REVIEW

Use of topical bromfenac for treating ocular pain and inflammation beyond cataract surgery: a review of published studies

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Abstract: Topical ophthalmic nonsteroidal anti-inflammatory drugs (NSAIDs) are commonly used to treat postoperative inflammation and pain following cataract surgery and for treatment and prophylaxis of pseudophakic cystoid macular edema (CME). Bromfenac is a brominated NSAID with strong in vitro anti-inflammatory potency. Like other ophthalmic NSAIDs, bromfenac is often used outside of the cataract surgery setting. This paper provides an overview of bromfenac's preclinical ocular pharmacology and pharmacokinetics, followed by a review of 23 published clinical studies in which various marketed bromfenac formulations were used for conditions other than cataract surgery or pseudophakic CME. These include: post-refractive eye surgery; macular edema associated with diabetes, uveitis, or retinal vein occlusion; inflammation associated with age-related macular degeneration; pain related to intravitreal injections; and other ocular anterior segment and surface disorders with an inflammatory component. The published evidence reviewed supports the safety and effectiveness of bromfenac in these additional ophthalmic indications. Bromfenac was well tolerated when given alone or in combination with intravitreal anti-vascular endothelial growth factor agents, topical corticosteroids, or topical mast-cell stabilizers. The most common adverse event reported was ocular irritation. No serious adverse events (ie, corneal epithelial disorders) were reported, although the majority of studies did not systematically evaluate potential side effects. Corneal complications, such as melts reported with diclofenac and ketorolac, were not observed with bromfenac in the studies. In summary, published study data support the clinical utility of bromfenac in various ocular disorders beyond post-cataract surgery. Additional studies are warranted to further define the potential role of bromfenac ophthalmic solution in clinical practice.

Keywords: bromfenac, clinical studies, ocular inflammation, pain, safety

Introduction

Nonsteroidal anti-inflammatory drugs (NSAIDs) are increasingly being utilized in ophthalmology for a variety of indications. NSAIDs exert anti-inflammatory and analgesic activity primarily through nonselective inhibition of cyclooxygenase-1 (COX-1) and cyclooxygenase-2 (COX-2) enzymes. 1-4 By inhibiting the COX pathway, NSAIDs limit conversion of arachidonic acid to prostaglandins. 1,3,5 Prostaglandins play a key role in stimulating pain and promoting miosis, vasodilation, disruption of the blood-ocular barrier, and leukocyte migration. Thus, prevention of prostaglandin formation by NSAIDs results in anti-inflammatory and other clinically beneficial effects (Figure 1). Reduction of vascular endothelial growth

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Figure I Cyclooxygenase pathway of prostaglandin production from membrane bound arachidonic acid and site of action of NSAIDs.

Adapted from Adv Ther, Voolume 27/ Edition 10, Schechter BA, Trattler W, Efficacy and safety of bromfenac for the treatment of corneal ulcer pain, 756–761, Copyright (2010), with permission from Adis, part of Springer Science+Business Media.

factor (VEGF) activity via inhibition of COX-2 may also contribute to beneficial effects of NSAIDs for some therapeutic indications in the retina.¹

Topical administration of NSAIDs is preferred for most ophthalmic indications as it results in high ocular drug concentrations, inclusive of therapeutic levels in posterior ocular tissues, with minimal systemic adverse events (AEs) when compared to systemic administration.^{6,7} Currently, several ophthalmic NSAIDs are indicated for the management of post-cataract surgery pain and inflammation (nepafenac, bromfenac) or post-cataract surgery inflammation (ketorolac, diclofenac). 8-11 Other FDA-approved indications include the temporary relief of pain and photophobia in patients undergoing corneal refractive surgery (diclofenac)¹¹ and the temporary relief of itching due to seasonal allergic conjunctivitis (SAC) (ketorolac). 12 Beyond these approved indications, and due to their ability to inhibit COX activity, other indications have been explored, including prevention and treatment of cystoid macular edema (CME) after cataract or refractive surgery; reduction of pain, discomfort, and inflammation following refractive surgery; treatment of macular edema associated with diabetes and ocular diseases, such as uveitis, branch vein retinal occlusion, and corneal ulcer pain. 1,13-25 Topical NSAIDs may also be beneficial in the treatment of age-related macular degeneration (AMD), which has a complex pathogenesis that includes a major inflammatory component. ²⁶ In addition, the anti-inflammatory properties of these agents may alleviate some of the signs and symptoms of anterior segment and ocular inflammatory surface disorders. ^{27–33}

Bromfenac is an NSAID developed and marketed for topical ophthalmic use. A twice-daily (BID) bromfenac 0.1% ophthalmic solution was approved in Japan in 2000 (Bronuck[®]; Senju Pharmaceutical Co., Ltd., Osaka, Japan) as a treatment for pain and inflammation associated with inflammatory eye disorders. The same BID formulation was subsequently FDA-approved in 2005 as bromfenac ophthalmic solution 0.09% (Xibrom®; ISTA Pharmaceuticals Inc., Irvine, CA, USA); equivalent in dose to the Japanese formulation (which is labeled as the salt) for the treatment of postoperative inflammation following cataract surgery. This same formulation is marketed as Yellox® in Europe. The US product was discontinued in 2011 following the introduction in 2010 of a once-daily (OD) formulation (bromfenac 0.09%; Bromday[®], ISTA Pharmaceuticals Inc.) indicated for treating postoperative inflammation and pain following cataract Dovepress Schechter

extraction with posterior chamber intraocular lens (IOL) implantation. A newer QD formulation of bromfenac (Prolensa®; Bausch & Lomb Incorporated), FDA-approved in 2013, has a lower and more physiologic pH (7.8 versus 8.3 for Xibrom), allowing for a reduced concentration (0.07%) while maintaining bioavailability. 10,34 Because bromfenac is a weak acid, reducing the pH decreases the proportion of ionized drug, facilitating penetration into targeted ocular tissues and allowing for the lower concentration to be utilized. 35,36 Bromsite® (bromfenac ophthalmic solution 0.075%; Sun Pharmaceutical Industries, Cranbury, NJ, USA) was approved in 2016 for treatment of postoperative inflammation and prevention of ocular pain in patients undergoing cataract surgery and is administered BID.³⁷ BromSite is formulated in DuraSite® (Sun Pharmaceutical Industries), a mucoadhesive matrix known to retain drug on the ocular surface.38

Bromfenac is the only brominated ophthalmic NSAID available (Figure 2). Halogenation of the molecule through the addition of a bromine atom in the 4-position of the benzoyl ring confers bromfenac with increased potency against COX-1 and COX-2 compared to other NSAIDs. ^{39–44} Multiple studies have evaluated the IC₅₀ (the drug concentration required to inhibit COX enzyme activity by 50%) of bromfenac with that of other NSAIDs using a variety of enzyme sources, including tissue extracts and/or recombinant COX enzyme preparations. The studies demonstrated that the potency of bromfenac in inhibiting both COX-1 and COX-2 was improved over that of other NSAIDs, except for ketorolac, for which the IC₅₀ against COX-1 was lower in one study (Table 1).

Bromination also increases the lipophilicity of bromfenac, which, in turn, may facilitate ocular tissue

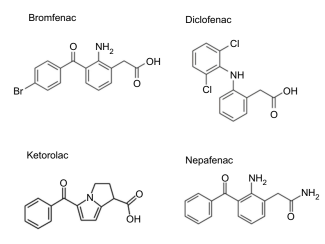


Figure 2 Chemical structures of bromfenac and other ophthalmic NSAIDs.

