Topical Nutraceutical Optixcare EH Ameliorates Experimental Ocular Oxidative Stress in Rats

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Abstract

Purpose: Based on the hypothesis that oral nutraceuticals do not adequately reach all ocular tissues in the anterior segment, we evaluated the ability of a 3% concentration of the ingredients in a topical nutraceutical antioxidant formulation called Optixcare Eye Health (Optixcare EH) to ameliorate oxidative stress in rat models of age-related ocular diseases.

Methods: Diabetes was induced by tail-vein injection of streptozotocin, and the development of cataracts was monitored by slit lamp. Young rats were exposed to ultraviolet (UV) light, and the reduction in lens glutathione (GSH) levels and increase in 4-hydroxynonenol (4-HNE) were measured. Oxidative stress in the neural retina was generated by exposure of dark-adapted rats to 1,000 lx of light, and oxidative stress markers were measured. Dry eye was induced in rats by twice daily (b.i.d.) subcutaneous scopolamine injections. Topical Optixcare EH was administered b.i.d. and compared in select experiments to the multifunctional antioxidant JHX-4, the topical aldose reductase inhibitor (ARI) Kinostat/C212, oral Ocu-GLO/C212, and the topical ocular comfort agents Optixcare Eye Lube, Optixcare Eye Lube + Hyaluron, and Idrop Vet Plus hyaluronic acid.

Results: In diabetic rats, topical ARI treatment prevented cataract formation while the nutraceuticals delayed their development with Optixcare EH > Ocu-GLO. In UV-exposed rats, the reduction of GSH and increase in 4-HNE in the lens were normalized in order JHX-4 > Optixcare EH > Ocu-GLO. In the retina, oxidative stress markers were reduced better by oral JHX-4 compared with topical Optixcare EH. In the scopolamine-induced dry-eye rats, tear flow was maintained by Optixcare EH treatment, while none of the comfort agents examined altered tear flow.

Conclusions: Topical administration of a 3% concentration of the ingredients in Optixcare EH reduces experimentally induced reactive oxygen species in rats exposed to several sources of ocular oxidative stress. In addition, Optixcare EH maintains tear volume in scopolamine-induced dry eye. This suggests that in the anterior segment, the ingredients in Optixcare EH may have clinical potential against ocular oxidative stress.

Introduction

Population aging is rapidly occurring in many parts of the industrialized world. In the United States, the population of those aged 65 years or older is expected to increase from 40.2 million in 2010 to 88.5 million in 2050. Accompanying this increase in population aging is the prevalence of age-related ocular diseases that include cataracts, macular degeneration (AMD), and keratoconjunctivitis sicca, also known as dry eye disease (DED). Cataracts account for the majority of blindness in the world today, while AMD is the leading cause of visual impairment and irreversible blindness in industrialized countries. DED, which affects the majority of people over the age of 65 years, results in significant ocular discomfort and visual dysfunction. While DED generally does not result in vision loss, superficial keratitis, if left untreated, can lead to corneal ulcers or scars, and result in pain.

At present, there is no established pharmaceutical treatment for cataracts, and vision in patients with cataracts can only be restored by surgery. Cataract surgery has become one of the most frequently performed procedures in ophthalmology, accounting for $6.8 billion of the $16.2 billion spent in direct medical costs in 2006 on visual disorders among US adults aged 40 years or older. Delaying the onset and progression of cataracts through pharmaceutical intervention by 10 years has been estimated to reduce the need for cataract surgery by 50%. The need for the development of a nonsurgical treatment for delaying cataract is supported by the recent finding that cataract surgery increases the
incidence of late AMD. Similarly, at present, there is no medical treatment for preventing the onset or reducing the enlargement of geographic atrophy (dry AMD).

