Update on corticosteroids for diabetic macular edema

Abstract: Diabetic macular edema (DME) remains an important cause of visual loss. Although anti-vascular endothelial growth factor (VEGF) agents are generally used as first-line treatments for patients with center-involving DME, there is an important role for corticosteroids as well. Corticosteroids may be especially useful in pseudophakic patients poorly responsive to anti-VEGF therapies, in patients wishing to reduce the number of required injections, and in pregnant patients. Intravitreal triamcinolone acetonide has been used for many years but is not approved for this indication. An extended-release bioerodable dexamethasone delivery system and an extended-release nonbioerodable fluocinolone acetonide insert have both achieved regulatory approval for the treatment of DME. All intravitreal corticosteroids are associated with risks of cataract progression, elevation of intraocular pressure, and endophthalmitis. There is no current consensus regarding the use of corticosteroids, but they are valuable for selected patients with center-involving DME.

Keywords: diabetic macular edema, vascular endothelial growth factor, triamcinolone acetonide, dexamethasone, fluocinolone acetonide, randomized clinical trial

Introduction
Diabetic macular edema (DME) remains a major cause of vision loss in developed countries despite continued improvements in the care of both diabetes mellitus and diabetic retinopathy.1 The Early Treatment Diabetic Retinopathy Study reported that immediate focal/grid photocoagulation significantly decreases the likelihood of losing 15 or more letters of best corrected visual acuity (BCVA) due to “clinically significant” DME for up to 3 years.2 Based on these results, focal/grid photocoagulation became a standard treatment for DME, though recently introduced intravitreal pharmacotherapies have largely replaced photocoagulation for patients with center-involving DME.3

Drugs that inhibit the actions of vascular endothelial growth factor (anti-VEGF agents) have become consensus first-line therapy for the treatment of center-involving DME. In the pivotal Phase III registration trials, monthly intravitreal injections of ranibizumab (Lucentis®; Genentech, South San Francisco, CA, USA)4 or aflibercept (Eylea®; Regeneron, Tarrytown, NY, USA)5 improved BCVA significantly more than sham injections/photocoagulation. These trials convinced the US Food and Drug Administration (FDA) to approve each drug for the treatment of center-involving DME accompanied by loss in BCVA. In Phase II randomized clinical trials (RCTs), bevacizumab (Avastin®, Genentech)6 and pegaptanib (Macugen®, Bausch + Lomb, Bridgewater, NJ, USA)7 each improved DME, but neither drug has been approved by the FDA for this indication. The Diabetic Retinopathy Clinical Research Network (DRCR.net) performed an RCT and reported that aflibercept, bevacizumab,
and ranibizumab all effectively improve DME in most patients for at least 2 years.8,9

Most patients with DME improve with anti-VEGF treatment and many will also benefit from treatment with corticosteroids.10 The mechanism by which corticosteroids improve DME is unclear but several have been postulated. Corticosteroids stabilize and reconstitute the blood–retinal barrier,11 favorably alter the Starling’s equilibrium by constricting capillaries, and downregulate VEGF expression.12 Three fluorinated synthetic corticosteroids that lack mineralocorticoid activity – triamcinolone acetonide, dexamethasone phosphate, and fluocinolone acetonide – are currently used for the treatment of DME. They differ according to their glucocorticoid-receptor binding affinities (dexamethasone > triamcinolone > fluocinolone) and their lipophilicity (triamcinolone > fluocinolone > dexamethasone), characteristics that may partially explain their relative potencies (triamcinolone =5, dexamethasone =25, fluocinolone =25, compared to cortisol =1).

