Current Trends in Ophthalmology

Patients Preference of Diquafosol or Rebamipide Eye Drops for the Treatment of Mild Dry Eye: A Pilot Study

Maria Ohtani¹, Yukihiro Horie¹, Yoshiaki Tagawa², Susumu Ishida² and Nobuyoshi Kitaichi¹,²

¹Department of Ophthalmology, Health Sciences University of Hokkaido, Sapporo, Japan
²Department of Ophthalmology, Hokkaido University Graduate School of Medicine, Sapporo, Japan

*Correspondence: Nobuyoshi Kitaichi, Ainosato 2-5, Kita-ku, Sapporo 002-8072, Japan, Tel: +81-11-778-7575; Fax: +81-11-770-5034; E-mail: nobukita@hoku-iryo-u.ac.jp

Received: May 17, 2018; Accepted: July 09, 2018; Published: July 11, 2018

Abstract

Purpose: To investigate the preference rates of recently developed eye drops, 3% diquafosol ophthalmic solution and 2% rebamipide ophthalmic suspension, as a randomized clinical trial in patients with dry eye.

Methods: Eighteen patients (mean age: 59.7 years old, females: 77.8%) with reduced tear film break-up time (TBUT ≤ 5 sec) were enrolled in this study. They were treated with either diquafosol or rebamipide for 1 week, and treated for an additional week after switching to the other eye drop. This treatment was repeated one more time. Then, the patients were treated with one of either eye drop for an additional 4 weeks. Patients’ preferences for eye drops were investigated using a questionnaire at the end of 4-week treatment. To confirm the effectiveness of the treatments, objective and subjective assessments including superficial punctate keratopathy (SPK) score, TBUT measurement, and Dry Eye-Related Quality-of-Life Score (DEQS) were performed at baseline, and 4 and 8 weeks.

Results: SPK score was improved after treatment (P < 0.05). Patients tended to prefer diquafosol (64.7%) to rebamipide (35.4%), however there was no significant difference in two eye drops (P = 0.23). Bitterness was reported significantly more frequently after rebamipide administration (66.7%) than diquafosol administration (5.6%, P < 0.001). The diquafosol bottle was considered “ease to use” by a larger percentage of users (94.1%) as compared to the rebamipide bottle (58.8%, P < 0.05). The effectiveness of the signs and symptoms were similar for both groups at 8 weeks.

Conclusions: Diquafosol and Rebamipide were both effective in treating dry eye and 65% of the patients choose Diquafosol. Bitterness after administration and ease to use of the bottles seemed to affect the patients’ preferences for these two eye drops.

Keywords: Dry eye; Diquafosol; Rebamipide

Introduction

Dry eye is a multifactorial disease of the tear film and ocular surface that results in symptoms of discomfort, visual disturbance and tears film instability with potential damage to the ocular surface [1]. It has a high prevalence worldwide, with the reported incidence rates varying from approximately 5% to 35%, depending on the population studied[2]. Older age and female sex are widely recognized as the two most common risk factors for dry eye, based on the results of multiple epidemiological studies [2-5]. The pathophysiology of dry eye includes two mutually reinforcing mechanisms; the vicious circle between tear film instability and the ocular surface epithelial damage [6]. Keratoconjunctival epithelial damage and tear film instability eventually lead to aggravation of dry eye, and also to subjective symptoms, such as eye discomfort,
foreign body sensation or visual disturbance [7], thereby lowering the patient's Quality of Life (QOL) [8] or Quality of Vision (QOV) [9].

Currently, artificial tears, sodium hyaluronate solutions, corticosteroids and cyclosporine were utilized as conventional therapies for dry eye [10-12], however, these therapies have limited efficacy and safety. “Artificial tears” refers to temporary water and electrolyte supplementation [13]. Sodium hyaluronate solution has shown some effectiveness in patients with dry eye [14-16], however it is considered ineffective against conjunctival disorders caused by mucin layer damage [14]. Corticosteroids and cyclosporine should be avoided in some subjects for reasons like increased intraocular pressure, cataract, infection, and insurance system in Japan.