Table I Comparison of cyclooxygenase inhibitory activity of topical NSAIDs

Study	NSAID	COX-I IC ₅₀ (μ M)	COX-2 IC ₅₀ (μM)
Kida 2014 ⁴²	Bromfenac	0.00556	0.00745
	Diclofenac	0.0555	0.0307
	Amfenac	0.0153	0.0204
Walters 2007 ⁴³	Bromfenac	0.0864	0.0112
	Ketorolac	0.0139	0.0911
	Amfenac	0.138	0.00177
	Nepafenac	82.3	>1000
Waterbury 2006 ⁴⁴	Bromfenac Ketorolac Diclofenac	0.210 0.020 0.0079	0.0066 0.12 NA

Notes: Reprinted from PLoS One, Volume 9/ Edition 5, Kida T, Kozai S, Takahashi H, Isaka M, Tokushige H, Sakamoto T, Pharmacokinetics and efficacy of topically applied nonsteroidal anti-inflammatory drugs in retinochoroidal tissues in rabbits, e96481, Copyright (2014), permission from Public Library of Science (PLoS) is not needed as this manuscript is an open-access article distributed under the terms of the Creative Commons Attributions License. ⁴² Reprinted from J Cataract Refract Surg, Volume 33/ Edition 9, Walters T, Raizman M, Ernest P, Gayton J, Lehmann R, In vivo pharmacokinetics and in vitro pharmacodynamics of nepafenac, amfenac, ketorolac, and bromfenac, 1539–1545, Copyright (2007), with permission from Elsevier. ⁴³ Reprinted from Curr Med Res Opin, Volume 22/ Edition 6, Waterbury LD, Sillman D, Jolas T, Comparison of cyclooxygenase inhibitory activity and ocular nati-inflammatory effects of ketorolac tromethamine and bromfenac sodium, 1133–1140, Copyright (2006), with permission from Taylor & Francis Ltd (http://www.tandfonline.com). ⁴⁴

Abbreviation: IC₅₀, half maximal inhibitory concentration; NA, not applicable.

penetration through the cornea's layers. 41 The pharmacokinetics of various bromfenac ophthalmic solutions have been studied in rabbit ocular tissues, while in humans, aqueous and vitreous humor concentrations have been evaluated at several time points post-instillation. In rabbits, a single 50-µL ocular dose of bromfenac 0.09% solution resulted in detection of bromfenac in all ocular tissues (conjunctiva, cornea, lens, iris-ciliary body, aqueous humor, choroid, retina, and sclera) except for the vitreous humor, with levels sustained over 24 hrs, as determined by HPLC. 45 Similarly, following a single 50-μL topical dose of ¹⁴C-bromfenac 0.07% solution, bromfenac was detected using liquid scintillation chromatography in ocular tissues for up to 24 hrs with a similar distribution as bromfenac 0.09% (single 50-µL dose).³⁴ In another rabbit PK study comparing the ocular penetration of bromfenac following a QD 40-µL topical dose of either bromfenac 0.075% solution formulated in DuraSite or bromfenac 0.07% solution for a total of 9 days, there were no apparent differences in drug concentrations in either the anterior and posterior ocular tissues as measured by HPLC. 46 As expected, bromfenac

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0.075% in DuraSite dosed BID (the approved clinical dosing regimen) achieved significantly higher ocular concentrations compared to bromfenac 0.07% dosed QD.⁴⁶ However, both formulations reached tissue levels in excess of the reported IC_{50} for bromfenac. Concentrations of bromfenac in ocular surface tissues (cornea, conjunctiva) were not evaluated in this study, but, based on a previous study evaluating the ocular distribution of azithromycin formulated in DuraSite, 47 ocular surface concentrations are expected to be higher following instillation of bromfenac 0.075% in DuraSite as compared to bromfenac 0.07%. While there were no safety signals reported in a 16-day clinical safety and efficacy study of bromfenac 0.075% in DuraSite in patients undergoing cataract surgery, 48 the impact of NSAIDs on ocular surface integrity, particularly in patients with underlying corneal disease, is a potential class concern, and further studies are needed to establish ocular surface levels of the new DuraSite formulation.

In humans, topical ocular instillation of one drop of bromfenac 0.09% prior to cataract surgery resulted in peak levels of bromfenac in the aqueous humor at the time of surgery which were several-fold higher than the IC_{50} for COX-2, or 0.0112 μ M.⁴³ A single 50- μ L drop of bromfenac 0.1% instilled prior to the initiation of cataract surgery resulted in maintenance of clinically effective aqueous humor levels (above the IC50 value for COX-2) for more than 12 hrs. 49 Reductions in aqueous prostaglandin E2 concentrations at the time of surgery have also been demonstrated with bromfenac 0.09%, administered either BID 2 days preoperatively or BID 1 day preoperatively plus 1 drop in the morning before surgery, in patients undergoing cataract surgery. 50,51 Bromfenac 0.075% in DuraSite and bromfenac 0.09% dosed QD for 2 days and then 3 hrs prior to cataract surgery both achieved aqueous humor concentrations in excess of reported IC₅₀ values (Table 1), with a higher concentration achieved with the bromfenac/DuraSite formulation. 52 In one study of patients undergoing vitrectomy (not related to a vitreous hemorrhage), penetration of topical bromfenac 0.09% into the vitreous humor was observed at a concentration sufficient to significantly reduce vitreous prostaglandin E2 concentrations;⁵³ however these results were inconsistent with those of an earlier smaller study in vitrectomy patients, which did not observe a significant reduction in vitreous prostaglandin E2 concentrations following topical bromfenac 0.09% administration.⁵⁴

The safety and efficacy of bromfenac for the management of pain and inflammation following cataract surgery, its approved indication, has been reviewed extensively elsewhere, ^{55–57} as has its use for prevention and treatment of CME following cataract surgery. ⁵⁸ The purpose of the current paper was to provide a comprehensive review of published studies which have evaluated the use of bromfenac for the treatment of anterior or posterior ocular inflammatory conditions beyond the post-cataract surgery setting.

Search strategy

A search was conducted of the MEDLINE® database using the search terms ("bromfenac" NOT "oral") with no time constraints. This search returned 194 citations. All citations were hand-screened and included if they described a clinical study of any type (prospective, retrospective, case series, masked, open label) addressing the use of bromfenac for an ocular indication other than post-cataract surgery (ie, treatment of inflammation/pain or treatment/ prevention of CME). Case reports were not included. Foreign-language papers with English abstracts were eligible for inclusion. Of the 194 citations, 173 were excluded for the following reasons: pertained to cataract surgery/CME prevention (n=56); animal/in vitro research (n=36); reviews (n=26); pertained to systemic bromfenac pharmacokinetic/pharmacodynamic (n=11); case reports (n=6); comments/letters (n=5); studies in which bromfenac was not the focus of the research (n=5); and studies involving healthy volunteers (n=3). An additional three citations were identified through scanning bibliographies of published papers. A supplemental search of EMBASE using the terms "bromfenac NOT (cataract surgery)" did not identify any additional relevant citations. Thus, 23 citations were included in this review.

Post-refractive eye surgery

Photorefractive keratectomy (PRK), laser in situ keratomileusis (LASIK), and laser-assisted subepithelial keratectomy (LASEK) are effective surgical procedures used to correct low to moderate myopia and hyperopia. While safe, each technique can induce some degree of ocular pain, inflammation, and/or discomfort. For instance, ablation of the corneal epithelium and associated sensory nerves is a source of postoperative pain in PRK. ^{59,60} LASIK uses a microkeratome to raise a corneal flap, which is replaced following laser ablation, but disruption of nerve fibers below the surface in the corneal stroma can

cause significant discomfort. LASEK is a modification of LASIK in which the flap is more superficial and does not include anterior stromal tissue, thus maintaining the cornea's biomechanical stability and lowering the risk of flaprelated complications, yet patients undergoing this technique can still experience pain or discomfort. Although LASEK would theoretically be expected to cause less postoperative pain than PRK, a meta-analysis of 11 randomized, controlled trials did not find consistent differences between these procedures in pain scores at 1, 2, or 3 days postsurgery. Topical NSAIDs have been used to reduce postoperative pain and inflammation following refractive eye surgeries. 16–22