Corticosteroids also differ according to the proteins that they regulate. Triamcinolone, dexamethasone, and fluocinolone each upregulate over 6,000 proteins in two types of retinal pigment epithelial cells, but only 15%–25% of these proteins are upregulated by all the three corticosteroids. Within trabecular meshwork (TM) 86 cells, dexamethasone upregulates transcripts associated with RNA postranscriptional modifications, fluocinolone affects lipid metabolism, and triamcinolone affects cell morphology; within TM 93 cells, dexamethasone affects histone methylation, fluocinolone affects the cell cycle, and triamcinolone affects cell adhesion. None of the drugs undergoes intraocular metabolism and clearance from the vitreous proceeds according to first-order pharmacokinetics with elimination half-lives that range from hours (dexamethasone) to days (triamcinolone).13 To our knowledge, similar expression data have not been reported from retinal capillary endothelial cells.

The preferred method for the administration of corticosteroids in the treatment of DME is intravitreal injection. Systemic corticosteroids are associated with systemic risks including hyperglycemia. Topical and periocular corticosteroids do not achieve sufficient intraocular penetration to the posterior segment. Intraocular corticosteroids are effective but are associated with increased risks of posterior subcapsular cataract development14 and elevation of intraocular pressure (IOP).15 Severe16 and intractable17–18 elevation of IOP constitutes the major impediment to widespread steroid use. The cause of IOP elevation is unknown, but the contribution of genetic factors has long been suspected.19 A genome-wide association study linked two intergenic quantitative trait loci affecting a novel mucin gene, HCG22 (HLA complex group 22), with increased IOP following treatment with intravitreal triamcinolone acetonide.20

Intravitreal corticosteroids are also associated with increased risks of infectious endophthalmitis21 and noninfectious endophthalmitis.22 These two entities may appear similar clinically and distinguishing between them can be challenging.23 Infectious endophthalmitis may be more common with corticosteroids than with anti-VEGF agents. In a series of 406,380 injections, intravitreal triamcinolone acetonide and dexamethasone phosphate were associated with a significantly higher rate of endophthalmitis (0.13%) than were anti-VEGF agents (0.019%).24 Since most patients are treated with a series of intravitreal injections, the cumulative per-patient risk of endophthalmitis is higher than the per-injection risk. Reducing this lifelong cumulative risk by decreasing the number of injections represents one of the major benefits of extended-release drug preparations.25

**Triamcinolone acetonide**

Intravitreal triamcinolone acetonide, a slow-release crystal structure formulation, has been used to treat DME for many years,26–28 but it has been largely replaced by the lower risk anti-VEGF agents. Triamcinolone acetonide remains clinically active for 3 months following a single intravitreal injection.29 Triamcinolone is cleared more rapidly from eyes that have undergone vitrectomy, but the clinical responses in these eyes may still be favorable.30 A single-dose preparation of preservative-free triamcinolone acetonide (Triesence®; Alcon Laboratories, Inc., Fort Worth, TX, USA) has been approved by the FDA to enhance visualization of the vitreous during pars plana vitrectomy31 and to treat some posterior segment inflammatory diseases, but it has not been approved for the treatment of DME.

The DRCR.net Protocol B RCT compared intravitreal injections of triamcinolone acetonide 1 mg, triamcinolone acetonide 4 mg, and focal/grid photocoagulation for the treatment of DME. Treatment was allowed every 4 months for persistent or recurrent edema. At the 2-year primary temporal end point, mean BCVA was significantly better in the patients treated with photocoagulation. In the subgroup of patients with poor visual acuity at presentation (20/200–20/320), those treated with triamcinolone acetonide 4 mg (compared to those treated with photocoagulation) had better median improvements in BCVA (+21 vs +7 letters), and rates of ≥10-letter worsening (0% vs 17%) and ≥10-letter improvement (77% vs 42%).32
At 3 years in Protocol B, patients receiving focal/grid photocoagulation had greater mean improvement in BCVA than those receiving triamcinolone acetonide. The prevalence of cataract surgery was highest in patients treated with triamcinolone acetonide 4 mg (83%), followed by triamcinolone acetonide 1 mg (46%), and focal/grid photocoagulation (31%). IOP increases of ≥10 mmHg were most common among patients treated with triamcinolone acetonide 4 mg (33%), followed by triamcinolone acetonide 1 mg (18%), and focal/grid photocoagulation (4%).