Recently, two pharmacological agents Diquafosol [17], an agonist of the P2Y2 receptor involved in water and mucin secretion, and Rebamipide [18] which can increase corneal and conjunctival mucin-like substances, were launched in Japan. Diquafosol ophthalmic solution is reported to significantly improve objective markers of dry eye disease such as corneal and conjunctival fluorescein staining in several randomized controlled trials, and improve tear film break-up time and Schirmer test scores in some studies [19-21]. It was reported that diquafosol sodium improved Schirimer test value and mucin-like substances among patients with dry eye [22,23]. Now diquafosol eye drops can be prescribed in China and Korea for patients with dry eye following Japan.

Rebamipide was initially developed and approved for use in treating gastric ulcers and lesions associated with gastritis. For the ophthalmic application, rebamipide ophthalmic suspension can effectively treat tear deficiency and mucin-related corneal epithelial damage, and restore the microstructure responsible for tear stability [18]. It was reported that the mucin-like substance on conjunctiva and cornea was increased by using Rebamipide in rabbit model [24]. Conjunctival goblet cell density significantly increased in patients with dry eye by using rebamipide [25]. Due to these favorable data, diquafosol and rebamipide are now widely used options for dry eye treatment in Japan.

To accumulate the evidences and to identify the eligible way of using these agents are very important clinically. In clinical practice, patient preference is of great significance because it strongly affects the treatment adherence of patients with dry eye who require long-term therapy. Therefore, we conducted a survey to investigate the preferences of patients with dry eye with respect to specific eye drops, diquafosol ophthalmic solution and rebamipide ophthalmic suspension, in a clinical study.

**Materials and Methods**

**Study Design**

This is a prospective, randomized controlled crossover study. Envelope method was approved as a sampling technique. Subjective and objective findings were assessed by the different observers. This study consisted of a survey to investigate patient preferences with respect to 3% diquafosol ophthalmic solution (Diquas ophthalmic solution 3%, Santen Pharmaceutical Co., Ltd, Osaka, Japan) and 2% rebamipide ophthalmic suspension (Mucosta ophthalmic suspension UD2%, Otsuka Pharmaceutical Co., Ltd, Tokyo, Japan) for the treatment of dry eye. After a screening period, patients were treated with either diquafosol or rebamipide for 1 week, and treated for an additional week after switching to the other eye drop. This treatment cycle was repeated one more time, and then patient preferences of eye drops were investigated with a questionnaire given at the end of the 4-week treatment. The patients selected their favorite one of two, and were treated with it for an additional 4 weeks. The patients were instructed to place a drop into the eye 4 times daily. The effectiveness of the eye drops was also confirmed at week 4 and 8 (Figure 1).

**Figure 1:** Study design. SPK; superficial punctate keratopathy, TBUT; tear film break-up time, DEQS; Dry Eye-Related Quality-of-Life Score.

Institutional Review Board of the Institute for Personalized Medicine, Health Sciences University of Hokkaido approved the protocol before the study initiation (No. 2013-001). This research was conducted in compliance with the Declaration of Helsinki, and prior written informed consent was obtained from all study patients.

**Patients**

The study subjects were adult patients (aged 20 years old or older) who complained dry eye-related symptoms, and visited the Department of Ophthalmology, Health...
Sciences University of Hokkaido Hospital, between June and October 2013, and were subsequently diagnosed as having dry eye disease according to the dry eye diagnostic criteria defined in 2006 by the Japanese Dry Eye Society [26]. Fluorescein paper was used to evaluate BUT and SPK. Corneal fluorescein staining was scored based on the modified criteria established by von Bijsterveld; each eye is divided into 3 sections (temporal conjunctiva, cornea and nasal conjunctiva) and scored from 0 to 3 (0, negative; 1, scattered minute; 2, moderate spotty; 3, diffuse blotchy staining). Patients with poorly-controlled diabetes, severe allergic conjunctivitis, significant conjunctival chalasis, and serious epithelial or endothelial damages were excluded from this study. No other subjects were participated as mild dry eye cases.