The pathogenesis of cataracts and AMD have not been elucidated. Both are multifactorial diseases where increased oxidative damage, accompanied by increased levels of iron, influences their development and progression. Risk factors for both cataracts and AMD include smoking, which is associated with both the generation of reactive oxygen species (ROS) and increased tissue iron levels. Cataracts are also associated with increased iron levels and iron disorders such as hyperferritinemia. ROS generated in the presence of iron results in hydroxyl radical formation through the Fenton reaction and mitochondrial dysfunction. Dysregulation of iron metabolism and iron handling proteins are also linked to the pathogenesis of AMD, with increased chelatable iron observed in the retinal pigmented epithelial cells (RPE) and Bruchs membrane in retinas from AMD patients. Iron overload is associated with a maculopathy that clinically resembles AMD, including the development and progression of multiple subretinal yellowish-white lesions and RPE cell atrophy in patients with aceruloplasminemia. Smoking also increases the risk of dry-eye syndrome and exacerbates existing dry-eye conditions. Similarly, systemic administration of the iron chelator deferiprone protects against light-induced retinal degeneration and diminishes oxidative stress in the mouse retina. Melatonin, a neurohormone with free radical scavenger activity, protects cultured human RPE against hydrogen peroxide-induced cell death. The AREDS antioxidant formulation reduces the risk of progression to advanced AMD by 25%. Several large clinical studies have also examined whether antioxidants can clinically reduce cataract progression. However, these studies suggest that any effect of antioxidants on cataract development is likely to be very small and probably is of no clinical or public-health significance.

Targeting oxidative pathways associated with these age-related diseases suggests therapeutic potential. For example, the clinical administration of a combined formulation of antioxidants and omega-3 essential fatty acids significantly improves the subjective symptoms of dry eye, while topical administration of the antioxidant and chelator epigallocatechin gallate reduces the clinical signs and inflammatory changes of DED in murine dry eye. Similarly, systemic administration of the iron chelator deferiprone protects against light-induced retinal degeneration and diminishes oxidative stress in the mouse retina.

ROS is also reduced by lowering tissue levels of iron, which can generate reactive hydroxyl radicals through the Fenton reaction. Studies with the deferoxamine suggest that the use of chelating agents to reduce lenticular ROS may be beneficial because patients with β-thalassemia receiving more chelation therapy had a lower incidence of cataracts. Similarly, in rats, cataracts from tobacco smoke exposure were decreased by deferoxamine treatment.

Kador has synthesized novel multifunctional antioxidants with the ability to scavenge free radicals and chelate iron independently, and demonstrated that by oral administration, they both delay cataract formation and protect the retina against light damage, a model for AMD.

Based on the hypothesis that oral administration of antioxidants generally fail to achieve adequate levels of antioxidants to the lens, a topical nutraceutical suspension called Optixcare Eye Health (Optixcare EH) is being developed. Composed of 4 unique antioxidants that are designed to reduce oxidative stress through free radical scavenging and chelating activity, these nutraceuticals are topically delivered in a viscous proprietary carbomer vehicle with established topical adhesion properties that have been shown to enhance the ocular uptake of select drugs. Using rat models associated with cataract formation, light-induced retinal degeneration, and scopolamine-induced dry eye, we demonstrate that the topical administration of a 3% suspension of the nutraceutical components in this formulation has therapeutic potential.

Methods

All reagents employed were of reagent grade. Optixcare Eye Lube (Optixcare) and Optixcare Eye Lube + Hyaluron (Optixcare + Hyaluron) were obtained from Aventix (Burlington, ON, Canada). The topical nutraceutical referred to here as Optixcare EH was prepared by blending 3% w/w of each of the 4 unique chemical grade antioxidants in Optixcare EH with Optixcare Eye Lube. Idrop Vet Plus was purchased from imed Pharma (Dollard-des-Ormeaux, QC, Canada), and the oral antioxidant formulation Ocu-GLO was purchased from Animal Health (Bellingham, WA). The aldose reductase inhibitor Kinostat was obtained from Therapeutic Vision, Inc. The multifunctional antioxidant JHX-4 (4-(5-hydroxypyrimidin-2-yl)-N,N-dimethyl-3,5-dioxopiperazin-1-sulfonamide (99% pure by HPLC)) was synthesized as previously described. Topical JHX-4 was prepared by suspending 3% (w/w) of compound in Optixcare Eye Lube. RIPA buffer was purchased from Cell Signaling Technology (Danvers, MA), and Halt Protease and Phosphatase Inhibitor Cocktail was obtained from VWR International (West Chester, PA). Antibodies against thiorodoxin (TRX) were purchased from Cell Signaling Technology, and antibodies against thiorodoxin reductase (TRXR) were purchased from Abcam, Inc. (Cambridge, MA). Chemiluminescent reagent and peroxide, biotinylated protein ladder, and prestained protein markers were obtained from Cell Signaling Technology. Polyacrylamide Ready Gels (4%–15% Tris-HCl) and Trans-Blot nitrocellulose membrane were purchased from Bio-Rad Laboratories (Hercules, CA). OxiSelect HNE-His Adduct enzyme-linked immunosorbent assay (ELISA) kit, OxiSelect Nitrotyrosine ELISA kit, and the OxiSelect HNE Adduct Competitive ELISA Assay were purchased from Cell Biolabs, Inc. (San Diego, CA). Zone-Quick Phenol Red Threads were purchased from FCI Ophthalmics (Pembroke, MA).