The DRCR.net Protocol I RCT compared ranibizumab plus prompt or deferred photocoagulation with triamcinolone acetonide 4 mg plus prompt photocoagulation and with sham injection plus prompt photocoagulation. Retreatments were based on a complex, predetermined algorithm and the primary temporal end point was at 1 year. Ranibizumab plus either prompt or deferred photocoagulation produced significantly greater mean improvements in BCVA (+9 letters for each group) than did sham injection plus prompt photocoagulation (+3 letters) and triamcinolone acetonide plus prompt photocoagulation (+4 letters). Among patients who were pseudophakic at baseline, triamcinolone acetonide plus prompt photocoagulation produced a similar improvement in mean BCVA (+8 letters) as ranibizumab plus prompt (+8 letters) or deferred (+7 letters) photocoagulation. This suggests that the underperformance of the entire triamcinolone arm may have been due to the development of cataract.

At 2 years in Protocol I, the ranibizumab groups continued to enjoy better visual outcomes than the triamcinolone acetonide group. Patients in the triamcinolone arm were eligible to receive as-needed ranibizumab from 1.5 to 3 years (depending upon enrollment dates) into the trial, but by 5 years the BCVA improvements in this group still trailed those initially treated with ranibizumab.

The DRCR.net Protocol E RCT compared peribulbar triamcinolone acetonide with laser photocoagulation in patients with mild DME and BCVA of 20/40 or better. Patients were randomized to receive focal/grid photocoagulation, anterior sub-Tenon’s triamcinolone acetonide, anterior sub-Tenon’s triamcinolone acetonide plus photocoagulation, posterior sub-Tenon’s triamcinolone acetonide, or posterior sub-Tenon’s triamcinolone acetonide plus photocoagulation. Visual outcomes were similar in all the five groups, suggesting that peribulbar triamcinolone acetonide was ineffective.

Intravitreal triamcinolone acetonide effectively treats DME, but its poor overall performance in clinical trials can probably be attributed to the high incidence of cataracts. Triamcinolone acetonide is still used by many clinicians to treat DME, particularly in pseudophakic eyes, eyes unresponsive to anti-VEGF therapies, and eyes that present with poor visual acuity.

Dexamethasone

Off-label, intravitreal injections of dexamethasone phosphate are ineffective for DME, probably because of rapid clearance from the vitreous. An extended-release bioerodible dexamethasone delivery system (Ozurdex; Allergan, Inc., Irvine, CA, USA), however, has been approved by the FDA for the treatment of DME (Figure 1A and B), macular edema secondary to retinal vein occlusions, and noninfectious intermediate or posterior uveitis. Injection of the dexamethasone insert with a single-use, preloaded 22-gauge injector is performed in the outpatient clinic.

In the pivotal Phase III registration RCTs, patients received a 0.35 mg dexamethasone implant, a 0.7 mg dexamethasone implant, or a sham injection, with retreatments using predetermined criteria no more frequently than every 6 months. An improvement in visual acuity of at least 15 letters was reported in 21.5% of patients treated with the 0.7 mg implant, 18.4% treated with the 0.35 mg implant, and 12% treated with sham. In the subgroup of patients with DME previously treated with focal/grid photocoagulation, anti-VEGF agents, intravitreal triamcinolone acetonide, or a combination of these treatments, an improvement in visual acuity of at least 15 letters was reported in 21.5% of patients treated with the 0.7 mg implant versus 11.1% of patients treated with sham.

According to postmarketing clinical data, the dexamethasone delivery system is effective for 3–6 months in most patients. It is equally effective in eyes that have previously undergone vitrectomy and it was reported to be safe and effective in five pregnant patients.