Several studies have evaluated bromfenac for the treatment of ocular discomfort following refractive surgery. In a prospective, double-masked, placebo-controlled case series, 64 patients (120 eyes) were treated with one of four regimens of bromfenac 0.07% and/or artificial tears (AT) to reduce perioperative ocular discomfort related to LASIK surgery (one drop immediately before and after the LASIK procedure): presurgical and postsurgical bromfenac; presurgical bromfenac and postsurgical AT; presurgical AT and postsurgical bromfenac; or presurgical and postsurgical AT (control).⁶¹ Mean postoperative eye pain rating and an averaged all-symptoms rating performed by patients (eye pain, tearing, itching, foreign body sensation, photophobia, discharge, haziness) using the Ocular Comfort Grading Assessment (OCGA) were both lower 1 hr postsurgery in subjects treated with presurgical bromfenac/ postsurgical AT (P=0.02). Among patients who received bromfenac both pre- and postsurgically, an all-symptoms OCGA mean score of zero, indicating no eye pain or other symptoms of discomfort, was reported at the 2-hr postsurgical assessment (P<0.05 vs control). Ocular discharge and tearing were the symptoms most improved by the bromfenac-containing regimens. Significant reductions in tearing vs control (AT only) were observed with presurgical bromfenac/postsurgical AT (1 and 3 hrs), pre- and postsurgical bromfenac (2 and 5 hrs), and presurgical AT/postsurgical bromfenac (5 hrs). Significant reductions in discharge vs control were observed at all time points through 5 hrs with pre- and postsurgical bromfenac and presurgical bromfenac/postsurgical AT; presurgical AT/postsurgical bromfenac was significantly different from control only at the 2-hr evaluation. Visual acuity (VA) was similar between all treatments at day 1 and 3 months postoperatively. No clinically significant AEs were observed.

An open-label study in 60 patients (120 eyes) who had undergone LASEK surgery followed by a 7-day course of topical dexamethasone 0.1% four times daily (QID)

compared bromfenac 0.1% BID to a tapered regimen of fluorometholone 0.1% (three times daily [TID] for 3 weeks, then BID for 4 weeks, then QD for 4 weeks) administered over the subsequent 11 weeks.⁶² The bromfenac group demonstrated improved VA compared to the fluorometholone group at 1 month (P<0.05) and similar VA at other time points. At 6 months, all bromfenactreated eyes achieved preoperative best corrected VA, while three eyes in the fluorometholone group failed to achieve preoperative best corrected VA. There were no significant differences between groups in intraocular pressure (IOP) or corneal topography at any time point. Haze was noted in two bromfenac-treated eyes at 1 month and at 2 months, and two fluorometholone-treated eyes at 1 month, but resolved with more frequent NSAID instillation and did not affect VA. Overall, bromfenac was reported to be well tolerated, and no patients discontinued treatment.

Four studies compared bromfenac to ketorolac following refractive eye surgery. 63-66 A prospective, randomized study compared bromfenac 0.1% BID (32 patients/eyes) with 0.5% ketorolac QID (32 patients/eyes), each given for 3 days before and 1 day after LASEK surgery. 63 Although the duration of postoperative irritative symptoms (a single outcome assessing symptoms such as pain, photophobia, foreign body sensation, and tearing) was significantly shorter in the bromfenac group (median, 2.0 vs 14.0 days; P=0.004), there were no significant differences between groups in corneal healing time or VA as measured from day 1 through month 3. A majority (87.5%; 28/32) of patients receiving ketorolac reported eye irritation and burning after administration (eg, pain and burning sensation), while no discomfort was reported in the bromfenac group.

In a prospective, double-masked study, 29 patients (58 eyes) undergoing PRK were randomized to treatment with nepafenac 0.1% (n=15 patients/30 eyes), bromfenac 0.09% (n=6 patients/12 eyes), or ketorolac 0.4% (n=8 patients/16 eyes); each topical NSAID was administered TID beginning 1 day preoperatively and continuing for 1 week postoperatively. By day 3, all groups showed a significant reduction in pain score compared to before NSAID use (P<0.05). Time to reepithelialization (complete closure) occurred significantly sooner with nepafenac (mean \pm SD, 5.5 \pm 1.59 days) and ketorolac (5.6 \pm 1.23 days) compared to bromfenac (7.3 \pm 2.53 days; P<0.05). Time to healing in this study was longer than reported in a previous study comparing nepafenac and ketorolac. The authors

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Table 2 Summary of studies

Study	Study Design	Efficacy Findings	Safety Findings
Post-refractive eye surgery	surgery		
Cleaveland 2017 ⁶¹	Prospective, double-masked, placebo-controlled, case series, post-LASIK Bromfenac 0.07% vs AT Four groups: 1. presurgical bromfenac/ postsurgical bromfenac 2. presurgical bromfenac/ 3. presurgical AT/postsurgical AT 4. presurgical AT/postsurgical AT	(i) Eye pain and the averaged all-symptoms ratings at I-hour were reduced compared with baseline in the pre-bromfenac/post-AT group only (P=0.02). (ii) Patients in the pre-bromfenac/post-bromfenac group had no discomfort at 2-hours (P<0.001). (iii) In the pre-bromfenac/post-AT group, patients had less tearing (P=0.02), foreign body sensation (P=0.01), photophobia (P<0.001), and discharge (P=0.01) at the I-hour time-point. (iv) Subjects treated with pre-bromfenac-/post-bromfenac had less tearing (P=0.01), foreign body	No notable AEs.
	All treatments given as one drop immediately before/after surgery N=64 (120 eyes)	sensation (P=0.03), photophobia (P<0.001), discharge (P<0.001) and haziness (P=0.02) at the 2-hour time-point. (v) Incidence of DLK was 8.5% (7/82) eyes treated peri-op with bromfenac vs. 15.8% (6/38) eyes treated with AT (P>0.05). (vi) VA was similar between treatments at 1 day and 3 months postoperatively.	
Chen 2015 ⁶²	Open-label post LASEK surgery Dexamethasone 0.1% QID for 7 days postsurgery, then 11 more weeks of either bromfenac 0.1% BID or FML 0.1% (tapered regimen) N=60 (120 eyes)	(i) No significant difference between groups in corneal topography. (ii) Improved VA in bromfenac group compared to FML at one month (P<0.05). No significant difference at other time points. (iii) No significant difference in mean IOP between groups at all time points.	Cases of grade -0.5 haze in both groups (2-bromfenac; 2-FML) which disappeared I month after treatment ended and did not affect VA. Ocular turgidity and blurred vision in I bromfenac and 3 FML-treated eyes; all treated with IOP-lowering agents.

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Table 2 (Continued).

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Study	Study Design	Efficacy Findings	Safety Findings
Dong 2015 ⁶³	Prospective, randomized study in LASEK patients	(i) No significant difference in postoperative irritation symptoms between groups	Transient local discomfort including stabbing pain and burning sensation after instillation of ketoro-
	Bromfenac 0.1% BID vs 0.5% ketorolac trometha-	(ii) Duration of postoperative irritation symptoms	lac; no discomfort noted with bromfenac.
	mine QID, each given 3 days before and I day after	was shorter in the bromfenac group (median, 2.0	
	Surgery N=64 patients/eyes	(iii) No significant difference in corneal healing	
		time and VA between groups.	
Durrie 2007 ⁶⁴	Prospective, randomized, double-masked, post-	(i) Reepithelialization occurred sooner with nepa-	No treatment-related AEs or evidence of corneal
	PRK	fenac and ketorolac compared to bromfenac	toxicity.
	Bromfenac 0.09% vs nepafenac 0.1% vs ketorolac	(ii) Significant reductions in pain scores were	
	0.4% (each given TID beginning I day preopera-	observed with nepafenac 0.1% on day 1	
	tively and continuing for one week	(P=0.0146) and day 3 (P<0.0001), ketorolac 0.4%	
	postoperatively)	on day 3 (P=0.0341), and bromfenac 0.09% on day	
	() () () () () () () () () ()	J (L-0.0369).	
	N=29 (58 eyes)		
Sher 2009 ⁶⁵	Open label post-PRK study	(i) No significant difference between groups in	No treatment-related AEs.
		pain, burning, photophobia, foreign body sensation over any of the days.	
	Bromfenac 0.09% BID vs ketorolac 0.4% QID;	(ii) No significant difference in epithelial healing	
	treatment continued until healing occurred	rate between groups.	
	N=149 patients (212 eyes)		
Wang 2011 ⁶⁶	Prospective, randomized, post LASEK or Epi-	(i) Bromfenac-treated eyes had lower pain scores	Eight patients (20%) reported temporary dry eye
	LASIK	than ketorolac-treated eyes on the surgery day	and 2 (5%) reported temporary irritation in both
	Bromfenac 0.09% in one eye vs ketorolac 0.5% in	and unlough a days postober advery (1 ~0.01). (ii) No differences in blurriness between groups	eyes. No serious AEs noted.
	the other (15 minutes and one minute prior to	on day of surgery or postoperatively through day	
	surgery, 2 and 4 hours postoperatively and daily	4 (P>0.01).	
	through day 4)	(iii) No difference in uncorrected VA between the	
	N=40 (80 eyes)	two groups (P>0.01).	

