regulation property of NFC.

Figure 4b exhibits the variation of storage modulus and loss modulus of hydrogel reaction solution along with the increase in NFC contents. When strain values exceeded $10^{-1}\%$, $G'$ and $G''$ of intelligent PDMAA hydrogel reaction solution with various NFC contents decreased and increased, respectively, which exhibited the intersection. The intersection indicated the existence of gel–sol transition point of hydrogels. Before the transition point, the hydrogel was in the gel state. After the transition point, the hydrogel was in the liquid state. The prepared PDMAA hydrogel reaction solution owned high rheology characteristics for 3D printing, which proved the feasibility of regulation role of NFC and provided printable ink for 3D printing of hydrogel actuators. Among the NFC contents, besides the perfect viscosity, storage modulus, and loss modulus, NFC10 hydrogel reaction solution owned the highest crosslinking density. Based on the printable ability and high swelling property, NFC10 was selected as the NFC content for 3D printing of intelligent hydrogel actuators.

Combined with Figures 1–4, it can be found that besides mechanical strength enhancement, addition of NFC realized the printable property of intelligent hydrogels from the point view of rheology characteristics. Therefore, to realize the 3D printing preparation and investigate the intelligent deformation characteristics, the structure models are built in Figure 5. The printing structures including $90^\circ/0^\circ$ and $45^\circ/135^\circ$ configurations and length–width ratios of 3:1, 4:1, and 6:1 were treated as 3D printing parameters. Figure 5a shows the $90^\circ/0^\circ$ configuration model with sample dimensions of $60 \times 10$ mm (length × width). Figure 5b shows the $45^\circ/135^\circ$ configuration model with sample dimensions of $60 \times 10$ mm (length × width). Figure 5c exhibits the $45^\circ/135^\circ$ configuration model with sample dimensions of $30 \times 10$ mm (length × width). Figure 5d shows the $45^\circ/135^\circ$ configuration model with sample dimensions of $60 \times 15$ mm (length × width). Figure 5a and b provides the influence of structure model for the deformation of PDMAA hydrogels. Figure 5c and d provides the influence of length–width ratio for the deformation of intelligent PDMAA hydrogels. All structure models were bilayer structure. Attributed to the inner diameter of 6 mm, the thickness of printed bilayer structure was 1.2 mm.

![Figure 4: Rheology characteristics of (a) apparent viscosity and (b) storage modulus $G'$ and loss modulus $G''$ of intelligent PDMAA hydrogels with various NFC contents.](image)
To realize the effective polymerization of hydrogel reaction solution, the in situ free radical polymerization under vacuum was adopted on the reference of mold preparation of intelligent hydrogels.

Figure 6 shows the effect of structure models including 90°/0° and 45°/135° configurations on deformation characteristics of intelligent PDMAA hydrogels. At the initial state, the original shape of intelligent hydrogel actuator with 60 × 10 mm and 90°/0° configuration was straight, as shown in Figure 6a-0. When the swelling time reached 329 s, the hydrogel actuator exhibited bending deformation along with the length direction in Figure 6a-1. With the increase in swelling time, the bending degree of hydrogel actuators increased, as shown in Figure 6a-2 and a-3. When the swelling time reached 1,206 s, the hydrogel actuator bent to the final circle state in Figure 6a-4. The initial state of hydrogel actuator with 60 × 10 mm and 45°/135° configuration was also straight in Figure 6b-0. The hydrogel actuator exhibited the torsion deformation at 105 s, which was different from Figure 6a-1. With the increase of swelling time, the spiral degree increased, as shown in Figure 6b-2 and b-3. When the swelling time reached 300 s, the final deformation pattern of hydrogel actuator was the spiral hollow cylinder, as shown in Figure 6b-4. Compared with Figure 6a-4 and b-4, it can be found that the 3D printing structure model significantly affected the intelligent deformation patterns. The 90°/0° and 45°/135° configurations exhibited the bending and spiral deformation, respectively, which indicated the high efficient printable ability of hydrogels and feasibility of 3D printing preparation.