Animals

All procedures involving live animals were performed in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Vision Research. Sprague Dawley rats were purchased from Sasco (Wilmington, MA); Wistar rats and Lewis rats were obtained from Charles River Laboratories (Wilmington, MA).

Drug administration

The topicals Optixcare Eye Lube, Optixcare EyeLube + Hyaluron, Optixcare EH, Idrop Vet Plus, Kinostat, and topical JHX-4 were administered twice daily (b.i.d) to each
eye at approximately 8 am and 5 pm. The contents in the capsules of Ocu-GLO were dissolved in an appropriate amount of Tween 80 so that administration of 1 mL of the mixture delivered the manufacturer’s suggested amount of nutraceutical formulation per weight of the animal. This 1 mL amount was administered daily by gavage at approximately 9 am. Oral JHX-4 was administered as 0.05% (w/w) in standard rat chow as previously described.34,35

Diabetic cataracts

Diabetes was induced in 24 young (100 g) Sprague Dawley rats by tail-vein injection of 75 mg/kg of streptozotocin as previously described.34 All rats with blood glucose levels >300 mg/dL were then equally divided into 4 groups of 6 rats each. The first group served as untreated controls, while the second group received 1 drop of the topical aldose reductase inhibitor Kinostat to each eye b.i.d., the third group topically received 1 drop of the nutraceutical Optixcare EH to each eye b.i.d., and the fourth group received 1 mL of Tween 80 Ocu-GLO solution once daily orally as described above. Lens changes were monitored weekly by hand-held slit lamp following dilation of each eye with 1% tropicamide ophthalmic solution, USP (Falcon Pharmaceuticals Ltd., Fort Worth, TX). Lens changes were evaluated on a scale of 0 to 3, with 0 corresponding to no lens changes, 0.5 to suture accentuation, 1 to vacuole formation, 2 to cortical opacity, and 3 to mature cataract. The study was terminated 8 weeks after induction of diabetes, and the lenses were carefully removed from each enucleated eye using a posterior approach and photo-documented. Blood glucose levels at the onset of the study were determined using a commercial glucometer (Freestyle by Therasense, Alameda, CA). HbA1C levels were determined at the end of the study using the Bayer Metrika A1cNOW Plus System test kit (San Diego, CA).

UV irradiation of one eye only

Twenty-five female Sprague Dawley rats (100 g) were divided into 5 equal groups. One group served as the nonirradiated control, while the second group served as the untreated irradiated control. The third group received topical JHX-4, the fourth received topical Optixcare EH, and the fifth group received 1 mL of the oral Ocu-GLO as described above. Treatments began 3 days prior to irradiation and continued until each experiment was terminated. Prior to irradiation, each rat was anaesthetized by intraperitoneal injection of a mixture of 60 mg/kg of ketamine and 10 mg/kg of xylazine, and the right eye was dilated with 1% tropicamide ophthalmic solution. The contralateral eye was covered with an eye patch that was held in place with black tape. The dilated eye was then exposed for 15 min at a distance of 20 cm to 1,600 μW/cm² of UV-B as above. Two days after irradiation, the experiment was terminated. Each rat was euthanized with carbon dioxide asphyxiation. Both eyes were enucleated, and the lenses were carefully removed by a posterior approach.

Oxidative stress analysis. Following morphological examination of each lens under retroillumination with a dissecting microscope, each lens was homogenized in a ground glass homogenizer with ice-cold lysis buffer (pH 7.5), and the insoluble proteins were removed by centrifugation at 4°C. Protein levels in an aliquot from each supernatant were measured according to Bradford Assay.37 Aliquots of the supernatant were then deproteinized with equal volumes of 20% TCA, and the glutathione (GSH) levels in the deproteinized supernatants were measured at 412 nm according to the DTNB method.38 4-Hydroxynonenal (4-HNE) levels were then determined by competitive ELISA in the remaining homogenate supernatants by using the OxiSelect HNE Adduct Competitive ELISA Assay Kit (Cell Biolabs, Inc.) according to the manufacturer’s instructions. All assays were done in triplicate.