(Continued)

Table 2 (Continued).

Study	Study Design	Efficacy Findings	Safety Findings
Xu 2013 ⁶⁷	Prospective, case-controlled study in patients undergoing sub-Bowman's keratomileusis Bromfenac 0.1% BID for 10-14 days (depending on myopic severity) vs FML 0.1% for 16 days postoperatively N=74 (140 eyes)	(i) Postoperatively, there was a significant difference in favor of bromfenac in low-moderate myopic patients for mean corneal curvature K2 at I and 3 months (P<0.05), and in high myopic patients for VA at 3 months (P=0.015). (ii) No other postoperative differences between treatment groups.	In patients with low-moderate myopia, mean postoperative IOP at 1 month and 3 months was lower in bromfenac-treated patients as compared to FML-treated patients (7.8 vs 9.4 mm Hg and 8.1 vs 9.5 mm Hg, respectively; both P<0.05).
Macular Edema As	Macular Edema Associated with Uveitis, Diabetes, and RVO		
Pinna 2017 ⁶⁸	Open label pilot study in patients with newly diagnosed diabetic macular edema Bromfenac 0.09% BID for 30 days	(i) Mean CMT decreased from a baseline of 465.41 (118.47) µm to 388.88 (152.63) µm (P=0.02). (ii) Mean macular volume decreased numerically, however the decrease did not reach significance (P=0.06). (iii) Best-corrected VA was unchanged (P=0.72).	No AEs reported.
Radwan 2013 ⁶⁹	Retrospective, comparative case series, uveitic macular edema Bromfenac ^a BID monotherapy vs bromfenac BID +single IVB injection vs bromfenac plus single IVTA injection N=55 (67 eyes)	(i) A decrease in CMT and improvement in VA was observed in the IVB/bromfenac and IVTA/bromfenac groups compared to baseline at 1 and 3 months (P≤0.031). (ii) CMT in the IVTA/bromfenac group was lower than in the IVB/bromfenac and bromfenac monotherapy groups at 1 month (P≤0.041). (iii) No significant effects on CMT or VA in the bromfenac monotherapy group at 1 or 3 months compared to baseline.	Safety outcomes not reported.
			(Continued)

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Table 2 (Continued).

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Shimura 2015 ⁷⁰	Prospective, case-controlled, pilot study in patients with macular edema secondary to branch RVO All patients received initial IVB injection, then randomized to bromfenac 0.1% QID or AT QID for 48 weeks; IVB repeated as necessary for recurrence of macular edema N=44 (48 eyes)	(i) No significant differences in foveal thickness or VA between bromfenac and AT-treated eyes. (ii) Mean±SD number of IVB injections required was 3.8±1.1 for bromfenac vs 4.8±1.2 for control (P=0.014).	No AEs related to the topical bromfenac.
Age Related Macular Degeneration	lar Degeneration		
Grant 2008 ⁷¹	Retrospective case series in patients with wet AMD IVR vs IVR+bromfenac ^a BID for 6 months N=30 IVR; N=30 IVR+bromfenac BID	(i) Patients treated with bromfenac/IVR required significantly fewer IVR injections (1.6±0.69 ^b) than patients receiving IVR alone (4.5±0.41; P=0.0002). (ii) Improvement in VA was not significantly different between treatment regimens. (iii) Similar trends were observed for optical coherence tomography and fluorescein angiography data.	No AEs associated with topical bromfenac use.
Flaxel 2012 ⁷²	Prospective, randomized, open label comparative study in patients with wet AMD IVR monthly for 4 months, then monthly as needed, alone vs +bromfenac 0.09% BID for 12 months N=30	(i) No statistically significant difference in the number of IVR injections received over 12 months. (ii) Greater reduction in mean 12-month change in CMT in the combination group vs monotherapy group (P=0.03). (iii) Proportion of eyes experiencing >50 µm decrease in CMT was higher in the combination therapy group (P=0.046).	No significant difference in AEs between groups. Most common AEs in patients receiving bromfenac/IVR were burning/stinging (40% vs 20% with IVR alone), itchy eye (30% vs 50% with IVR alone), and headache (30% vs 40% with IVR alone). All AEs were mild.
Gomi 2012 ⁷³	Single-center, double-masked, placebo-controlled study in patients with wet AMD Bromfenac 0.1% vs AT BID for 6 months N=16 bromfenac; N=22 AT	(i) Mean number of ranibizumab injections over 6 months was lower (P=0.0274) in the bromfenac group (2.2) compared to the control group (3.2). (ii) Change in VA did not differ between groups (P=0.314). (iii) Trend towards a reduction in CRT in the bromfenac group (P=0.060).	No serious AEs or side effects attributed to bromfenac.

(Continued)

Table 2 (Continued).

Study Study Wygledowska-Prosp Promienska AMD 2014 ⁷⁴			
dowska- inska	Study Design	Efficacy Findings	Safety Findings
	Prospective, open-label study in patients with AMD	(i) VA improved in the IVB/ bromfenac group at 6 months vs baseline (P=0.001) but not with IVB alone.	No significant difference in IOP between groups.
8 N N N N N N N N N N N N N N N N N N N	IVB (3 injections then as needed)+bromfenac 0.09% for 3 months following first IVB injection vs IVB alone N=52 (26 in each group)	(ii) Median height of subretinal fluid measured by OCT decreased significantly with IVB/ bromfenac but increased in the IVB only group. (iii) More IVB injections were required in the IVB alone group than in the IVB/ bromfenac group (5.8 vs 6.9).	No severe AEs. No difference in ocular AEs between groups.
Wygledowska- Prosp Promienska 2015 ⁷⁵ IVA (3 + + + + + + + + + + + + + + + + + +	Prospective, open-label study in patients with AMD IVA (3 injections per month then as needed) +bromfenac 0.09% for 3 months following first IVB injection vs IVB alone N=54 (27 in each group)	(i) VA was improved in the IVA/ bromfenac group after 4 (P=0.001) and 6 months (P<0.001) while there was no significant improvement with affibercept alone. (ii) CRT did not change significantly between visits in either group. (iii) Mean height of subretinal fluid decreased in both groups between baseline and 6 months but no significant difference between groups.	No AEs observed.
Pain Related to Intravitreal Injections	eal Injections		
Georgakopoulos Singl 2017 ⁷⁶ blinc patie VEG treat ary t com minu rece givee N=6 bror	Single-center, prospective, randomized, doubleblind, placebo-controlled crossover study in patients requiring intravitreal treatment with antiveGF agents (ranibizumab or aflibercept) for treatment of AMD, DME, macular edema secondary to central and branch RVO, or angiod streaks complicated by choroidal neovascularization Bromfenac 0.09% or placebo in one eye 30-45 minutes before IVA or IVR; treatment not received during first intravitreal injection was given during consecutive injection in study eye N=65 (each patient served as own control, bromfenac vs vehicle)	(i) Pain perception was significantly lower immediately after intravitreal injection when patients received pretreatment with bromfenac vs placebo as assessed by the VAS score (P=0.002) and the main component of the short form of the SF-MPQ (P=0.001). (ii) Six hours after intravitreal injection, pain was significantly lower with bromfenac versus vehicle pretreatment as assessed by the VAS score (P<0.001) and the SF-MPQ (P<0.001). (iii) The median present pain intensity score was significantly lower in patients pretreated with bromfenac vs placebo 6 hours post-intravitreal injection.	No AEs observed.
Anterior Segment/Ocular Surface Disorders	r Surface Disorders		

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Table 2 (Continued).

