A novel in situ gel base of deacetylase gellan gum for sustained ophthalmic drug delivery of ketotifen: in vitro and in vivo evaluation

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Abstract: In this study, an ion-activated ketotifen ophthalmic delivery system was developed by using a natural polysaccharide, deacetylase gellan gum. Its rheological characteristics, stability, in vitro gelation, release in vitro, and pharmacodynamic activity in vivo were investigated. The formulation had an optimum viscosity that will allow easy drop as a liquid, which then underwent a rapid sol–gel transition due to ionic interaction. There were negligible alterations in the initial values of viscosity of the formulations over a storage period of 180 days. The in vitro release profiles indicated that the release of ketotifen from in situ gels exhibited a sustained feature. Scintigraphic studies indicated that deacetylase gellan gum could increase the residence time of the formulation. At the same dose, in situ gels demonstrated a typical sustained and prolonged drug-effects behavior compared with the common drops.

Keywords: ophthalmic delivery, deacetylase gellan gum, sol–gel transition, scintigraphic studies, pharmacodynamic

Introduction
In conventional ophthalmic delivery system, liquor solution was widely used because of the simple preparation process, accurate dosage, easiness to use, and high medication compliance among patients.1 However, due to the tear secretion and blinking, liquor solution had very short residence time (~5 minutes) on the surface of the cornea. Once the liquid entered the corneal capsule, it was immediately diluted or caused corneal erosion and increased the need for frequent drug times to improve efficacy.2 Although some bio-adhesive formulations such as ocular gel were able to extend the residence time of the drug in the conjunctiva of the eye, they had higher viscosity, inaccurate dosage, and higher risk of adverse reactions. Meanwhile, they may severely limit the development of the dosage form.3,4 Therefore, it is important to improve the efficacy of ophthalmic preparations by selecting suitable polymeric materials as a thickener and for an anti-tear clearance of drugs.

Ketotifen fumarate (KF) is a drug for histamine H1 receptor antagonist and mast cells. It can inhibit type I allergic or immediate-type allergy-related cells (mast cells, eosinophils, basophils, and neutrophil cells). The release of inflammatory mediators can also inhibit the chemotaxis, activation, and degranulation of eosinophils.5 Clinically, KF ophthalmic solution was mainly used for the treatment of seasonal allergic conjunctivitis following a dosage of one to two drops per time and four times a day (during the morning, noon, night, and before sleep). However, the discontinuity of drug use by many clinical patients often affected the final effects.6,7 Therefore, our
group intended to develop a suitable ophthalmic preparation to improve the efficacy of KF.

In situ gel is a polymer solution that changes in accordance with temperature of the applied place, ion strength, or pH and forms a viscoelastic semisolid gel system. The formation mechanism is to use polymer materials in response to external stimuli so that the polymer undergoes reversible changes in the dispersion state or conformation under physiological conditions, and then completes the transformation from solution to gel. During the preparation, gelation phenomenon occurs under appropriate physiological conditions and at the medication area, leading to extended residence time of the drug and improved bioavailability. Combining advantages of liquid and gel, the gel has a wide-ranging application prospect in drug delivery field, for instance, in ophthalmic administration, oral administration, implants, and nasal system. In recent years, the in situ gel delivery system has attracted great attention from scholars and become one of the focus areas of pharmacy.

In situ gel can be divided into three types according to its phase change performance: temperature sensitive, pH sensitive, and ionic strength sensitive. Temperature-sensitive materials mainly include block copolymer and poloxamer. pH-sensitive materials include cellulose acetate phthalate and acrylic acid polymers, which, by changing the pH value of the environment, can facilitate phase transition. Among ion-sensitive materials, sodium alginate is the most commonly used. But in recent years, deacetylase gellan gum (DGG) starts to be considered more frequently.

DGG is an extracellular polysaccharide, produced from *Pseudomonas elodea*, a Gram-negative bacterium separated from water lily. It has a parallel double-helix structure, and the whole gel chain is formed by four basic units via repeat polymerization; when formulated into solution with a certain concentration, it exhibits the feature of cation-induced gelation. Because of this unique nature, it has been used for oral, ocular, and nasal delivery system. But there is no report about KF in ophthalmic drug delivery systems.

In this study, an ion-activated ketotifen ophthalmic delivery system was developed by using a natural polysaccharide, DGG. Its rheological characteristics, stability, in vitro gelation, release in vitro, and pharmacodynamic activity in vivo were investigated.