Several adverse events, particularly cataract development and elevated IOP, are associated with the dexamethasone delivery system. Among patients treated with the 0.7 mg insert (the dose that was eventually approved by the FDA) in the 36-month registration trials, 41.5% required topical IOP lowering medications, but only 0.3% required incisional glaucoma surgery. The dexamethasone delivery system may migrate into the anterior chamber in pseudophakic or aphakic eyes, which requires surgical removal (Figure 2). Less frequently reported complications include endophthalmitis, acute retinal necrosis due to varicella zoster virus, and retinal and vitreous hemorrhage due to retinal impact during injection.

In a prospective study of eyes with DME, aqueous propermeability factors were measured before and after...
Fluocinolone acetonide

Two distinct extended-release formulations of fluocinolone acetonide have been approved by the FDA for intravitreal use: a larger (0.59 mg), nonbioerodable implant (Retisert®; Bausch + Lomb), which is implanted surgically through a scleral incision in an operating room; and a smaller (0.2 mg), nonbioerodable insert (Iluvien®; Alimera, Alpharetta, GA, USA), which is inserted with a single-use, preloaded 25-gauge injector in the outpatient clinic.

The larger surgical implant is approved for the treatment of chronic noninfectious posterior segment uveitis.

In a RCT conducted on patients with DME that persisted or recurred following focal/grid photocoagulation, the implant was compared against standard therapy (additional photocoagulation or observation). A BCVA improvement of three or more lines was more common in patients treated with the implant than in those receiving standard therapy at 6 months (16.8% vs 1.4%), 1 year (16.4% vs 8.1%), 2 years (31.8% vs 9.3%), and 3 years (31.1% vs 20%). However, the implant was associated with high rates of IOP elevation and cataract. IOPs of at least 30 mmHg were seen in 61.4% of implant eyes (vs 5.8% of standard therapy eyes), and 31.5%...
of implant patients underwent incisional glaucoma surgery within 4 years. Among phakic eyes, cataract extraction was performed in 91% (vs 20% of standard therapy eyes) within 4 years.\textsuperscript{58} Based on these disappointing results, the implant was not approved for the treatment of DME.

The smaller injectable insert has been approved by the FDA for the treatment of DME in patients treated previously with corticosteroids without a clinically significant increase in IOP,\textsuperscript{59} though neither the route of corticosteroid delivery nor the duration of treatment is specified on the label. A single insert is effective for up to 3 years.\textsuperscript{60}

In the pivotal Phase III registration RCTs, patients with persistent DME after focal/grid photocoagulation were randomized to receive a 0.2 \( \mu \)g/day insert, a 0.5 \( \mu \)g/day insert, or sham injection. At 2 years, 15-letter improvements in BCVA were achieved in 28.7% of patients treated with the 0.2 \( \mu \)g/day insert (which was eventually FDA approved), 28.6% with the 0.5 \( \mu \)g/day insert, and 16.2% with sham injection.\textsuperscript{61} Among patients receiving the 0.2 \( \mu \)g/day insert, 38.4% were treated with IOP-lowering medication (vs 14.1% of sham-treated patients) and 4.8% were treated with incisional glaucoma surgery (vs 0.5% of sham-treated patients) within 3 years.\textsuperscript{62} The rates of incisional glaucoma surgery in patients previously treated and not treated with triamcinolone acetonide were 0% and 6.1%, respectively,\textsuperscript{63} suggesting that restricting the implant to patients known to be nonresponders would improve outcomes and decrease complication rates. A review of fundus photographs obtained during the trials concluded that clinically significant glaucomatous optic nerve damage was not associated with IOP elevation.\textsuperscript{64}

Similar to the dexamethasone delivery system, the fluocinolone insert may migrate into the anterior chamber\textsuperscript{65} and require surgical repositioning.\textsuperscript{66}

**Summary**

Currently, surgeons have three intravitreal corticosteroid options for the treatment of DME: the dexamethasone delivery system, the fluocinolone acetonide insert, and off-label intravitreal triamcinolone acetonide (Table 1). All three agents, as well as the larger fluocinolone acetonide surgical implant, are associated with risks of cataract progression and IOP elevation (Table 2). There is no current consensus regarding the indications for intravitreal corticosteroids,\textsuperscript{67} but they are generally regarded as second-line agents for patients with center-involving DME who respond insufficiently to a series of anti-VEGF injections.