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Study	Study Design	Efficacy Findings	Safety Findings
Sawa 1997 ³⁰	Multicenter, randomized, double-blind, placebo-controlled study in patients with ocular surface inflammatory disorders (blepharitis, conjunctivitis, scleritis, and episcleritis) Bromfenac 0.1% BID or panoprofen 0.1% QID for 2 weeks; placebo used for masking to maintain double-blind Safety: N=207 Efficacy: N=188	(i) Ratings of very effective or effective established for both bromfenac (63.4%) and pranoprofen (54.7%), with no significant difference between treatment groups. (ii) Excellent rates of efficacy demonstrated for blepharitis, conjunctivitis, scleritis, and episcleritis.	Bromfenac group: 4 patients with AEs (3.9%): increased eyelid inflammation, increased blepharitis, blood engorgement/follicle deterioration, sensitivity to infusion. Pranoprofen: 1 patient with an AE (1.0%): sensitivity to infusion. IOP decreased by 0.4 mm Hg in both groups but remained within normal limits.
Miyake-Kashima 2004 ⁷⁷	Investigator-masked, randomized study in patients with SAC Bromfenac 0.1% BID in one eye vs pemirolast 0.1% BID in contralateral eye for 1 week	(i) Subjective symptoms were not significantly improved in either group. (ii) Bromfenac improved signs of conjunctival injection (P=0.002), papilla formation (P=0.03), edema (P=0.002), and swelling of eyelid (P=0.05). (iii) Pemirolast improved signs of conjunctival injection (P=0.0002), papilla formation (P=0.003), and edema (P=0.004). (iv) No significant differences between treatments for subjective or objective symptoms.	No serious AEs were observed.
Fujishima 2009 ⁷⁸	Multicenter, investigator-masked, randomized study Bromfenac 0.1% BID+DSCG 2% QID in one eye vs FML 0.02% QID+DSCG 2% QID in the contralateral eye N=86	(i) All subjective symptom scores were significantly improved in both groups by I week (P<0.05). (ii) All objective signs and symptoms were significantly improved after I-week treatment with concomitant use of bromfenac or FML with DSCG vs. baseline (P<0.05), with no differences between treatments. (iii) Patient-assessed global evaluation was equivalent between the treatment groups.	No AEs reported and no reports of corneal epithelitis.

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 Table 2 (Continued).

Study	Study Design	Efficacy Findings	Safety Findings
U chio 2007 ⁷⁹	Prospective, randomized, double-masked study in patients with vernal keratoconjunctivitis Bromfenac 0.1% BID or saline eye drops; all patients received concomitant FML QID+tranilast QID vs placebo; mean follow-up=20.9 months N=22	The proportion of patients without recurrence (ie, "survival") at I year was 90.9% in the bromfenac group compared with 56.3% in the placebo group and at 2 years was 90.9% in the bromfenac group compared with II.3% in the placebo group; the difference in survival was statistically significant (P=0.01).	No clinically important side effects noted.
Fujishima 2015 ⁸⁰	Open-label, single-arm study in patients with dry eye disease who experienced no symptomatic improvement after I month of treatment with AT Bromfenac 0.1% BID+AT QID for I month, followed by 3 months of treatment with AT QID alone N=26	(i) Dryness scores significantly improved from pre-bromfenac to the end of bromfenac treatment (P<0.001) and then worsened by 3 months after discontinuation of bromfenac vs the end of bromfenac treatment (P<0.001). (ii) No significant changes in Schirmer scores were observed during the study. (iii) Tear film breakup time significantly improved from before bromfenac to the end of bromfenac therapy (P<0.001) and then worsened from the end of bromfenac discontinuation (P<0.001). (iv). Superficial punctate keratopathy improved from pre-bromfenac to the end of bromfenac therapy (P<0.001) and then worsened by 3 months after discontinuation of bromfenac (P<0.001).	No AEs observed.
Usui 1997 ⁸¹	Multicenter, open-label study in patients with anterior uveitis Bromfenac 0.1% BID for 2 weeks (short-term assessment) or up to 12 weeks (long-term assessment) Short-term evaluation: efficacy, n=40; safety, n=17 Long-term evaluation: efficacy, n=13; safety, n=17	(i) Based on physician evaluation of anterior protein, the percentage of patients with a rating of effective or very effective was 62.5% for shortterm treatment and 76.9% for long-term treatment. (ii) Based on evaluation of flare, the effectiveness rate was 51.9% for short-term treatment and 80.0% forlong-term treatment.	AEs during short-term treatment: I case of superficial punctate keratitis and 2 cases of sensitivity to instillation. No additional AEs occurred with long-term administration.

Table 2 (Continued).

Study	Study Design	Efficacy Findings	Safety Findings
Schechter 2010 ²³	Prospective, non-randomized trial in patients with (i) Primary endpoint of time to healing did not	(i) Primary endpoint of time to healing did not	No corneal AEs were observed.
	corneal ulcer pain whose eyes had bacterial or	differ significantly between treatment groups.	
	fungal infiltrates	(ii) Significantly more eyes treated with bromfenac	
	Bromfenac 0.9% BID (n=25 eyes) plus standard	had no pain by day 3 vs control eyes (52% versus	
	anti-infective agents or standard anti-infective	0%, P=0.023).	
	agents alone (controls; n=10 eyes)	(iii) By day 20, epithelial healing observed in 68%	
		of bromfenac-treated eyes vs 10% of control eyes	
		(P=0.040).	
		(iv) Bromfenac treatment was associated with a	
		shorter mean time to return to regular daily	
		activities (3.2 days versus 25.8 days in controls,	
		P<0.001) and a higher percentage of patients	
		returning to normal activities within 2 days of	
		starting treatment (71% vs 0% of controls,	
		P=0.018).	

Notes: ^aSource does not specify bromfenac concentration. ^bSource does not specify as standard deviation or standard error.

Abbreviations: AE, adverse event; AMD, age-related macular degeneration; AT, artificial tears; BID, twice-daily; CMT, central macular retinds thickness; CRT, central retinal thickness; DLK, diffuse lamellar keratitis; DME, diabetic macular edema; DSCG, disodium cromoglicate; FML, fluorometholone; IVA, intravitreal affibercept; IVB, intravitreal bevacizumab; IVR, intravitreal transitizumab; IVTA, intravitreal traindicolone acetonide; LASEK, laser epithelial keratomileusis; OCT, ocular coherence tomography; PRK, photorefractive keratectomy; QID, four times daily; RVO, retinal vein occlusion; SAC, seasonal allergic conjunctivitis; SF-MPQ, short form of the McGill Pain Questionnaire; TID, three times daily; VA, visual acuity; VEGF, vascular endothelial growth factor.

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suggested that this could have been a consequence of their study procedures, which included removing the bandage contact lens (BCL) and application of a drop of fluorescein sodium ophthalmic solution 1% at each postoperative visit. Physical manipulation of the BCL and application of fluorescein, which is known to be toxic to epithelial cells, may have delayed reepithelialization. Moreover, the concurrent use of different antibiotics (moxifloxacin 0.5% in nepafenac eyes and gatifloxacin 0.3% in all other eyes) along with differences in sample size may have confounded study results further. Nevertheless, there were no AEs reported and no evidence of corneal toxicity in any treatment group.