**Materials and methods**

**Materials**

KF was given by Xingyin Chemical Ltd. (Hubei, People’s Republic of China; patch number: 10209). KF eye drops (0.5 mg/mL) was purchased from Jiminkexin Group Co., Ltd. (Wuxi, People’s Republic of China; patch number: 090515). DGG was obtained from Kelco Co., Ltd. (Atlanta, Georgia) (molecular weight 500 kDa, 95% deacetylation degree; USA; patch number: 20050929). Egg albumin (EA) was purchased from Sigma Aldrich Co., Ltd (Shanghai, People’s Republic of China). Artificial tears (AT) including NaCl 6.78 g, CaCl$_2$·2H$_2$O 0.084 g, KCl 1.38 g, and NaHCO$_3$ 2.18 g in 1,000 mL of purified water was prepared according to previous report. Purified water from Milli-Q system (Millipore, Bedford, MA, USA) was used throughout the experiment. All other reagents were of commercially analytical grade.

**Preparation of in situ gel**

Briefly, a certain amount of DGG (0.25%, 0.6%, 1.25%; w/v) was added to deionized water and dissolved by heating to 60°C with moderate stirring. Once completely dissolved, the solution was cooled to a temperature below 30°C. Then, KF (0.05%, w/v), chlorhexidine acetate (0.01%, w/v), and glycerol (2%, w/v) were slowly added to the system and mixed well for at least 30 minutes. The pH of the formulation was between 6.0 and 8.0. In this formulation, DGG served as a gel base, and chlorhexidine acetate, a preservative.

**In vitro gelation and rheological characteristics**

The three types of in situ gels prepared were mixed with AT at the ratio of 25:7, and the temperature was maintained at 34°C±0.5°C, simulating a real environment in vivo. Then, the viscosity of the different DGG formulations, either in solution or in gel made with AT, was determined with a rotational viscometer using a proper sample. Measurements were performed using suitable spindle number at different rotation rates. The viscosity was read directly from the viscometer display. All measurements were made in triplicate.
HPLC analysis
High-performance liquid chromatography (HPLC)-variable wavelength detector was used in the determination of KF in vitro samples, and specific chromatographic conditions were as follows: column: Kromasil C18 column (150 mm × 4.6 mm, 5 μm); mobile phase: methanol and water (1,000 mL, 65:35, V/V), where triethylamine of 0.35 mL was added; column temperature: 30°C; flow rate: 1 mL/min; detection wavelength: 300 nm; and injection volume: 20 μL.

Stability studies
The proposal of stability study was modified mainly based on the guiding principles of the Chinese Pharmacopoeia. KF in situ gel (0.6% DGG) was placed in three stable boxes kept at 4°C, room temperature, and 40°C, and saturated sodium chloride solution was added and maintained at a relative humidity of 75%±5%. Then, at 0 day, 30 days, 60 days, 90 days, and 180 days of the test, they were observed to find if the gel content, appearance, gelation, and viscosity were changed or not.

In vitro release behavior
Preparation of release media
In order to maintain the biological activity of cornea in vitro, this study used glutathione media buffer for release. This buffer consisted of two parts: 1) NaCl 12.4 g/L, KCl 0.716 g/L, NaH₂PO₄·H₂O 0.206 g/L, and NaHCO₃ 4.908 g/L; and 2) CaCl₂·2H₂O 0.23 g/L, MgCl₂·6H₂O 0.318 g/L, glucose 1.8 g/L, and glutathione (oxidized form) 0.184 g/L.²³ Solutions of the two parts are placed at 4°C in a refrigerator and equally mixed before use.

Preparation of in vitro cornea
Air was injected into the ear vein of a number of New Zealand white rabbits, causing their death. Thirty minutes after their death, their corneas were separated using ophthalmic surgical blades and tweezers. The sclera, iris, ciliary body, and other excess tissues were removed and washed with distilled water for several times, and water attached to the stand was sucked using filter paper.

In vitro drug release
The studies on the in vitro release of KF from in situ gels were carried out using the Franz diffusion cell method. The stirring speed was set at 50 rpm, and the temperature was maintained at 37°C±0.5°C. Aliquots of 200 μL of KF in situ gels (or 200 μL of KF eye drops) were directly placed into cornea and then mixed with 56 μL of AT. Samples of 0.2 mL were extracted from the tubes at 0.25 hour, 0.5 hour, 1 hour, 2 hours, 4 hours, 6 hours, 8 hours, 12 hours, and 24 hours after centrifugation at 4,000 rpm for 5 minutes. Then, the KF was analyzed by HPLC. The accumulative release was calculated, and the results were shown as mean ± SD (n=3).