Intravitreal corticosteroids are more appropriate for pseudophakic eyes or in patients being considered for cataract surgery in the near future. Corticosteroids may be preferable to anti-VEGF agents in patients who wish to reduce the number of required injections and in pregnant patients who require treatment of the DME before delivery. As corticosteroid use expands and new data emerge from randomized, controlled

<table>
<thead>
<tr>
<th>Trial</th>
<th>Therapies studied</th>
<th>Summary of results</th>
</tr>
</thead>
<tbody>
<tr>
<td>DRCR.net Protocol B, 2-year results\textsuperscript{32}</td>
<td>Triamcinolone vs photocoagulation</td>
<td>Mean BCVA better in patients treated with photocoagulation</td>
</tr>
<tr>
<td>DRCR.net Protocol B, 3-year results\textsuperscript{33}</td>
<td>Triamcinolone vs photocoagulation</td>
<td>Greater mean improvement in BCVA in patients treated with photocoagulation</td>
</tr>
<tr>
<td>DRCR.net Protocol I, 1-year results\textsuperscript{34}</td>
<td>Triamcinolone plus photocoagulation vs ranibizumab plus photocoagulation vs sham injection plus photocoagulation</td>
<td>Ranibizumab plus photocoagulation produced greatest mean improvements in BCVA (in pseudophakic eyes, ranibizumab similar to triamcinolone)</td>
</tr>
<tr>
<td>DRCR.net Protocol I, 2-year results\textsuperscript{35}</td>
<td>Triamcinolone plus photocoagulation vs sham injection plus photocoagulation</td>
<td>Ranibizumab associated with better visual outcomes than triamcinolone</td>
</tr>
<tr>
<td>DRCR.net Protocol I, 5-year results\textsuperscript{36}</td>
<td>Triamcinolone plus photocoagulation vs sham injection plus photocoagulation</td>
<td>Ranibizumab associated with better visual outcomes than triamcinolone</td>
</tr>
<tr>
<td>Boyer et al\textsuperscript{44}</td>
<td>Dexamethasone insert vs sham injection</td>
<td>15-letter improvement in BCVA: 22.2% with dexamethasone 0.7 mg, 18.4% with dexamethasone 0.35 mg, 12% with sham</td>
</tr>
<tr>
<td>Augustin et al\textsuperscript{45}</td>
<td>Dexamethasone insert vs sham injection – subgroup with previous other treatments</td>
<td>15-letter improvement in BCVA: 21.5% with dexamethasone 0.7 mg, 11.1% with sham</td>
</tr>
<tr>
<td>Pearson et al\textsuperscript{46}</td>
<td>Fluocinolone surgical implant vs standard therapy</td>
<td>Three-line improvement in BCVA vs standard therapy: 16.8% vs 1.4% at 6 months, 16.4% vs 8.1% at 1 year, 31.8% vs 9.3% at 2 years, 31.1% vs 20% at 3 years</td>
</tr>
<tr>
<td>Campochiaro et al\textsuperscript{47}</td>
<td>Fluocinolone insert vs sham injection</td>
<td>15-letter improvement in BCVA at 2 years: 28.7% with 0.2 ( \mu )g/day insert, 28.6% with 0.5 ( \mu )g/day insert, 16.2% with sham</td>
</tr>
</tbody>
</table>

**Abbreviations:** BCVA, best corrected visual acuity; DRCR.net, Diabetic Retinopathy Clinical Research Network.
clinical trials, a consensus regarding their optimal use may ultimately emerge.

Acknowledgment
This article was partially supported by the NIH Center Core Grant P30EY014801 and Research to Prevent Blindness Unrestricted Grant to the University of Miami.

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

References