An open-label study compared the use of topical bromfenac 0.09% BID (n=105 eyes) with ketorolac 0.4% QID (n=107 eyes), each given postoperatively until epithelial healing occurred, for management of pain, discomfort, and photophobia following PRK.65 Patients undergoing unilateral surgery were randomly assigned to a treatment group; those having surgery on both eyes used one treatment in each eye. Following placement of a BCL on the operated eye, all patients received their assigned NSAID in addition to prednisolone acetate 1% eye drops and gatifloxacin 0.3% ophthalmic solution. No significant differences were noted with regard to postoperative pain, burning, foreign body sensation, photophobia, or epithelial healing rate between the two treatment groups at any time point, with defects fully resolved in both groups at day 5. There were no AEs that were considered related to either treatment.

Bromfenac 0.09% BID and ketorolac 0.5% OID were compared in patients undergoing LASEK (n=32) or epi-LASEK (n=8) in a prospective, randomized study.⁶⁶ For each patient, the right eye was randomized to either bromfenac or ketorolac, and the left eye was treated with the alternate NSAID. One drop of study medication was instilled in each eye 15 mins prior to surgery and 2 and 4 hrs after surgery, then patients were instructed to instill one drop QID for 4 days. For masking purposes, numbered eye-specific medication vials were provided to patients such that two of the daily bromfenac group instillations were AT. Patient-rated pain scores for the bromfenac-treated eves were lower as compared to ketorolactreated eyes on the day of surgery and every day through 4 days postsurgery (P<0.01 at each time point). Scores for visual blurriness were not statistically different between treatments at any time point. Epithelial healing was complete by day 6 for all eyes. Overall, AEs were similar between treatment groups; eight patients (20%) reported temporary dry eye in both eyes, and two patients (5%) reported temporary irritation in both eyes.

Overall, these studies suggest that bromfenac provides relief of pain and discomfort similar to that of other topical anti-inflammatories without apparent adverse consequences in patients following refractive eye surgery. While most studies comparing bromfenac with ketorolac reported no difference in epithelial healing rate, one study reported a shorter time to reepithelialization with nepafenac or ketorolac compared with bromfenac.

Macular edema associated with uveitis, diabetes, and retinal vein occlusion (RVO)

Macular edema is a build-up of fluid in the macula that can develop secondary to a number of conditions, including uveitis or as a complication of diabetes or branch RVO.83 Prostaglandins can contribute to the process by causing vasodilation and partial disruption of the blood-ocular barrier. 84,85 Macular edema can result in impairment of central vision and is a significant cause of vision loss in patients with metabolic, vascular, and inflammatory retinal diseases. Treatment of macular edema typically includes topical antiinflammatory agents, including corticosteroids NSAIDs. 86,87 Inhibitors of VEGF are also often employed since VEGF has been implicated in the pathogenesis of macular edema via multiple mechanisms.⁸⁷ The potential benefits of NSAIDs in macular edema relate to their ability to inhibit COX-2, resulting in reduced production of prostaglandins, although additional mechanisms of action have been proposed, including effects on leukotriene formation and modulation of chloride movement, thus affecting fluid movement through the retinal pigment epithelium.87 Additionally, NSAIDs may downregulate VEGF via inhibition of COX-2 activity, 1,88 and this mechanism of action may potentially contribute to the beneficial effects of NSAIDs in treating macular edema. The ability of topically applied NSAIDs to achieve therapeutic levels in the retina is relevant to the treatment of macular edema.⁶ A 2015 Cochrane Database systematic review did not identify any randomized, controlled trials of NSAIDs ongoing or completed studies for the treatment of diabetic CME.89

Three studies have evaluated the use of topical bromfenac alone or administered adjunctively with other medications in the treatment of macular edema (Table 2). A pilot study evaluated the effect of bromfenac 0.09%, instilled BID for 30 days in the affected eye, in 17 patients

with newly diagnosed diabetic macular edema (DME). Mean central macular thickness (CMT) decreased from a baseline of 465.41 to 388.88 μ m (P=0.02). Macular volume and best corrected VA did not change significantly over the course of the study, although there was a trend toward reduced macular volume (P=0.06). Bromfenac was well tolerated and no AEs were reported.

A retrospective, comparative case series compared outcomes among patients with uveitic macular edema who were treated with either bromfenac BID monotherapy (n=34), bromfenac BID in combination with a single intravitreal injection of bevacizumab (IVB) (n=21), or bromfenac BID in combination with a single intravitreal injection of triamcinolone (IVTA) (n=12).⁶⁹ Change (improvement) from baseline in VA (primary outcome) was noted in all treatment groups at 3 months, but was only significant for the bromfenac/IVB and bromfenac/IVTA groups ($P \le 0.017$). Likewise, CMT decreased in all groups, but only significantly in the bromfenac/IVB and bromfenac/IVTA groups (P≤0.009). It should be noted that mean baseline CMT was significantly (P=0.018) higher in the IVB and IVTA treatment groups vs the bromfenac monotherapy group; further, baseline mean VA was numerically lower (better) in the bromfenac monotherapy group, suggesting that the patients treated with IVB and IVTA were more severe and likely to demonstrate greater changes from baseline. Mean VA scores at 3 months were similar in the bromfenac monotherapy, bromfenac/IVB, and bromfenac IVTA groups (0.31, 0.35, 0.33, respectively; P=0.928).

The efficacy of bromfenac 0.1% in combination with IVB in 44 patients (48 eyes) with macular edema related to branch RVO was studied in a prospective case-control pilot study. Patients received IVB initially and repeated the treatment when macular edema (defined as foveal thickness \geq 300 µm) recurred. Prior to the second IVB injection, patients were randomized to treatment with topical bromfenac QID (n=24 eyes) or AT QID (n=24 eyes) for 48 weeks. On average, patients in the bromfenac group required fewer injections (mean±SD: 3.8 ± 1.1) of IVB than those in the saline group (mean±SD: 4.8 ± 1.2) throughout their clinical course (P=0.014). However, no significant differences in foveal thickness or VA were observed between bromfenac- and saline-treated eyes. There were no reported AEs related to bromfenac administration.

Overall, these small studies indicate that bromfenac may be of benefit alone in patients with DME or in conjunction with intravitreal corticosteroids/VEGF inhibitors for the treatment of uveitic macular edema or with VEGF inhibitors for the treatment of macular edema associated with RVO. Further investigation through more robust, controlled studies are necessary to evaluate the role of bromfenac monotherapy and combination regimens with corticosteroids or VEGF inhibitors in these conditions.

Inflammation associated with AMD

AMD is a leading cause of blindness worldwide. ⁹⁰ Intravitreal injections of anti-VEGF agents such as bevacizumab and ranibizumab are commonly used to treat neovascular (wet) AMD. ⁹¹ As inflammation plays an important role in AMD, the adjunctive use of topical NSAIDs has been studied in these patients. ²⁶ Possible mechanisms of action of NSAIDs in AMD include a reduction of inflammation and edema and minimization of COX-2-facilitated choroidal neovascularization. ^{72,92} Further, studies suggest that COX-2 promotes angiogenesis by modulating the expression of the VEGF receptor and thus inhibition of COX-2 is thought to downregulate VEGF. ^{1,88} Topical administration of NSAIDs for the treatment of AMD has several potential benefits, including minimal risk of complications/adverse effects and achievement of therapeutic levels in the retina. ^{1,72}

One retrospective case series and four prospective studies have evaluated bromfenac in combination with anti-VEGF agents in patients with AMD (Table 2). A retrospective case series compared 30 patients receiving intravitreal ranibizumab (IVR) for the treatment of choroidal neovascular membrane associated with neovascular AMD with 30 patients who received IVR in combination with bromfenac 0.09% BID over a period of 6 months.⁷¹ The addition of bromfenac significantly reduced the number of ranibizumab injections required to control choroidal neovascular membrane (1.6±0.69 vs 4.5±0.41, P=0.0002). Improvement in VA did not differ between the two treatment regimens, and similar trends were observed for optical coherence tomography (OCT) and fluorescein angiography data. Extended topical administration of bromfenac was not associated with any AEs.

