The long-awaited medical glaucoma revival is finally here, fueled by the development of new medications and emerging drug delivery systems. Glaucoma medications have not seen a class innovation since the introduction of prostaglandin analogues (PGAs) in the 1990s. Research and development efforts have bestowed on us new drugs with novel mechanisms of action. Additionally, pharmaceutical companies are devising ways to improve drug delivery systems in order to create more efficacious, simple, and tolerable medications, with the ultimate goal of improving patient adherence.

Traditional glaucoma medications fit into several categories, including beta blockers, alpha agonists, carbonic anhydrase inhibitors, parasympathomimetics, and PGAs. These drug categories control IOP through different mechanisms: by (1) increasing aqueous humor outflow via the trabecular meshwork pathway, (2) increasing aqueous humor egress via the uveoscleral outflow pathway, or (3) decreasing production of aqueous humor.

Latanoprostene bunod ophthalmic solution 0.024% (Vyzulta, Bausch + Lomb) and netarsudil ophthalmic solution 0.02% (Rhopressa, Aerie Pharmaceuticals) are new antihypertensives that lower IOP via novel mechanisms of action.

**VYZULTA**

Vyzulta was approved by the FDA in November for the reduction of IOP in open-angle glaucoma and ocular hypertension. This once-daily medication has a dual mechanism of action, metabolizing into two components: latanoprost acid and butanediol mononitrate (Figure 1). Latanoprost acid, a prostaglandin F2-alpha analogue, increases uveoscleral outflow. Butanediol mononitrate breaks down into nitric oxide (NO), which effectually increases outflow through the trabecular meshwork and Schlemm canal.1

NO is a well-known mediator in smooth muscle relaxation and is found in various ocular tissues, including the trabecular meshwork and Schlemm canal.2 Studies demonstrate that, in eyes with open-angle glaucoma, there are lower levels of NO activity in the trabecular meshwork, Schlemm canal, and ciliary musculature than in normal eyes.3 In the phase 3 trials APOLLO and LUNAR, treatment with Vyzulta demonstrated a mean IOP reduction of 7.5 to 9.1 mm Hg from baseline between 2 and 12 weeks of treatment.4,5 No significant safety findings were reported, and side effects were similar to those associated with other PGAs.4,5

Figure 1. Vyzulta’s dual mechanism of action. Latanoprost acid increases uveoscleral outflow and butanediol mononitrate increases trabecular outflow via the release of nitric oxide.
Glaucoma was a reality for me from the time I was a child, when my maternal grandmother was diagnosed with the disease. Despite bilateral glaucoma surgery and chronic use of pilocarpine drops, her vision was exceedingly poor. I was flattered when she asked me to shop with her, always clutching my hand tightly. What I saw as her being protective was, in reality, a sign she was legally blind, probably with severe optic nerve atrophy, leaving her with limited central vision and a small tunnel field of vision.

From my teenage years on, I was determined to be an ophthalmologist. After medical school, residency, and fellowship, I decided to focus on cataracts, glaucoma, and corneal and external diseases. Due to my familial interest and research experience, I joined a private practice in 1987 as the office’s glaucoma specialist. In my first year of practice, I saw about 50 patients with guide dogs. The etiology for their legal or total blindness was exceedingly poor. I was flattered when she asked me to shop with her, always clutching my hand tightly. What I saw as her being protective was, in reality, a sign she was legally blind, probably with severe optic nerve atrophy, leaving her with limited central vision and a small tunnel field of vision.

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Hampering our diagnosis was the variability in results from Goldmann visual field (VF) testing, the gold standard diagnostic of that era. Depending on the size of the stimulus used, the skill and methodology of the examiner, and the patient’s endurance and frame of mind during the test, there was a high degree of variability. We instead depended on highly magnified, excellent-quality stereo disc photos and assigned a greater level of importance to changes noted there.

When drops were no longer a viable option, the next step was surgical intervention with trabeculectomy. Debates then focused on the optimal size and shape of the trapdoor, the desired size and elevation of the bleb, the benefits versus risks of antimetabolites such as 5-fluorouracil, and the use of devices such as a Simmons shell to help the bleb remain a certain size. A tougher decision, in patients with cataract, was whether to combine trabeculectomy with cataract surgery so that future cataract surgery would not interfere with the bleb. Failed blebs were dreaded. The rare case of endophthalmitis precipitated deep concern and an inclination to be surgically conservative so as to avoid this nightmarish complication.

While I was in training, beta blockers had gained an increasingly strong foothold; however, they were not used in private practice because primary care physicians were concerned about double-dosing patients who required a systemic antihypertensive. Eventually, this myth was dispelled, enabling ophthalmologists to prescribe timolol to all patients. Most patients were thrilled—especially those who had lived with the debilitating pilocarpine—thereby increasing patient compliance, enabling better glaucoma control, and significantly reducing the need for argon laser trabecuoplasty.