Light-induced retinal damage

This study was conducted as previously described in detail.35 Briefly, groups containing twelve 4-5-week-old male Wistar rats receiving either normal rat chow (untreated), normal rat chow and topically treated with Optixcare EH b.i.d., or an oral diet containing 0.05% of the multifunctional antioxidant JHX-4 as described above were dark adapted for 2 weeks. Rats from each group were then individually placed into acrylic, flat-bottom rodent restraints (Plas-Labs, Inc., Lansing, MI) in a light-box apparatus and exposed for 3 h to 1,000 lx of cool white fluorescent light (Lights of America, Los Angeles, CA; light-damaged rats, LD). An additional group of nontreated rats was also placed into the light-box apparatus for 3 h, but not exposed to light (nonlight-damaged rats, NLD). Immediately following light exposure, 8 LD rats from each group and 8 NLD rats were immediately euthanized, and the neural retinas were carefully dissected from the enucleated eye and frozen on dry ice. The remaining rats in each group were returned to the dark environment for 7 additional days. Each rat was then euthanized by carbon dioxide asphyxiation, and the dorsal ventral orientation of each eye was marked with a cautery tool. The eyes were then enucleated and fixed in a 3:1 (v/v) methanol:acetic acid solution overnight. The fixed samples were then processed by the UNMC Tissue Sciences Facility (www.unmc.edu/tissuesciences/) for subsequent morphological analyses.

Oxidative stress analysis. Oxidative stress markers were analyzed as previously described.35 Briefly, thawed retinas were homogenized in lysis buffer for ELISA determinations of 4-HNE-His Adduct and Nitrotyrosine using ELISA kits. Additional thawed retinas were homogenized in RIPA buffer containing Halt Protease and Phosphatase Inhibitor Cocktail for Western blot analysis of thioredoxin (TRx) and thioredoxin reductase (TRxR) expression levels. Protein samples were separated by SDS-PAGE and transferred onto nitrocellulose membranes. After blocking the membrane with 5% dry milk in 0.1% Tris-buffed saline with Tween (TBST), the membranes were then incubated overnight at 4°C with the primary antibodies anti-TRx (1:1,000 dilution) or anti-TRxR (1:2,000 dilution). Subsequently, after washing with TBST, the membranes were incubated with a horseradish peroxidase (HPR)-conjugated secondary antibody for 1 h. The protein bands were then visualized by chemiluminescence and exposure to X-ray film. Finally, the membranes were reprobed with antibody against glyceraldehyde 3-phosphate dehydrogenase (GAPDH) as an internal control. All visualized protein levels were quantified using the NIH ImageJ program (http://rsbweb.nih.gov/ij/).
Morphometric analysis. Digital images of the hematoxylin and eosin (H&E)-stained sections were obtained with a Zeiss AxioPlan microscope (Carl Zeiss MicroImaging, Inc., Thornwood, NY) using Axiovision software. The outer nuclear layer (ONL) thickness was measured at 400-µm intervals, starting at the optic nerve head and extending toward the superior and inferior ora serrata using Image-Pro Plus software (Media Cybernetics, Silver Spring, MD).

Dry-eye studies

Thirty 180–200-g female Lewis rats were divided into 5 groups; one served as an untreated control, while each of the other groups received 1 drop of Optixcare, Optixcare H, Optixcare EH, or I-Drop Vet Plus Ophthalmic Solution in each eye b.i.d. Dry eye was induced in all rats by subcutaneous injection of a 0.1 mL aqueous solution b.i.d. containing 24 mg of scopolamine (48 mg total dose per day). Tear flow was measured weekly prior to dosing using ZoneQuick™ Phenol Red Threads (FCI Ophthalmics, Pembroke, MA) by anesthetizing each rat with isoflurane. The lower eyelid was pulled down and, using forceps, the folded 3-mm end of each thread was placed into the palpebral conjunctiva at a point approximately one third of the distance from the lateral canthus. The string was left in the open eye for 15 s and then removed by gently pulling the lower eyelid down and removing the thread with an upward motion. The length of red-stained thread was then immediately measured. At the end of the study, all eyes were enucleated, and the grading of each lens was verified by examination of the subsequently dissected lens under a dissecting microscope. As summarized in Fig. 1, all young diabetic rats rapidly developed cataracts over an 8-week period, and this formation was inhibited by the topical administration of the aldose reductase inhibitor (ARI) Kinostat, which is currently undergoing a Food and Drug Administration (FDA)-approved clinical trial for the prevention of cataracts in diabetic dogs. The onset and progression of cataracts in these diabetic rats was delayed by the administration of nutraceuticals with the topical administration of Optixcare EH demonstrating a greater delay than the oral administration of the Ocu-GLO.