Figure 7 shows the effect of length–width ratios of 3:1 and 4:1 on deformation characteristics of intelligent PDMAA hydrogels. Attributed to the same 45°/135° configuration, the intelligent hydrogels with length–width ratios of 3:1 and 4:1 exhibited the spiral deformations. The initial state of hydrogel actuator was straight, as shown in Figure 7a-0. With an increase in swelling time, both sides of hydrogel actuator twisted together, as shown in Figure 7a-1–a-3. When the swelling time reached 120 s, the spiral hollow cylinder is realized in Figure 7a-4. Both sides of the straight hydrogel actuator with relative high length–width ratio spiraled together, as shown in Figure 7b-0 and b-1. With an increase in swelling time, the spiral degree was enhanced, which can be found in Figure 7b-2–b-4. The variation of length–width ratio also significantly affected the deformation patterns. The pitch that represented the intersection of the same helix and the diameter line can be used to describe the spiral deformation difference between Figure 7a-4 and b-4. The pitch of Figure 7a-4 and b-4 was 2.6 and 3.3 cm, respectively. Namely, the hydrogel actuator with larger length–width ratio owned the lower pitch value, which exhibited higher spiral degree.

Combined with Figures 6 and 7, it can be found that the 3D printed bilayer structure built the swelling deformation base of hydrogel actuators. The anisotropic swelling property of layered structure was the intelligent deformation base of 3D printing hydrogel actuators. The addition of NFC into intelligent PDMAA hydrogels realized the 3D printing of hydrogel actuators, which provided selectable intelligent soft materials for the preparation of intelligent soft actuators with high mechanical strength and self-driven deformations.
In this paper, a series of intelligent NFC-reinforced PDMAA hydrogels with high mechanical strength and 3D printable property were prepared. As the reinforcement, NFC changed the crosslinking density and maintained the existence of hydrophilic acylamino, which provided the mechanical strength and swelling function base. With the increase of NFC content, stress and strain values increased. Besides the regulation role of micro-pore size, the addition of NFC significantly changed the rheology characteristics including apparent viscosity, storage modulus, and loss modulus. With the increase in NFC content, the viscosity of hydrogel reaction solutions with non-Newtonian fluid property increased. The existence of the gel–sol transition point exhibited the printable property of hydrogels. Moreover, combined with the swelling rate, 10 mg/mL was selected as the optimal reinforcement content for 3D printing. To investigate the effect of 3D printing parameters including structure model and length–width ratio on the deformation characteristics of hydrogel actuators, the 90°/0° and

4 Conclusion

In this paper, a series of intelligent NFC-reinforced PDMAA hydrogels with high mechanical strength and 3D printable property were prepared. As the reinforcement, NFC changed the crosslinking density and maintained the existence of hydrophilic acylamino, which provided the mechanical strength and swelling function base. With the increase of NFC content, stress and strain values increased. Besides the regulation role of micro-pore size, the addition of NFC significantly changed the rheology characteristics including apparent viscosity, storage modulus, and loss modulus. With the increase in NFC content, the viscosity of hydrogel reaction solutions with non-Newtonian fluid property increased. The existence of the gel–sol transition point exhibited the printable property of hydrogels. Moreover, combined with the swelling rate, 10 mg/mL was selected as the optimal reinforcement content for 3D printing. To investigate the effect of 3D printing parameters including structure model and length–width ratio on the deformation characteristics of hydrogel actuators, the 90°/0° and
$45^\circ/135^\circ$ configurations and 3:1 and 4:1 were designed. Based on the bilayer structure with precise anisotropic swelling property, the printed hydrogel actuators owned swelling intelligent self-driven deformations. The hydrogel actuators with $90^\circ/0^\circ$ and $45^\circ/135^\circ$ configurations exhibited bending and spiral deformations, respectively. The hydrogel actuators with a larger length–width ratio owned the lower pitch value. The self-driven deformation property proved the feasibility of material composition and 3D printing preparation, which provided the selective materials and preparation method candidates for intelligent soft actuators.

**Acknowledgments:** This work was supported by National Key R&D Program of China (2018YFB1105100).

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