Design and Fabrication of Photo-Responsive Hydrogel for the Application of Functional Contact Lens

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Traditional hydrogel has usually had restricted application in the biomedical field due to lack of a control unit. Herein, Azobenzene (AZO), as a typical kind of photo-response molecule, was introduced into hydrogel by polymerization to provide a control unit for hydrogel. Simultaneously, the immobilization of AZO end group would solve the problems of uncontrollable recovery and light fatigue for AZO small molecules according to previous research. In the research, two kinds of crosslinkable AZO molecules, namely AZO crosslinker and AZO monomer, were synthesized as functional monomers or crosslinkers. Further, AZO copolymerized hydrogel and AZO crosslinked hydrogel were obtained. In order to estimate fundamental properties for contact lenses, gelation time and swelling ratio of hydrogels as a function of AZO concentration as well as crosslinking type were investigated. Moreover, synthesized hydrogels exhibited typical porous morphology, but their size and homogeneity were dependent on the type of hydrogel. In order to evaluate photo-responsive performance, detailed photo induced isomerization of both AZO copolymerized hydrogel and crosslinked hydrogel were tracked by UV spectra. The results confirmed the reversible, stable, and controllable photo responsive process. In vitro evaluation was used to investigate drug release behavior using ofloxacin and puerarin as model drugs. It was found that AZO addition as well as hydrogel type would influence puerarin release, but would have little effect on the ofloxacin release behavior in hydrogels. Furthermore, the isomer type in the hydrogel would have some effects on drug release behavior including ofloxacin and puerarin.

Keywords: hydrogel, azobenzene, hydrogel contact lens, drug delivery, isomerization

INTRODUCTION

Since the emergence of poly(hydroxyethyl methacrylate) (pHEMA) hydrogel as soft contact lenses in the 1960s, the swollen soft materials became favorites in the materials field on account of having a similar microenvironment to the biological system (Hao et al., 2011; Hu and Gong, 2016; Hoteling et al., 2018; Samsom et al., 2018; Lorente-Velazquez et al., 2019; Wei et al., 2019). Traditional hydrogel contact lenses were synthesized by some hydrophilic monomers (containing such as hydroxy, carboxyl groups, different from hydrophobic groups like alkyl groups) like HEMA, vinyl pyrrolidone (NVP), acrylamide (AA) with excellent transparency, controllable water content, and certain non-deformability (Hoteling et al., 2018; Samsom et al., 2018; Lorente-Velazquez et al., 2019; Tong et al., 2019; Wei et al., 2019; Guan et al., 2020; Wei et al., 2020). However, these early
synthesized hydrogels had some original drawbacks due to their structure and properties. Typically, an aqueous microenvironment also benefits bacterial reproduction, which would induce eye infection (Hoteling et al., 2018; Samsom et al., 2018; Lorente-Velazquez et al., 2019; Wei et al., 2019). Recently, with the development of regeneration medicine, traditional hydrogels were explored for new applications as drug carriers/hydrogel scaffolds (Hao et al., 2011; Hu and Gong, 2016; Hoteling et al., 2018; Samsom et al., 2018; Lorente-Velazquez et al., 2019; Wei et al., 2019). While these new applications put forward new requests for tradition hydrogels, among these requests, the control unit seems especially relevant for its application.

Generally, the control unit is composed of a stimuli-response structure or material. Until now, the pH-responsive unit, photo-responsive unit, electric-responsive unit, magnetic responsive unit, and chemical responsive unit have been investigated as control units (Dong and Chen, 2011; Guragain et al., 2015; Kim et al., 2015; Liu et al., 2015; Lorenzo et al., 2015; Wang et al., 2015; Bian et al., 2016a; Cao and Wang, 2016; Dong et al., 2017; Xiao et al., 2019). Azobenzene (AZO) has photoisomerizing characteristic, which provides possibilities as a photo switch (Li et al., 2014; Pang et al., 2014). However, uncontrollability and instability of photoisomerization for azobenzene and its small derivatives restricted applications of the chemicals as photo switches. Therefore, many measures have been made to improve the controllability and stability of photoisomerization including fixing one end of molecule to a solid surface, grafting onto a polymer chain, copolymerization with other monomer, and introduction of the long chain to one or both ends of AZO (Deka et al., 2015; Bian et al., 2016a; Bian et al., 2016b; Pang et al., 2018; Gao et al., 2019; Pang et al., 2020; Pang et al., 2020b; Chen et al., 2020). In these contributions, our research mainly focused on the introduction of AZO to polymer by grafting, copolymerization, and AZO initiated ring opening polymerization (ROP) (Chen et al., 2020). Once AZO molecules were combined with polymer, photoisomerization of AZO domain could be adjusted by either the external environment of macromolecules including solvent, temperature, light density, light source, or internal structure including properties of copolymer monomer and chain length of polymer (Pang et al., 2018; Gao et al., 2019; Pang et al., 2020b). Furthermore, AZO based macromolecule could be endowed with biocompatibility according to the properties of the main chain (Pang et al., 2018). When it comes to application of photo switches, material is an indispensable carrier. Therefore, the AZO based molecules have been fabricated into nanoparticles, films, hydrogels, etc.