These slit-lamp observations were confirmed at the end of the study by microscopic examination of the dissected lenses (Fig. 2). While no lens opacities were evident in lenses from the ARI-treated rats, both the untreated and orally treated Ocu-GLO rats developed lens changes ranging from cortical to mature cataracts (Fig. 2). In contrast, primarily suture accentuation, which is the initial stage of sugar cataract development, was present in the Optixcare EH-treated diabetic rats. These differences in sugar cataract formation were not linked to differences in the severity of diabetes because all groups were similarly hyperglycemic over the course of this study, as indicated by their glycosylated hemoglobin levels (HbA1c, mean ± SEM untreated control 10.7 ± 0.7; Kinostat 11.48 ± 0.2; Optixcare EH 10.6 ± 0.7; Ocu-GLO 10.4 ± 0.76).

Another major risk factor of cataractogenesis is UV irradiation, which induces lens damage through the generation of ROS. Following the induction of diabetes with streptozotocin, lens changes were monitored weekly by hand-held slit lamp and graded on a scale of 0 (no lens change) to 3 (mature cataract). At the end of the study, all eyes were enucleated, and the grading of each lens was verified by examination of the subsequently dissected lens under a dissecting microscope. As summarized in Fig. 1, all young diabetic rats rapidly developed cataracts over an 8-week period, and this formation was inhibited by the topical administration of the aldose reductase inhibitor (ARI) Kinostat, which is currently undergoing a Food and Drug Administration (FDA)-approved clinical trial for the prevention of cataracts in diabetic dogs. The onset and progression of cataracts in these diabetic rats was delayed by the administration of nutraceuticals with the topical administration of Optixcare EH demonstrating a greater delay than the oral administration of the Ocu-GLO.

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of ROS that leads to decreased GSH levels and subsequent lens opacification. Experimental exposure of young male albino rats to UV irradiation results within 2–3 days led to a rapid decrease in lens GSH levels and the appearance of anterior lens vacuoles. For these studies, each rat was anesthetized and then placed on its side with only one eye exposed for 15 min to 1,600 mW/cm² of UV light (280–360 nm, UVmax = 306 nm), while the covered contralateral eye was down and away from UV exposure. In rats exposed to UV radiation by this method, lens GSH levels were significantly lower (P<0.05) in the exposed eyes compared with GSH levels in their unexposed contralateral lenses (Fig. 3). Similar irradiation of rats pretreated for 3 days with topical 3% JHX-4 b.i.d., Optixcare EH b.i.d., or oral Ocu-GLO administered daily by gavage resulted in GSH levels being significantly reduced only in the Ocu-GLO-treated rats. GSH levels in irradiated lenses from rats topically treated with either Optixcare EH or 3% JHX-4 were not significantly lower (Fig. 3). Nevertheless, all exposed eyes developed anterior vacuoles independent of treatment, and vacuoles were only absent in nonexposed eyes (Fig. 4).

To confirm that the reduced GSH levels after UV exposure were linked to increased levels of ROS, the lenticular levels of the oxidized membrane lipid 4-hydroxynonenol (4-HNE) were also measured. As summarized in Fig. 5, UV exposure resulted in an increase in 4-HNE levels, which were reduced by antioxidant treatments in the order: topical JHX-4 > topical Optixcare EH > oral Ocu-GLO. Compared with 4-HNE levels in lenses from untreated rats exposed to UV irradiation, 4-HNE levels were significantly lower in UV-exposed lenses from rats treated with JHX-4 (P<0.01) and Optixcare EH (P<0.05). 4-HNE levels in irradiated lenses from the Ocu-GLO-treated rats were not significantly different from levels in lenses from untreated rats exposed to UV radiation. 4-HNE lens levels from the UV exposed and unexposed contralateral eyes were statistically similar only in the JHX-4-treated rats.