An open-label pilot study evaluated the addition of bromfenac 0.09% BID for 12 months to treatment with IVR (administered monthly for 4 months, then monthly as needed) in patients with wet AMD.⁷² Patients who received combination therapy (n=20) were compared with those who received IVR alone (n=10). There were no significant differences between treatment groups in baseline characteristics, including CMT, VA, number of patients with newly diagnosed exudative AMD, and lesion type. In both groups, there was a significant reduction

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from baseline in CMT at 12 months (P<0.005). Compared with monotherapy, combination treatment with bromfenac plus IVR was associated with a greater mean reduction in CMT at 12 months ($-81.56~\mu M~vs~-42.50~\mu M,~P$ =0.03). Further, the proportion of eyes experiencing clinically relevant improvement (>50 $\mu M~decesion$) are greater for the combination group (P=0.046). There were no significant differences between treatment groups in the number of IVR injections received and no significant changes in VA between study groups at any time point. The overall frequency of ocular AEs, all mild in severity, was similar between treatment groups, although reports of burning/stinging occurred more often among patients receiving bromfenac.

A single-center, randomized, double-masked, placebo-controlled study evaluated topical bromfenac in patients with AMD with lesions <2 disk diameters. Patients were randomized 2:3 to receive either topical bromfenac sodium 0.1% (n=16) or AT BID (n=22) for 6 months in addition to at least one 0.5-mg IVR injection. Additional IVR injections were administered if there was evidence of intraretinal or subretinal fluid in the macula at visits at 3 and 6 months. The mean \pm SD number of IVR injections required over 6 months was lower in the bromfenac group (2.2 \pm 1.3) versus the AT group (3.2 \pm 1.5; P=0.027). Change in VA did not differ between groups (P=0.314), while a nonsignificant (P=0.060) trend toward a reduction in central retinal thickness in the bromfenac group was observed. One patient in the AT group discontinued treatment due to an unpleasant sensation.

In a prospective open-label study, patients with exudative AMD were treated with topical bromfenac 0.09% and IVB injections (n=26) or IVB injections alone (n=26).⁷⁴ Both groups received three initial IVB injections, with additional injections given each time an increase in subretinal fluid was noted on OCT examination. In the combination treatment group, bromfenac was administered BID for 3 months beginning after the first dose of IVB. At 6 months, VA improved significantly compared to baseline in the bromfenac group (P=0.001) but not in the group that received IVB alone. More IVB injections were required in the IVB-alone group than in the IVB+bromfenac group (5.8 [95% CI, 4.5-6.5] vs 6.9 [95%, 5.0-8.0). All AEs were mild or moderate in severity with no difference in the overall incidence of ocular AEs between treatment groups. Similar results were observed in a study by the same authors with similar methodology, except the intravitreal injections comprised aflibercept (IVA).⁷⁵ While VA improved significantly from baseline in the bromfenac+IVA group (n=27) after 4 months (P=0.001) and 6 months (P<0.001), there was no significant improvement in the group managed with IVA alone (n=27). Central retinal thickness did not change significantly between visits in either group. Median height of subretinal fluid (in μ m) on OCT decreased in both treatment groups between baseline and 6 months but was not significantly different between groups. No AEs were observed.

In summary, the adjunctive use of bromfenac for the treatment of AMD has demonstrated some clinical benefits compared with anti-VEGF therapy alone with no apparent safety issues. In some studies, the use of bromfenac led to a significant reduction in the number of intravitreal anti-VEGF injections required.

Pretreatment for pain related to intravitreal injections

Intravitreal injection is an invasive procedure that may cause pain during and after injections, which may contribute to pre-procedural anxiety.⁹³ Topical NSAIDS, including ketorolac and nepafenac, have been shown to reduce ocular pain during and/or after intravitreal injections.^{76,94,95}

A single-center, prospective, randomized, double-blind, placebo-controlled crossover study evaluated the efficacy and safety of pretreatment with bromfenac for pain in 65 patients requiring intravitreal injections with an anti-VEGF agent (Table 2). 76 Patients in this study were receiving IVR or IVA for the treatment of AMD, DME, macular edema secondary to central and branch RVO, or angioid streaks complicated by choroidal neovascularization. A single drop of bromfenac 0.09% or placebo (AT) was administered topically into one eye 30-45 mins before IVA or IVR; patients crossed over to receive the alternative treatment (bromfenac or placebo) during the next consecutive injection of IVR or IVA. Immediately after intravitreal injection, pain perception was significantly lower following pretreatment with bromfenac vs placebo based on both the visual analogue scale (VAS)score (P=0.002) and the main component of the short form of the McGill Pain Questionnaire (SF-MPQ; P=0.001). Similarly, at 6 hrs post-intravitreal injection, bromfenac treatment was associated with significantly lower pain scores compared with placebo on the VAS and the SF-MPQ (P<0.001 for both comparisons). The median pain intensity score was significantly lower in patients pretreated with bromfenac compared with placebo 6 hrs post-intravitreal injection (P=0.001) but not immediately after intravitreal injection. No AEs were observed during the study. These findings suggest that topical bromfenac may have a beneficial effect in reducing postinjection pain for up to 6 hrs following intravitreal injection.

Inflammation associated with ocular anterior segment/surface disorders

A number of ocular anterior segment and surface disorders have an inflammatory component and thus topical NSAIDs may be a suitable treatment option for these conditions. 96-98 Accordingly, topical NSAIDs have been found to be effective in patients with SAC, 27-29,99-101 vernal keratoconjunctivitis, 31,32 acute anterior uveitis, 102 and dry eye disease.³³ Bromfenac has been evaluated for a variety of anterior segment and ocular surface disorders, including blepharitis, SAC, vernal keratoconjunctivitis, scleritis and episcleritis, dry eye disease, anterior uveitis, and corneal ulcer pain (Table 2).

A multicenter, randomized, double-masked study compared bromfenac 0.1% BID with pranoprofen 0.1% QID, administered over 2 weeks, in Japanese patients with a range of ocular surface inflammatory disorders, including blepharitis, conjunctivitis, corneal inflammation, scleritis, and episcleritis.³⁰ Placebo was used to mask assignment to bromfenac vs pranoprofen. Efficacy was evaluated in 188 patients (bromfenac, n=93; pranoprofen, n=95) and safety in 207 patients (bromfenac, n=102; pranoprofen, n=105). Patients were assessed prior to instillation and on days 3, 7, and 14, and after treatment. For the entire population, good efficacy (ratings of "very effective" or "effective" based on a scoring system incorporating clinical observations and subjective/objective symptom scores) was reported for 63.4% (59/93) of bromfenac-treated patients and 54.7% (52/95) of pranoprofen-treated patients, with no significant difference between treatment groups. Both treatment groups showed excellent rates of efficacy for blepharitis, conjunctivitis, scleritis, and episcleritis, whereas the number of cases was too small for complete evaluation of efficacy for corneal inflammation. Four AEs (incidence of 3.9%) were reported in the bromfenac group (increased eyelid inflammation, increased blepharitis, blood engorgement/follicle deterioration, sensitivity to infusion), and one AE was reported in the pranoprofen group (1.0%; sensitivity to infusion).

Two published studies have evaluated bromfenac in patients with SAC. A Japanese study compared bromfenac 0.1% BID to the topical mast-cell stabilizer pemirolast 0.1% BID in 22 patients with a minimum 1-year history of SAC.⁷⁷ Each patient instilled one of the study

medications in one eye and the other study medication in the contralateral eye for 1 week. As evaluated by masked investigators, both drugs significantly improved conjunctival injection, papilla formation, and edema at 1 week (P<0.05 for all comparisons). Neither drug significantly improved subjective symptoms (itching, lacrimation, ocular discharge, redness, nasal discharge) and there were no significant differences between treatments in either subjective or objective symptoms.