It was inferred from existing research that time-dependent restricted motion of polymers provides better controllability of AZO domain isomerization transformation and stability of AZO isomers, especially for cis-isomers. Besides time, spatial conditions were also inferred to be optimal elements for improving controllability and stability of photoisomerization. Therefore, the AZO unit was introduced to the traditional hydrogel network by copolymerization in the research. In one aspect, the AZO unit was restricted in a mesh of hydrogel network, which would improve the controllability and stability of AZO photoisomerization. In other aspects, AZO photoisomerization would bring the deformation of mesh in the hydrogel network, which would endow photo-response properties to tradition hydrogel. In order to realize the idea, AZO crosslinked hydrogel and AZO copolymeried hydrogel were synthesized and investigated elaborately in the research. Finally, the function of drug delivery was evaluated to explore new application of tradition hydrogel in the work.

**Experiment**

**MATERIALS**

Acryloyl chloride, p-aminoazobenzene (AZO), and azodianiline (DAZO) was purchased from Aladdin. Hydroxyethyl methacrylate (HEMA), N-Vinyl pyrrolidone (NVP) and puerarin were obtained from Shanghai Jingchun Industries Co. Ltd., China and distilled under vacuum before use. Dichloromethane (DCM), diethyl ether, tetrahydrofuran (THF), dioxane, triethylamine (TEA) and dimethylsulfone (DMSO), ammonium persulfate (APS), and N,N,N',N'-tetramethylthlenediamine (TEMED) were obtained from Shanghai Chemical Industries Co. Ltd., (China). Hyaluronic acid and fluorescein isothiocyanate were obtained from Sigma. Norfloxacain and puerarin were purchased from Jinling pharmac Industries Co. Ltd., China. All other reagents and solvents were of analytical grade and used as received.

**Synthesis of AZO Monomer and AZO Crosslinker**

AZO monomer and AZO crosslinker were synthesized by acyclorination (Gao et al., 2019). AZO monomer was synthesized according to the previous method. Briefly, 10 mM AZO was reacted with 11 mmol acryloyl chloride in 20 ml anhydrous DCM at stirred state at room temperature for 4–5 h, using TEA as an alkali absorber. After reaction, AZO monomer was obtained by the water washing purification procedure and reduced pressure distillation. AZO crosslinker was synthesized using a similar method. Briefly, 10 mM DAZO was reacted with 33 mmol acryloyl chloride in 20 ml DMF at stirred state at room temperature for 20 h, using TEA as an alkali absorber. Differently, the obtained product was collected by precipitation with water (pH4) and washed with saturated NaHCO3 solution and water. The final product was obtained by freeze-drying and characterized by 1H nuclear magnetic resonance (1H NMR, Bruker, AV300).

**Synthesis and Characterization of Hydrogels**

AZO copolymerized hydrogel and AZO crosslinked hydrogel were fabricated by the radical polymerization. Briefly, AZO monomer/AZO crosslinker with various ratios dissolved into 5 ml HEMA and 1.162 ml NVP by stirring, into which certain amounts of APS and TEMED with equal molar ratio were respectively added with final initiator concentration at 0.5 mol
%. The above mixture was injected into a circle model, and then polymerized at 60°C in oven. Gelation time was obtained by judgment of its fluidity. In order to ensure full crosslinking, 1 h polymerization time was used in the hydrogel product.