Scintigraphic studies
The ocular mucosal residence time of in situ gels was investigated using a modified scintigraphic method.²⁴ Briefly, 0.6% DGG KF in situ gels (prepared as described earlier) was incorporated with 99mTc-diethylene triamine pentaacetic acid (DTPA). A solution of 99mTc-DTPA dissolved in phosphate-buffered saline (PBS) was used as control. Then, six rabbits were divided into two groups, each receiving gels and solution formulations in a crossover design with at least 1 week of washout period. The rabbit was positioned 10 cm in front of the probe and 100 μL of the radiolabeled gels or solutions, which had been stored at 20°C for 30 minutes before use, was instilled into the ocular mucosal surface. Single-photon emission computing tomography (ZLC 3700, Münich, Germany) was autotuned to detect the 140 keV radioactivity of 99mTc-DTPA. Recording started 2 seconds after administration and continued for 60 minutes using a 156×156 pixel matrix. A total of 84 frames of dynamic images were recorded and analyzed by medical system ICONP workstation (Siemens, Münich, Germany).

Pharmacodynamic activity
Immunization and challenge
All rats were injected intraperitoneally with 100 μg of EA plus 20 mg of alum in 1 mL of PBS (pH 7.4) for immunization. Fourteen days later, rats were challenged topically with 10 μL of 10% EA in PBS. To obtain a remarkable ocular reaction, 20 μL of 1 mol/L dl-dithiothreitol and a mucolytic agent in PBS (freshly prepared before use) were applied to each eye 15 minutes before topical challenge as described by Calonge et al.²⁵ Evans Blue (EB; 2 mg/100 g) in 1 mL of PBS was intravenously injected immediately prior to challenge. Animals were anesthetized with intraperitoneal 5% chloral hydrate (approximately 0.2 mL/100 g).

Drug administration
Animals were divided into two groups (gels and solutions), 30 in each. One eye of each animal received one drop of relevant drug (0.1% KF formulation), and the fellow eye was untreated as control. All rats were killed by exsanguination 1 hour after challenge. Globes and adnexa were taken at 5 minutes, 10 minutes, 15 minutes, 30 minutes, 45 minutes,
and 60 minutes after ocular administration. Then, the tissues were exenterated, weighed, immersed in an extracting solution of 5 mL of sodium sulfate (0.5%) plus acetone (3:7, v/v), shaken strongly, and kept at room temperature. Twenty-four hours later, solutions were centrifuged at 1,000 rpm for 10 minutes. The color intensity of supernatant was detected spectrophotometrically at 620 nm.

Standard curve was generated to transform absorbance unit in microgram of EB per milliliter of solution, and EB extravasation in ocular tissues was calculated. Drug efficacy was expressed as percent inhibition of the reaction on the vehicle-treated eye. In the equation, IR (%) = (UCE – DTE)/UCE, where IR is the drug inhibitory rate, DTE is EB content in drug-treated eye, and UCE, in the untreated control eye.

A $P$-value of $\leq 0.05$ was considered significant.

**Statistical analysis**

Results were expressed as the mean value ± standard deviation. Statistical analysis was evaluated by two-tailed Student’s $t$-test or by two-way analysis of variance followed by Dunnett’s $t$-test, with a level of significance chosen at $P<0.05$.

**Results and discussion**

**In vitro gelation**

To gain a better understanding of the existence of gel, hydrogel formation in vitro was observed. The two main prerequisites of in situ gelling systems are optimum viscosity and gelling capacity. The formulation should have an optimum viscosity that will allow easy drop as a liquid, which then undergoes a rapid sol–gel transition due to ionic interaction. Figure 1 shows the prepared gels formed under the conditions of AT. Because of the microscale of ions required for gel formation, rapid gelation in vivo can be expected. In the selection of the concentration of the gelling polymer, a compromise is sought between a satisfactory gel strength for use as a delivery vehicle and an acceptable viscosity for easiness of spraying. As shown in Figure 2, three different DGG formulations demonstrated pseudoplastic behavior either in solution or in gel. The viscosity of the tested gels grew with the increase in DGG concentrations, and a great viscosity change was found when DGG underwent sol–gel transition at lower concentrations (0.25% and 0.6%). By contrast, with a relatively high original viscous solution obtained with 1.25% DGG, viscosity change observed after gel formation was slight. It was proposed that as the concentration of DGG increased, the polymer chains approached closer and the number of interactions between the polymer chains increased, leading to a denser three-dimensional network structure.