A 1-week multicenter, randomized, investigatormasked study compared bromfenac 0.1% BID to fluorometholone 0.02% OID in 86 patients with SAC. 78 All patients used the mast-cell stabilizer disodium cromoglicate 2% QID in both eyes and were randomized to additional treatment with either bromfenac or fluorometholone in the right eye and the other study medication in the left eye. After 1 week of treatment, conjunctival itching, lacrimation, discharge, foreign body sensation, and conjunctival injection were significantly improved from baseline in both groups (P<0.05). There were no significant differences between treatments. No treatment-related AEs were observed, and there were no reports of corneal epithelitis.

A randomized, double-masked, placebo-controlled study evaluated long-term use of bromfenac 0.1% BID as adjunctive therapy to fluorometholone OID and the mast-cell stabilizer tranilast QID in 22 patients (21 males, 1 female; mean age 14.5 years; range, 4–34 years) with vernal keratoconjunctivitis, an allergic ocular condition typically seen in young boys and in countries with a warm climate.⁷⁹ Patients were randomized to receive bromfenac 0.1% (n=11) or placebo (normal saline eye drops, n=11) BID. Patients used weekly diary cards to record severity of symptoms on a scale from 0 (none) to 3 (severe). Investigator assessments of disease severity (clinical assessment and slit lamp evaluation) were performed at clinic visits at months 1, 2, and 4, then approximately every 3 months thereafter. Clinical recurrence was defined as an increase of three or more points in the "clinical score," defined as the total score for three subjective (itching, eye discharge, photophobia) and seven objective (conjunctival hyperemia, chemosis, follicles, giant papillae, limbal edema, Trantas' dots, corneal lesions) findings. Mean duration of follow-up was 18.7 months in the bromfenac group and 23.7 months in the placebo group. A greater proportion of patients receiving bromfenac vs placebo eye drops were without clinical recurrence (ie, "survived") at 1 year (90.9% vs 56.3%) and at 2 years (90.9% vs 11.3%); the difference

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in survival was statistically significant (*P*=0.01). No clinically important AEs were observed.

An open-label study evaluated the effect of adding bromfenac 0.1% to AT on signs and symptoms of dry eye in 26 Japanese patients who experienced no symptomatic improvement after 1 month of AT alone. 80 Bromfenac BID was administered along with AT QID for 1 month, with AT treatment alone continued for the subsequent 3 months. Dryness scores at the end of bromfenac treatment were significantly improved versus pre-bromfenac scores (P<0.001) as well as dryness scores 3 months after discontinuation of bromfenac treatment (P<0.001). While no significant changes in Schirmer scores were observed during the study, the tear film breakup time (BUT) was significantly improved at the end of bromfenac therapy (mean ±SD: 4.4±2.3 s) compared with before initiation of bromfenac (mean±SD: 2.8±1.8 s, P<0.001); BUT declined significantly from the end of bromfenac therapy to 3 months after bromfenac discontinuation (P<0.001). Similarly, superficial punctate keratopathy improved significantly from pre-bromfenac to the end of bromfenac therapy (P<0.001) and then worsened by 3 months after discontinuation of bromfenac (P < 0.001). No AEs were reported.

A multicenter, open-label Japanese study assessed bromfenac 0.1% BID for the treatment of anterior uveitis.⁸¹ Both short-term (2 weeks; efficacy, n=40; safety, n=48) and long-term (12 weeks; efficacy, n=13; safety, n=17) administration were evaluated; patients continued treatment for up to 12 weeks if the investigator determined that control of inflammation was possible or necessary through continued administration of bromfenac. The percentage of patients with a rating of effective or very effective based on physician evaluation of anterior protein was 62.5% for short-term treatment and 76.9% for long-term treatment. The effectiveness rate based on evaluation of flare was 51.9% for shortterm treatment and 80.0% for long-term treatment. During short-term treatment with bromfenac, AEs included one case of superficial punctate keratitis and two cases of sensitivity to instillation. No additional AEs occurred with longterm administration.

Bromfenac has also been evaluated for the treatment of corneal ulcer pain in a prospective, non-randomized trial.²³ Eyes with bacterial or fungal infiltrates were treated either with bromfenac 0.9% BID (n=25 eyes) plus standard anti-infective agents or with standard anti-infective agents alone (controls; n=10 eyes). Although the primary endpoint of time to healing did not differ significantly between treatment groups, significantly more eyes treated with

bromfenac had no pain by day 3 compared with control eyes (52% vs 0%, P=0.023). The epithelium of bromfenac-treated eyes was also more likely to have healed by day 20 (68% vs 10% of control eyes, P=0.040). Additionally, bromfenac was associated with significant functional improvement, reflected by a shorter mean time to return to regular daily activities (3.2 days vs 25.8 days in controls, P<0.001) and a higher percentage of patients returning to normal activities within 2 days of starting treatment (71% vs 0% of controls, P=0.018). Bromfenac treatment was not associated with any corneal AEs, although larger studies are needed to establish the safety of bromfenac in this patient population. It should also be noted that NSAIDs may cause sterile corneal ulcers with continued use, especially in at-risk patients. ¹⁰

Results of these studies suggest that bromfenac may have benefits across a range of anterior segment and ocular surface disorders. In all studies, bromfenac was well tolerated with no clinically important side effects reported. It should be noted that use of bromfenac and other topical NSAIDs has been associated with adverse corneal sequelae such as corneal erosions, ulceration, and perforation, particularly in patients considered high risk (complicated oral surgeries, corneal denervation, corneal epithelial defects, diabetes mellitus, ocular surface diseases, rheumatoid arthritis, repeat ocular surgeries within a short time) and those treated with prolonged use after (>14 days) or before (>24 hrs) surgery.

Summary

Topical NSAIDs, while generally approved for management of postsurgical pain and inflammation following cataract surgery, are increasingly being investigated for their utility in other ocular conditions with an inflammatory component. The evidence reviewed here supports the safety and efficacy of bromfenac for a number of indications other than its approved indication (treatment of post-cataract surgery inflammation and pain) or the non-approved but clinically well-established use for prevention of post-cataract surgery CME. Several studies have demonstrated that bromfenac decreases eye pain and irritative symptoms following refractive surgery with no signals for increased rates of side effects or adverse changes in VA or healing/corneal epithelialization. 61-66 Bromfenac has shown promise in reducing retinal thickening in patients with DME and has been reported to be beneficial in the management of uveitic macular edema when used in combination with IVTA or IVB including improvements in CMT and VA⁶⁹ and

reduced frequency of intravitreal injections.⁷⁰ Results of several studies suggest that topical bromfenac may be beneficial in patients with anterior segment and/or inflammatory ocular surface disorders, including blepharitis,³⁰ conjunctivitis/SAC/vernal keratoconjunctivitis, scleritis/ episcleritis, 30,77-79 dry eye disease, 80 anterior uveitis, 81 and corneal ulcer pain.²³

The studies reviewed herein demonstrated that bromfenac was well tolerated when given alone or in combination with intravitreal anti-VEGF agents, topical corticosteroids, or topical mast-cell stabilizers, with the most common AEs being signs of irritation. Corneal complications have been **NSAIDs** (particularly with diclofenac, ketorolac)103,104 but were not reported in these reviewed studies. Overall, these findings are consistent with the generally good safety profile of topical ophthalmic NSAIDs. 104

In conclusion, a growing body of research suggests the potential utility of the topical NSAID bromfenac for various ocular disorders. Additional studies are warranted to further define the potential role of bromfenac ophthalmic solution in clinical practice.

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Disclosure

The author reports no conflicts of interest in this work.

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