Next, the fundamental properties of the hydrogels like swelling ratio and hydrogel morphology as contact lenses were characterized. Hydrogels were dried and weighed \( W_0 \). The dried hydrogels had been swollen to equilibrium in water and weighed \( W_1 \). The equilibrium swelling ratio of the hydrogels was defined as \( \frac{W_1}{W_0} \times 100\% \). The morphology of the hydrogels was characterized by scanning electron microscope (SEM, S-8100).

**Responsive and Recovery Performance of Hydrogels**

Hydrogels were tracked by UV spectroscopy (Cary 50). UV lamp (10 W) was used as a photo source to induce trans-to-cis transition of AZO domain in hydrogels. After UV irradiation, white light was used to induce cis-to-trans recovery at different conditions. Real-time UV spectra as a function of irradiation time and recovery were recorded to track the structural change of molecules. Repeated irradiation and recovery methods were used to investigate fatigue resistance of AZO domain in hydrogels. Moreover, factors of temperature and light density were applied to clarified photo-isomerization for AZO domain in hydrogel.

**In vitro Drug Delivery Evaluation**

Orfloxacin and puerarin were used as model drugs to be loaded into hydrogel by copolymerization in situ. Briefly, orfloxacin and puerarin was dissolved in monomers of HEMA and NVP with concentration of 3 mg/ml. Then hydrogel with drugs was formed using the above-mentioned method.

The above-prepared hydrogels were submerged in 4 ml PBS in a 5 ml centrifugal tube, which was placed under shake cultivation at specific temperature. At regular intervals, 2 ml of released solution were moved for detection. Simultaneously, 2 ml of fresh PBS were supplemented into the released system. At given times, the absorbance of removed drug solution at 280 nm for orfloxacin and at 355 nm for puerarin was recorded by UV spectroscopy. The drug concentration was obtained by dilution ratio and a calibration curve, which was constructed from known concentrations of drug solutions. The cumulative released amount was then calculated according to drug concentration and volume. UV light was used to excite isomerization of AZO crosslinked hydrogel in the process of drug release. After excitation, drug release was performed in a dark environment at 20°C.

**Statistical Analysis**

Data were analyzed using the t-test for differences. Results are reported as means ± standard deviation. The significant level was set at \( p < 0.05 \).

**RESULTS AND DISCUSSION**

**Fabrication and Characterization of Photo-Responsive Hydrogel**

The AZO crosslinker was synthesized by acylchlorination, which was characterized by \( ^1 \)H NMR spectrum in Figure 1. In detail, chemical shifts from 5.8 to 6.5 ppm were attributed to hydrogens of vinyl groups at a, b, and c positions, chemical shifts from 7.1 to 7.9 ppm are attributed to Hs of benzene ring, and chemical shifts at 10.5 ppm are attributed to H of imine groups. Besides qualitative analysis, quantitative calculation was applied to confirm the precise chemical structure of the AZO crosslinker. The integration of peak from 5.8 to 6.5 ppm was 3.0, the integration of peak from 7.1 to 7.9 ppm was 4.0, the integration of peak at 10.5 ppm was 1.0. These ratios were consistent with H numbers of groups, which are listed as 6 Hs for vinyl, 8 Hs for benzene ring, 2 Hs for imine. These results indicated that AZO crosslinker with two vinyls was successfully synthesized without any impurity.

In order to endow hydrogel photo-response property, two kinds of hydrogel were synthesized by radical polymerization, namely AZO copolymerized hydrogel and AZO crosslinked hydrogel as shown in Figures 2A,B. Formed hydrogels exhibited yellow transparent state (Figures 2A,B). Since traditional hydrogel was colorless, yellow color could demonstrate the successful synthesis of AZO copolymerized hydrogel and AZO crosslinked hydrogel from a side. In polymerization, AZO domain introduction would influence the gelation time and swelling ratio (Figures 2C,D). For AZO copolymerized hydrogel, the gelation time was prolonged largely by the increased AZO monomer (Figure 2C). Even the hydrogel could not be gelled when the AZO monomer concentration exceeded 4%. For AZO crosslinked hydrogel, the gelation time was also prolonged by the increased AZO crosslinker (Figure 2C). Similarly, the hydrogel could hardly be gelled when AZO crosslinker concentration exceeded 4%. Since gelation of pHMA kind hydrogel depended on the growth of branch and mutual crosslinking between branches, the steric hindrance of the AZO structure would hinder branch formation in polymerization. Thus, the gelation time would increase along with the AZO domain increasing. Furthermore, gelation time of AZO crosslinked hydrogel was shorter than that of AZO copolymerized hydrogel with significant difference on account of the larger double carbon bond density (Figure 2C). Differently, the swelling ratio of both AZO crosslinked hydrogel and AZO copolymerized hydrogel decreased with the increase of AZO domain in hydrogel (Figure 2D). When the AZO domain reached 4%, the swelling ratio of AZO crosslinked hydrogel was significantly smaller than that of AZO copolymerized hydrogel (Figure 2D). Since the AZO domain was not a hydration unit, the hydrogel with the AZO domain decreased sight with the increase of AZO domain. Moreover, crosslinker density was another factor to influence the swelling ratio of hydrogels. Therefore, the AZO crosslinked hydrogel possessed a smaller swelling ratio due to larger crosslinker density.
FIGURE 1 | $^1$H NMR spectrum of AZO crosslinker.