Study Assessments

Patient preferences for eye drops were investigated with a questionnaire at the end of 4-week treatment cycle. The questionnaires included “Which ophthalmic solution do you prefer?” and questions asking impression of each eye drop after use such as “Comfortable”, “Moistened”, “Be better” or “See clearly”, and feeling after use such as “Nothing”, “Irritation”, “Bitterness” or “Discharge”, and evaluation of each eye drop’s bottle such as “Easy to use”.

Objective assessment using SPK score or TBUT, and subjective assessment using Japanese version of DEQS [27] were also performed to confirm the effectiveness at baseline, week 4 and 8.

Statistical Analyses

For continuous variables, descriptive statistics included the number of patients (N), mean and standard deviation (SD). For categorical variables, the number and percentage of patients were calculated. Patient preferences derived from responses to “Which ophthalmic solution do you prefer?” were analyzed with chi-square test and the responses to impressions and feelings following each eye drop application, and evaluations of each eye drop’s bottle were analyzed with McNemar’s test. For efficacy evaluations, the changes from baseline variables at week 4 were analyzed using Wilcoxon signed-rank test and pair-wise comparisons between two eye drops at week 8 was analyzed using Wilcoxon rank-sum test (TBUT and DEQS). Similarly, the changes from baseline variables were analyzed using a paired t-test, and pair-wise comparison was analyzed using the Student’s t-test (SPK score). The significance level was set at 0.05 using two-sided tests.

Results

Demographics and Baseline Characteristics

Eighteen dry eye patients (mean age: 59.7 years old, 14 female: 77.8%) were enrolled in this study. They included 1 dentist, 1 nurse, 1 librarian, 1 office worker, 2 drivers, and 12 retired persons/housekeepers. The SPK score, TBUT and DEQS at baseline was 2.5 ± 2.2 (mean ± SD), 2.6 ± 1.2 sec. and 34.6 ± 28.0, respectively.

Patient Eye Drop preferences

Among the 18 patients, 17 patients answered the question related to their preferred eye drop among diquafosol and rebamipide (response rate: 94.4%) (Figure 2). Diquafosol (11 patients, 64.7%) tended to be preferred to rebamipide (6 patients, 35.3%), however, there was no statistical significance between two eye drops (P = 0.23, chi-square test).

![Figure 2: Questionnaire investigating the preferred eye drops among Diquafosol and Rebamipide. Patients answered the question ‘what ophthalmic solution do you prefer?’ P value was 0.225 (chi-square test).](image-url)

In the questionnaire survey, patients’ impressions and feelings of each eye drop after use, and patients’ evaluations of eye drop bottles were also investigated. Bitterness was reported significantly more often with rebamipide (12 patients, 66.7%) than diquafosol (1 patients, 5.6%, P < 0.001). A larger fraction of patients answered that the diquafosol bottle was easier to use (17 patients, 94.1%) as compared to the rebamipide bottle (10 patients, 58.8%, P < 0.05). No significant differences were observed for other questionnaire items related to patient impressions, such as “comfortable”, “moistened”, “be better”, “see clearly” or feelings after administration. Patients who answered that “the bottle is harder to use” for rebamipide were older (mean age ± SD: 65.3 ± 6.3 years old) than patients who answered “easy to use”.
(mean age ± SD; 54.8 ± 7.6, P < 0.01) (Figure 3).

Figure 3: Sub-analyses for the evaluations of each eye drop’s bottle based on patient's ages. The age of patients (Total N=17) who answered “The bottle is harder to use” or “easier to use” for (a) diquafosol and (b) rebamipide are plotted, and compared for each eye drop. P value was calculated by student's t-test.

Effectiveness

SPK scores significantly improved after 4 weeks of treatment with diquafosol and rebamipide compared to baseline (Figure 4a, P < 0.05). No significant difference in SPK score was observed at week 8 between the two groups (Figure 4a). TBUT tended to be prolonged at week 4 compared with baseline, but this was not significant, and the effectiveness of the two eye drops was similar at the end (Figure 4b). The patients’ QOL subjectively assessed using DEQS, also tended to be improved (Figure 4c).