Exposure of dark-adapted rats to light also results in the generation of ROS, iron dysregulation, and the release of iron from ferritin in the neural retina.46,47 In this animal model of AMD, Wistar rats were either orally treated with a diet containing 0.05% of the multifunctional antioxidant JHX-4 or topically treated b.i.d. with Optixcare EH with treatment beginning at the onset of 2-week dark adaption. As previously reported by Randazzo et al.,35 exposure of neural retinas to 1,000 lx of light for 3 h resulted in an eye was down and away from UV exposure. In rats exposed to UV radiation by this method, lens GSH levels were significantly lower (P<0.05) in the exposed eyes compared with GSH levels in their unexposed contralateral lenses (Fig. 3). Similar irradiation of rats pretreated for 3 days with topical 3% JHX-4 b.i.d., Optixcare EH b.i.d., or oral Ocu-GLO administered daily by gavage resulted in GSH levels being significantly reduced only in the Ocu-GLO-treated rats. GSH levels in irradiated lenses from rats topically treated with either Optixcare EH or 3% JHX-4 were not significantly lower (Fig. 3). Nevertheless, all exposed eyes developed anterior vacuoles independent of treatment, and vacuoles were only absent in nonexposed eyes (Fig. 4).

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increase of oxidative stress markers that include 4-HNE adducts formed with histidine following the nonenzymatic oxidation of polyunsaturated fatty acids and nitrotyrosine-modified proteins resulting from tyrosine residues being oxidized by peroxynitrite formed by the reaction of nitric oxide with superoxide anions. After light exposure, both of these markers significantly increased, while neither oxidative marker significantly increased in similar light-exposed rats treated with JHX-4 (Fig. 6). Rats treated with Optixcare EH showed partial increases in both oxidative markers. However, these increases were significantly less than those in untreated rats.

ROS and free radicals generated by retinal exposure to acute excessive light are also mediated by the thioredoxin (TRx)/thioredoxin reductase (TRxR) defense system. Therefore, the expression levels of both TRx and TRxR in the neural retinas of were also examined. Both TRx and TRxR expression increased in untreated rats exposed to light (Fig. 7). The expression of TRxR was significantly lowered \( (P<0.05) \) in the neural retinas of light-exposed rats treated with either JHX-4 or Optixcare EH. However, TRx expression was only reduced by JHX-4.

The reduction of oxidative stress by JHX-4 and Optixcare EH was confirmed by histological measurements of the photoreceptor layers, where intense light exposure resulted in photoreceptors thinning in the ONL, with damage in the superior hemisphere more pronounced than the inferior hemisphere. In rats treated with JHX-4 (Fig. 8A), the photoreceptor layer was only slightly decreased in both the inferior hemisphere and the superior hemisphere. In contrast, the photoreceptor layer was only partially protected in the Optixcare EH-treated rats (Fig. 8B). These results suggest that despite topical delivery, Optixcare EH can partially reduce oxidative stress in the retinas of light-damaged rats.

Optixcare is a carbomer-based eye lubricant that is used in the veterinary market to protect eyes from dehydration during anesthesia and as adjunctive support for dry eyes. Optixcare belongs to a class of ocular comfort agents that is composed of poly(acrylic acid) polymers that are designed to relieve ocular discomfort by augmenting the characteristics of the tear film so that the tear volume is stabilized and retained, while the ocular surface remains lubricated. Other such agents include hyaluronic acid (HA), which retains water. Based on this primary commercial use for Optixcare, we investigated whether the addition of nutraceutical antioxidants alter the ability of Optixcare to serve as an adjunctive treatment for dry eye. Dry eye was experimentally induced in adult Lewis rats by twice daily subcutaneous injections of scopolamine, and tear production was monitored with phenol red thread. As shown in Fig. 9, tear flow in each rat, determined as the average of both eyes, decreased over time in the untreated rats, and topical administration of either of the comfort agents Optixcare, Optixcare with HA, or i-Drop HA did not alter this decrease in tear flow. In contrast, tear flow was preserved in similar scopolamine-injected rats receiving the nutraceutical Optixcare EH. This
suggests that this nutraceutical formulation may actually be beneficial in dry eye by maintaining tear flow. The corneal surface was also clinically evaluated by slit lamp with Flurane/C212 stain prior to termination of the study. No sign of surface corneal or conjunctival erosion appeared to be present in eyes from either the nontreated or any of the treated groups, indicating that the scopolamine-induced dry eye was not clinically severe.