FIGURE 2 | Schematic formation of AZO crosslinked hydrogel (A) and AZO copolymerized hydrogel (B). (C) Gelation time of hydrogel as a function of AZO concentration. (D) Swelling ratio of hydrogel as a function of AZO concentration. *$p < 0.05$, **$p < 0.01$.

FIGURE 3 | SEM images of traditional hydrogel (A), AZO crosslinked hydrogel (B) and AZO copolymerized hydrogel (C). The scale bar is 200 µm.
Moreover, hydrogel morphology was characterized by SEM images to clarify hydrogel inner structure, which was shown in Figure 3. Hydrogels exhibited a typical porous structure, but porous size was different for different hydrogels. Namely, the traditional hydrogel possessed the largest pore size and an asymmetrical structure. The AZO crosslinked hydrogel exhibited the smallest pore size and a homogeneous structure. The AZO copolymerized hydrogel had a medium pore size and a relative homogeneous structure. The hydrophilic and soft chain nature of traditional hydrogel would collapse during freeze-drying, which resulted in asymmetrical structure. However, AZO domain introduction would increase the rigidity of the polymer chain, which prevented the hydrogel from collapsing during freeze-drying. For the AZO crosslinked hydrogel, larger crosslink density resulted in smaller mesh size in the hydrogel. These results were consistent with the above-mentioned discussion.

**Photo Responsive Performance of Hydrogel**

As a photo-responsive unit, the photo-response of hydrogel was evaluated by UV spectrum using UV light as a stimulus since the trans isomer and cis isomer of the AZO unit had different UV spectra from the verified theory and previous research (Pang et al., 2018; Gao et al., 2019; Pang et al., 2020a; Pang et al., 2020b).

UV spectra of AZO copolymerized hydrogel as a function of irradiation time and recovery time were shown in Figures 4A,B. Originally, a typical absorption peak at 350 nm belonging to $\pi-\pi^*$ transition and a small flat absorption peak at 430 nm belonging to $n-\pi^*$ transition exhibited the typical characteristic of trans isomer of AZO unit in Figure 4A, which also confirmed the successful synthesis of AZO copolymerized hydrogel from the other side. After UV irradiation, the maximum absorbance at 350 nm decreased greatly and shifted to 330 nm, simultaneously the absorbance at 430 nm increased a little with irradiation time until 90 s, which indicated that the trans isomer had been transferred to the cis isomer (Figure 4A). After removing light, the recovery process from cis isomer to trans isomer could be accelerated by white light irradiation according to previous research (Pang et al., 2018; Gao et al., 2019; Pang et al., 2020a; Pang et al., 2020b). In order to track the recovery process in a convenient and observable period, white light irradiation was used to induce the recovery process from cis to trans in Figure 4B. Upon white light irradiation, absorbances at 330 nm were gradually shifted to 350 nm and recovered to their respective origin value within 30 min, absorbance at 430 nm was also recovered to its origin value. These phenomena confirmed the reversible recovery of the trans structure (Figure 4B). Since absorbance at 350 nm/330 nm directly reflected the trans isomer content and response time reflected the efficiency of the light control unit, the two parameters were used to evaluate fatigue resistance of copolymer repeated UV/white light irradiation, which was important for the photo control unit in Figures 4C,D. From the recorded real-time absorbance at 350 nm/330 nm of AZO copolymerized hydrogel in Figure 4C, maximum absorbance at 350 nm, either initially or after recovery, was stabilized around 0.9 regardless of circle time, and minimum absorbance at 330 nm after UV irradiation was stabilized at 0.4–0.5 regardless of circle time. From the tracked response time in Figure 4D, irradiation response time was stable.
at 90 s and recovery response time was stable at 30 min regardless of cycle time. These results confirmed that UV could induce reversible isomerization for the AZO copolymerized hydrogel, and further the transition could be realized stably by specific light excitation source without any sign of fatigue or photobleaching.