Although stereo disc photos have been reliable for decades, diagnostic testing has taken a quantum leap in the past 2 decades. Automated VF testing machines are now more accurate, reproducible, patient-friendly, and reliable. Devices such as the Humphrey Visual Field Analyzer (Carl Zeiss Meditec) are invaluable for earlier detection of field defects and monitoring response to therapy. Additionally, optic nerve testing with OCT has dramatically enhanced early glaucoma detection as well as monitoring the health of the optic nerve and retinal nerve fiber layer.

DEVELOPMENTS IN DROP THERAPY

Over the decades, additional classes of glaucoma eye drops have been introduced, including combination drops containing two medications with different mechanisms of action. The last of the classes to be introduced was the prostaglandin analogues (PGAs), with the first, latanoprost, in 1996. This class of drugs works by decreasing uveoscleral outflow to reduce IOP by 25% to 35%. PGAs had a meteoric rise in popularity because they are effective; are safe with only topical, localized side effects; and require only once-daily dosing.

In November, the FDA approved netarsudil mesylate (Rhopressa, Aerie Pharmaceuticals), a once-daily Rho-associated protein kinase (ROCK) inhibitor. Rhopressa works by increasing trabecular outflow through inhibition of the Rho pathway, thereby precipitating cytoskeletal changes to effect the relaxation of cells in the trabecular meshwork and Schlemm canal. Aerie is also expected to release a combination therapy containing netarsudil along with latanoprost (Roclatan), offering another dual-action option.

CONCLUSION

With OCT, automated VF machines, disc photos, and excellent topical glaucoma medications, glaucoma diagnoses and treatment have advanced light-years in my 3 decades of practice. I have gone from having 50 patients with guide dogs in 1987 to one single patient whose blindness is a result of congenital deformities, unrelated to glaucoma.
**RHOPRESSA**

Rhopressa was approved by the FDA in December for the reduction of IOP in open-angle glaucoma and ocular hypertension (Figure 2). This drug, dosed once daily, is a Rho kinase (ROCK) inhibitor that lowers IOP through three mechanisms of action: (1) increasing outflow via the trabecular meshwork, (2) decreasing production of aqueous humor, and (3) decreasing episcleral venous pressure. The ROCKET-1 and ROCKET-2 trials demonstrated noninferiority to timolol, with the most common adverse reaction being conjunctival hyperemia. Patients treated with once-daily dosing of Rhopressa experienced a reduction in IOP ranging from 3.9 to 4.1 mm Hg.

**DRUG DELIVERY SYSTEMS**

Attention has also been focused on improving patient adherence. One study found that nearly 45% of glaucoma patients used their once-daily drops less than 75% of the time. Improving drug delivery systems may be the solution to this problem. By providing a depot or long-term elution of medication, the burden of daily medication use may be mitigated.

Recent drug delivery innovations include Allergan’s bimatoprost sustained-release intracameral implant (Bimatoprost SR; Figure 3) and sustained-release bimatoprost ocular ring, Ocular Therapeutix’s sustained-release travoprost punctal plug (OTX-TP; Figure 4), and Glaukos’ sustained-release travoprost implant (iDose Travoprost; Figure 5).

**COMBINATION THERAPIES**

Patient adherence may also be improved by using combination drop therapy. Combigan (timolol/brimonidine 0.2%/0.5%, Allergan), Cosopt (dorzolamide-timolol ophthalmic solution 2%/0.5%, Akorn), and Simbrinza (brinzolamide/brimonidine 1.625%/0.5%, Allergan) are commercially available combination drop therapies with dosing frequency of twice or thrice daily. Compounding companies such as Ocular Science and Imprimis Pharmaceuticals now offer formulations that combine two, three, and four eye drops (timolol and lantanoprost; timolol, brimonidine, and dorzolamide; and timolol, brimonidine, dorzolamide, and lantanoprost), reducing the burden and preservative load typically associated with daily use of multiple drops.

**CONCLUSION**

This is an exciting time for the management of glaucoma. With the development and release of multiple medications with novel mechanisms of action and simple once-daily dosing, sustained-release medications in the not-too-distant pipeline, and two-to-four medication combination drop therapy options now available, the future of glaucoma management is brighter than ever.

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4. Weinreb RN, Storlazzi CT, Vittitow J, Ludwig J. Latanoprostene bunod 0.024% versus timolol maleate 0.5% in subjects with open-angle glaucoma or ocular hypertension: the APOLLO study. Ophthalmology. 2016;123(5):950-957.

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