**Stability studies**

There were negligible alterations in the initial values of viscosity of the formulations over a storage period of 180 days. The samples were also analyzed for drug content by the HPLC method. Again, the drug degraded to a negligible extent, and the percentage of drug degradation is $5\%$. Many factors affected the stability of a pharmaceutical product, including the stability of the active ingredient(s), and the potential interaction between active and inactive ingredients. To calculate the shelf life of the formulation, extensive stability data are collected according to the International Conference on Harmonization guidelines. Because the overall degradation is $<5\%$, a tentative shelf life of 2 years may be assigned to the optimized formulation.

**In vitro release**

In vitro release of KF from different formulations was studied using Franz diffusion cell method at 37°C. The plot of drug release percentage versus time was drawn to exhibit the release profile (Figure 3). As described in Figure 3, $>50\%$ of KF was dissolved in release medium from KF eye drop.

![Figure 1](image1.png) **Figure 1** In vitro hydrogel formation with in situ gels (0.6% DGG, w/v) and artificial tears (25:7, v/v).

**Abbreviation:** DGG, deacetylase gellan gum.
within 30 minutes, and nearly 100% of drug was released after 2 hours. By contrast, KF in situ gels presented a sustained-release manner in glutathione media buffer. After incubated for 24 hours, approximately 78.4%, 66.2%, and 57% KF release from 0.25%, 0.6%, and 1.25% in situ gels were found, respectively. The in vitro release profiles indicated that KF was released from in situ gels in a sustained fashion. The phenomenon could be related to the fact that DGG underwent a rapid sol–gel transition when exposed to AT, which was also confirmed by the viscosity experiment. During the hydrogel formation, a portion of KF might be loaded into the hydrogel phase to slow down drug release. In addition, the release rate and DGG concentration were shown to be negatively correlated. At fixed drug concentrations, the higher the DGG concentration was, the lower the rate of drug release was. The release from various DGG formulations could be ranked as follows: 0.25%>0.6%>1.25%. The in vitro release data were kinetically analyzed according to zero-order, first-order, and the diffusion-controlled release mechanism. The relatively high correlation coefficient values obtained from the analysis of the amount of drug released versus the square root of time indicated that the release followed the Higuchi kinetic model (r=0.9993–0.9997). Overall, opposite to the fast drug release in control group, KF in situ gels performed a controlled-release character which was beneficial to persistent ocular therapy.

**Scintigraphic studies**

Gamma scintigraphy is a technique whereby the transit of a dosage form through its intended site of delivery can be
Table 1 Inhibition rates of the drugs on Evans Blue leakage in eyes (n=10)

<table>
<thead>
<tr>
<th>Inhibition rates of the drugs (%)</th>
<th>5 minutes</th>
<th>10 minutes</th>
<th>15 minutes</th>
<th>30 minutes</th>
<th>45 minutes</th>
<th>60 minutes</th>
</tr>
</thead>
<tbody>
<tr>
<td>KF eye drops</td>
<td>49.09±11.32</td>
<td>52.31±15.21</td>
<td>43.33±13.26</td>
<td>31.82±11.09</td>
<td>20.38±10.12</td>
<td>13.85±5.43</td>
</tr>
<tr>
<td>KF in situ gels</td>
<td>48.33±12.56</td>
<td>50.91±13.73</td>
<td>48.46±15.31</td>
<td>48.00±12.89*</td>
<td>42.37±13.29*</td>
<td>41.80±14.19*</td>
</tr>
</tbody>
</table>

Note: *P<0.05, KF in situ gels versus KF eye drops. Data are presented as mean ± standard deviation.
Abbreviation: KF, ketotifen fumarate.
Conclusion
In the present study, an ophthalmic delivery system of ion-activated in situ gel for ketotifen with DGG was developed, and its rheological characteristics, stability, in vitro gelation, release in vitro, and pharmacodynamic activity in vivo were investigated. The formulation had an optimum viscosity that will allow easy drop as a liquid, which then underwent a rapid sol–gel transition due to ionic interaction. There were negligible alterations in the initial values of viscosity of the formulations over a storage period of 180 days. In vitro release showed that the release of ketotifen from in situ gels was moderate without any burst effects. Scintigraphic studies indicated that DGG could increase the residence time of the formulation. At the same dose, in situ gels demonstrated a typical sustained and prolonged drug-effects behavior than the common drops.

Disclosure
The authors report no conflicts of interest in this work.

References