**Discussion**

Worldwide, nutraceuticals are being sold with the premise that they are beneficial in ameliorating age-related diseases such as cataracts, AMD, and dry eye. “Support” for their use is primarily based on theoretical mechanisms or in vitro “test tube” results rather than experimental evidence from in vivo animal studies. When clinical trials with nutraceutical antioxidants and vitamin supplements have been conducted, the results obtained have often been discouraging. For example, the Roche European American Cataract Trial (REACT), the Linxian cataract study, the AREDS study, Centrum use in the AREDS study, the α-tocopherol, β-carotene, and vitamin C prior to lens extraction showed no increase in intralenticular antioxidants despite substantially raised blood levels of these antioxidants. In contrast, topical application can result in increased lens levels as demonstrated by Ohta et al. who reported that lenticular levels of α-tocopherol can be raised by topical instillation of α-tocopherol containing liposomes and that this delays sugar cataract formation in rats.

The present studies suggest that topical administration of Optixcare EH, which contains 4 nutraceuticals suspended in a viscous carbomer vehicle that also contains 1% cetrimide as both a preservative and corneal permeation enhancer, may be beneficial in delaying lens changes in 2 rat cataract models associated with ROS. As anticipated in the diabetic rat model, topical application delayed the onset and progression of sugar cataracts. This is because osmotic changes from intracellular sorbitol formation are the primary initiator of sugar cataract formation in diabetic rats, and ROS generated through sorbitol-linked osmotic stress is only a secondary contributor to lens changes. This premise has been confirmed by a number of prevention studies in rats and dogs where adequate ARI administration prevents sugar cataract formation, while antioxidants only delay the onset and progression of sugar cataracts. This observation

![FIG. 6](image)

**FIG. 6.** Comparison of 4-HNE-histidine adduct levels (A) and nitrotyrosine adduct levels (B) in the neural retina of rats exposed to light after 2-week dark adaptation. HNE-histidine adduct levels, as determined by ELISA, in rats treated with or without the multifunctional antioxidant JHX-4 and Optixcare EH suggests that HNE levels were reduced by treatment. Similar conclusions were obtained with ELISA measurements of nitrotyrosine-modified proteins in rats treated with or without these antioxidants. The JHX-4 data have been previously published. Mean ± standard deviation (SD); n = 6. *P < 0.05 compared with the untreated, nonlight-damaged (NLD) group. #P < 0.05 compared with the untreated light damaged (LD) group. ELISA, enzyme-linked immunosorbent assay.

![FIG. 7](image)

**FIG. 7.** Comparison of expression of thioredoxin (TRx) (A) and thioredoxin reductase (TRxR) (B) in the neural retina of rats exposed to light after 2-week dark adaptation. Comparison of expression of TRx normalized to levels of glyceraldehyde 3-phosphate dehydrogenase (GAPDH) in rats treated with or without the multifunctional JHX-4 or topical Optixcare EH indicates that JHX-4 reduced the induction of TRx to a greater extent than Optixcare EH. Similarly, comparison of the expression of TRxR normalized to levels of GAPDH in rats treated with or without the multifunctional JHX-4 or topical Optixcare EH indicates that both JHX-4 and Optixcare reduced the induction of TRx compared with the levels observed in the untreated rats. The JHX-4 data have been previously published. Mean ± SD; n = 6. An asterisk (*) denotes a significant difference (P < 0.05) when compared to the untreated NLD group while the hatched symbol (#) denotes a significant difference (P < 0.05) compared to the untreated LD group.
has previously been reported by us by comparing the orally administered ARI AL1576 and orally administered multifunctional antioxidant JHX-4. The present study further confirms this premise by demonstrating that the topically administered ARI Kinostat, which is currently undergoing an FDA-approved clinical trial for the prevention of cataracts in diabetic dogs, also prevents cataract formation in rats, while the nutraceutical antioxidants Optixcare EH and Ocu-GLO delayed the onset and progression of sugar cataract development (Fig. 1). At the end of the study, little difference in cataract severity was observed between the untreated and orally Ocu-GLO-treated diabetic rats, while the topical Optixcare EH-treated rats primarily developed suture accentuation, which represents the initial stage of sugar cataract formation (Fig. 2). Many nutraceutical formulations also contain flavonoids that reduce AR activity. However, the ingredients in Optixcare EH do not directly inhibit AR. This suggests that the differences seen between Optixcare EH and Ocu-GLO are due to the strength of the antioxidant activities in the lens.