Discussion

Dry eye is recognized as a chronic disease, frequently requiring long-term management [11]. As with many therapies for chronic diseases, some patients with dry eye do not adhere to long-term treatment with eye drops. To improve adherence to dry eye therapies, patient education, including direct instructions from physicians as well as an additional discussion with clinical staff focusing on duration of therapy, potential ocular adverse effects and tolerability, is thought to be an essential for the initiation of repeated courses of topical application of eye drops [28].

Each parameter was compared with baseline at week 4, and between groups at week 8. (a) SPK and (b) DEQS scores were analyzed by Wilcoxon signed-rank test in comparing baseline to at week 4, and by Wilcoxon rank-sum test in comparing between groups at week 8. Similarly, the change in (c) TBUT from baseline value was analyzed at week 4 by paired t-test, and compared between groups at week 8 by student's t-test.

Diquafosol is now widely used in Japan, China, and Korea. These 2 new eye drops would be approved for clinical use in other Asian countries before too long. Our present results provide useful information
to ophthalmologists and patients in other upcoming regions.

There was no statistical significance in preference between diquafosol and rebamipide, but some impression of the patients was different. The results of this survey revealed that bitterness was reported significantly more often with rebamipide than with diquafosol, and more patients found the bottle of diquafosol easier to use than that of rebamipide. These results seemed to correlate with the ratio of patients’ preferences with respect to these eye drops.

Rebamipide was reported to be effective and well-tolerated in a randomized multicenter phase III study, however dysgeusia (bitter taste) was commonly observed as an adverse event [28]. This is consistent with our survey results in clinical practices. As for diquafosol, eye irritation occurred with the highest frequency in a randomized multicenter phase 3 studies [21]. The results of the present survey also confirm the clinical trial data, indicating that irritation was reported more frequently with diquafosol than with rebamipide, but with no significant difference.

Diquafosol ophthalmic solution and rebamipide ophthalmic suspension differ with respect to the clinically supplied bottle-type, as the former is contained in multidose bottles containing 5 mL of solution each, while the latter is contained in disposable single-dose bottles containing 0.35 mL of suspension without preservatives. We conclude that a multi-dose bottle is more convenient for 4 to 6 times daily administration, and this is why more patients preferred diquafosol. Furthermore, the aged patients who responded “hard to use” was significantly more than that of patients who answered “easy to use” for rebamipide. This result suggests that an eye drop with “easy-to-use” is favorable for dry eye therapy because the incidence of dry eye is especially high among older patients [2-5]. These results indicates that patient preferences varied from patient to patient, and understanding the practical characteristics of the eye drops is very important for proceeding with the treatment.

Clinical efficacy was also confirmed in this trial, generally satisfying with the previous evidences related to diquafosol and rebamipide in phase III studies [21,29] and post-marketing surveillance [30]. These two new eye drops are also effective in clinical practice as a long time use, reported recently [31].

This study is the first to report patients’ preferences with respect to two major, novel eye drops, diquafosol and rebamipide, in real-world dry eye patients based on a questionnaire. However, the present study has some limitations; (1) small number of enrolled patients, (2) single-center study design and short treatment period of a total 8 weeks, (3) bitterness information beforehand, and (4) no washout period. In the present study, the patients changed eye drops 4 times without washout period, because we placed importance on looking to the subjective usability of these eye drops and on continuing treatment with less emotional strain to them. No washout design may influence the results partway in this study. Further studies are needed to establish the patient preferences with respect to these eye drops.

In conclusion, diquafosol and rebamipide are similarly effective in treating dry eye subjectively and objectively in clinical practice, and there was no significant difference in the preference of them. Our survey also suggests that patient preferences vary depending on their individual conditions, bitterness from rebamipide and easiness in using the diquafosol bottle seemed to correlate with the trend of patient preferences with respect to these eye drops.

Acknowledgements

We thank Ms. Shino Katayama and Ms. Marina Tokoro for their technical assistance.

References

6. Perry HD. Dry eye disease: pathophysiology,