Besides the AZO copolymerized hydrogel, the reversible photo triggered isomerization of AZO crosslinked hydrogel was tracked by UV spectra in Figure 5. In its natural state, a typical absorption peak at 373 nm existed in the UV spectrum of AZO crosslinked hydrogel in Figure 5A, indicating the trans isomer of the AZO unit in hydrogel. After UV irradiation, the maximum absorbance decreased greatly until 150 s, which indicated that the trans isomer had been transferred to the cis isomer completely (Figure 5A). Upon white light irradiation, absorbances at 373 nm recovered to their respective origin value within 30 min, which confirmed the reversible recovery of the trans structure (Figure 5B). Moreover, maximum absorbance after white light irradiation was stabilized around 1.1 and minimum absorbance after UV irradiation was stabilized around 0.4 regardless of cycle number (Figure 5C). Simultaneously, irradiation response time was stable at 150 s and recovery response time was stable at 30 min regardless of circle time (Figure 5D). These results revealed that AZO crosslinked hydrogel possessed reversible and stable photo-responsive performance without photobleaching. Theoretically, the AZO domains existed in two ground states (trans and cis form) and one excited state. Transition between isomers must go through the excited state, although the excited state nearly could not be detected. Therefore, light excitation in fact provided energy between the ground state and excited state. Besides the energy factor, steric hindrance was also an important factor to influence isomerization transition from previous research (Pang et al., 2018; Gao et al., 2019; Pang et al., 2020a; Pang et al., 2020b). Although the transition energy of the AZO unit from trans to cis in the AZO crosslinked hydrogel is similar to that in the AZO copolymerized hydrogel, the AZO unit in the AZO crosslinked hydrogel with two fixed ends had a larger steric hindrance in the process of trans-to-cis transition than that of AZO copolymerized hydrogel with one fixed end. Therefore, in the same UV irradiated intensity, trans-to-cis transition of the AZO unit in the AZO crosslinked hydrogel needed a longer time (150 s) to be completed than that of the AZO copolymerized hydrogel (30 s).

Since the recovery process is an essential factor to realize the controllable and stable photo response properties of hydrogel, it was investigated considering the influencing factors of light intensity and temperature in Figure 6. For the AZO copolymerized hydrogel, recovery time decreased greatly with the increase of light intensity, especially when light intensity was lower than 500 LUX (Figure 6A). Likewise, the recovery time of the AZO crosslinked hydrogel as a function of light intensity exhibited similar trends, except the trend became sharper than that of the AZO copolymerized hydrogel (Figure 6B). Additionally, the cis structure in both AZO copolymerized hydrogel and AZO crosslinked hydrogel could not be recovered in the dark during the observed 3 days at 20°C, which was not listed in Figures 6A,B. Besides light intensity, effects of temperature were also presented in Figures 6C,D in consideration of two different light intensities (500 LUX and
1,000 LUX). For the AZO copolymerized hydrogel, recovery time decreases gradually with the increase of temperature either under 500 LUX white light irradiation or 1,000 LUX white light irradiation (Figure 6C). Likewise, recovery time of AZO crosslinked hydrogel as a function of temperature exhibited similar trends with a sharper trend (Figure 6D). From the above discussion, the cis-to-trans recovery process should overcome the barrier between the cis ground state and excited state. While heat/white light could provide needed energy from the cis structure ground state to excited state. Therefore, it was easy to understand that higher temperature and light intensity resulted in shorter recovery response time.

In summary, both exhibited controllable and stable photo responsive performance, which could be adjusted by environmental factors.