Exposure to UV light is a risk factor for cataracts. Experimental exposure to UV light also generates oxidative stress, which results in a decrease in reduced lens GSH levels and the subsequent development of lens opacification. In the present studies, rats with dilated eyes were then exposed to 1,600 μW/cm² of UV-B radiation at a distance of 20 cm for 15 min, while the contralateral eyes were covered and not exposed. This resulted in a significant decrease in the lenticular antioxidant levels of GSH and a significant increase in lens 4-HNE levels, a marker of lens lipid peroxidation products, in only the exposed eyes. Similar UV exposure of rats treated either orally or topically with the multifunctional antioxidant JHX-4 prevented these significant changes in GSH and 4-HNE. JHX-4 is a small molecule containing a unique 2-amino-5-hydroxyprymidine free radical scavenging ring system that readily reaches the lens and has been previously reported to delay the onset and progression of gamma-irradiation induced cataracts in rats. Topical administration of the ingredients in Optixcare EH to similar UV-exposed rats resulted in a similar, nonsignificant reduction of GSH levels compared with unexposed contralateral eyes and increases of 4-HNE levels that were significantly lower than those of similarly exposed eyes from untreated rats. In UV-exposed rats treated with Ocu-GLO, little antioxidant protection was observed. The decreases in GSH levels compared with their unexposed contralateral eyes were significant, while the 4-HNE levels compared with similarly UV-exposed eyes from untreated rats were not significantly lower. This suggests that the lenticular antioxidant levels were inadequately lower in the orally administered Ocu-GLO rats. Interestingly, none of the antioxidants afforded protection against the development of UV-induced anterior lens vacuoles.
Topical ocular administration of drugs generally does not result in adequate retinal levels of drug being obtained. Therefore, topical administration of Optixcare EH to light-exposed, dark-adapted rats was not anticipated to result in any significant protection of retinal oxidative stress compared with the oral administration of the multifunctional antioxidant JHX-4. Surprisingly, beneficial retinal effects of topically administered Optixcare EH were observed with 4-HNE histidine adduct levels and nitrotyrosine-modified protein levels being significantly reduced compared with untreated rats (Fig. 8) and some preservation of the ONL photoreceptor layer histologically observed (Fig. 9). However, compared with that of JHX-4, this protection was less, and Optixcare EH gave no protection against ERG changes (data not shown). This observation suggests that better retinal antioxidant protection by Optixcare EH may be achieved if the nutraceutical ingredients in Optixcare EH are also concomitantly administered orally.

Another age-related problem affecting the anterior segment is dry eye. This disease, which is also linked to oxidative stress and inflammation, is not only treated with immune suppressants but also with nutraceuticals such as combined antioxidants and omega-3 essential fatty acids formulations24 or topical epigallocatechin gallate25 and comfort agents such as viscous water-retaining carbomer gels and HA. Since Optixcare carbomer gel and Optixcare containing HA are used in the veterinary market for alleviating the clinical symptoms of dry eye, and the nutraceutical ingredients in Optixcare EH have been reported to demonstrate not only antioxidant but also anti-inflammatory activity, studies comparing the effects of these carbomer gel ingredients on rats with scopolamine-induced dry eye were investigated. Surprisingly, the nutraceutical Optixcare EH maintained tear flow in the dry eye–induced rats, while tear flow was not affected by either the administration of carbomer gel with HA or HA alone. Further studies are planned to determine the potential clinical effects of Optixcare EH on dry eye.

In summary, Optixcare EH is a carbomer gel that contains 4 nutraceuticals possessing antioxidant, anti-inflammatory, and chelating activity along with the preservative cetrimide, which enhances corneal permeability. In the limited prevention studies in rats reported here, topical application of this formulation protects against oxidative stress not only in the lens but to a lesser extent in the retina. Moreover, the formulation appears to maintain tear flow in an experimental dry-eye rat model. These results suggest that further clinical studies of Optixcare EH are warranted.

Author Disclosure Statement

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