**In vitro Drug Delivery Evaluation**

Since one important application of hydrogel contact lenses was as an ophthalmic drug carrier, performances of hydrogels as a kind of drug carrier were evaluated in vitro. Model drugs were incorporated into hydrogels by in situ polymerization. The cumulative drug release in different hydrogels is shown in
Figure 7. From Figure 7A, 70% ofloxacin was rapidly released from any of hydrogels at initial 30 min, and then another 20% ofloxacin was gradually released from any of hydrogels in the following 16–17 h. Further, the difference either between traditional hydrogel and AZO functional hydrogel or between AZO copolymerized hydrogel and AZO crosslinked hydrogel was not significant. It seemed the slight addition of the AZO unit in the hydrogel network had little effect on ofloxacin release behavior. When it came to puerarin, the condition was different in Figure 7B. Initially, 30% puerarin was rapidly released from any of hydrogels during the first 30 min. Next, another 40% puerarin was relatively rapidly released from the traditional hydrogel, another 30% puerarin was relatively rapidly released from the AZO copolymerized hydrogel, and another 20% puerarin was relatively rapidly released from the AZO crosslinked hydrogel in the following 4–5 h. Finally, an additional 10% puerarin was relatively slowly released from any of hydrogels. The delayed puerarin release in AZO containing hydrogel might be ascribed to the interaction between puerarin molecules and the AZO unit. The AZO crosslinked hydrogel possessed a more compact structure due to higher double carbon density, which resulted in a slower puerarin release rate. Furthermore, the AZO crosslinked hydrogel was chosen as a carrier for drug release behavior investigation since isomerization of the AZO crosslinker in the hydrogel would cause deformation of the hydrogel network. Before the beginning of drug release, one group of hydrogels was irradiated by UV to realize trans-to-cis transition. After irradiation, drug release was performed in a dark environment at room temperature in order to maintain the cis structure of the AZO unit in hydrogel since the cis isomer could not be recovered to the trans isomer within 3 days from the above-mentioned discussion. The drug release behavior of two groups of AZO crosslinked hydrogels with or without irradiation were shown in Figures 8A,B. Without irradiation, the cumulative ofloxacin release behavior at room temperature in Figure 8A was similar to that at 37°C in Figure 7A. But after irradiation, the ofloxacin release speed became slower and the cumulative ofloxacin release amount became obviously lower (Figure 8B). It was inferred from these results that trans-to-cis isomerization would make the crosslinked polymer network shrunken as Figure 8C showed, which resulted in a tighter mesh of the hydrogel network. Therefore, the cis-state of the AZO unit in the AZO crosslinked hydrogel delayed drug release.

CONCLUSION

Divinyl functional AZO, namely AZO crosslinker, was successfully synthesized without any impurities, just as the previously synthesized vinyl functional AZO (AZO monomer). Based on traditional hydrogel contact lenses, AZO functional hydrogels, namely AZO copolymerized hydrogel and AZO crosslinked hydrogel, were successfully copolymerized with an adjustable AZO unit (1–4% W/V). Both the formed AZO copolymerized hydrogel and AZO crosslinked hydrogel exhibited a yellow transparent state. In polymerization, the gelation time was prolonged largely by the increased AZO unit for both hydrogels, but the effects of AZO were more significant for AZO crosslinked hydrogel than for AZO copolymerized hydrogel, especially when AZO concentration was higher than 2%. The swelling ratio of both hydrogels decreased slightly with the increase of the AZO unit. Especially when the AZO domain reached 4%, AZO crosslinked hydrogel possessed a significantly lower swelling ratio than the AZO copolymerized hydrogel. Synthesized hydrogels exhibited typical porous morphology, but their size and homogeneity were dependent on the type of hydrogel. UV could induce reversible isomerization of the AZO unit in both the AZO copolymerized hydrogel and crosslinked hydrogel, and the transition could further be realized stably by a specific light excitation source without any sign of fatigue or photobleaching. For the cis-to-trans process, high temperature and light intensity would accelerate transition speed. In summary, both exhibited controllable and stable photo responsive performance, which could be adjusted by environmental factors. In the in vitro evaluation, it was found that the slight addition of the AZO unit in the hydrogel network had little effect on the ofloxacin release behavior, but AZO unit would slow and delay puerarin release in hydrogel. Additionally, the more compact structure of the AZO crosslinked hydrogel could
cause puerarin release more slowly. Furthermore, the cis state in the AZO crosslinked hydrogel would delay drug release either for orfloxacin and puerarin, which confirmed that the crosslinked polymer network would shrink in the cis state of AZO in the AZO crosslinked hydrogel.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/supplementary material, further inquiries can be directed to the corresponding author.

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AUTHOR CONTRIBUTIONS

XL prepared the hydrogel HZ investigated the properties of materials YQ evaluated the drug release properties XH give the whole idea.

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