The efficacy and safety outcomes of the 0.19 mg fluocinolone acetonide implant after prior treatment with the 0.7 mg dexamethasone implant in patients with diabetic macular edema

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Purpose: There are little or no published data comparing the outcomes of ILUVIEN® (0.19 mg fluocinolone acetonide [FAc]) and OZURDEX® (0.7 mg dexamethasone [DEX]) implants in patients with diabetic macular edema (DME), and this case sought to compare their outcomes.

Methods: This case was extracted from a monocentric audit involving a pool of 25 patients (33 eyes) with DME and treated with a single FAc implant between October 2013 and December 2016. This case, a 61-year-old male with a pseudophakic lens, is from a patient that had received 4 intravitreal injections of a DEX implant prior to FAc implant and then was monitored for 3 years until re-treatment with a second FAc implant. Parameters measured included visual acuity (VA), central retinal thickness (CRT), and intraocular pressure (IOP).

Results: After the DEX implants, CRT transiently improved. In March 2014, the decision was taken to administer an FAc implant, and this led to a reduction in CRT below 300 µm (from a baseline of 748 µm), and this was sustained for 30 months. VA remained above 65 Early Treatment Diabetic Retinopathy Study letters to month 36, after which time a second FAc implant (in April 2017) was administered due to recurrence of edema and CRT decreased to below 300 µm and VA improved to 70 letters. Side effects included elevated IOP, which was effectively managed with IOP-lowering drops.

Conclusion: A single injection of FAc implant led to sustained improvements in CRT and VA that lasted for between 30 and 36 months, which is in contrast to the DEX implant where re-treatment was generally required within 6–7 months. After 36 months, re-treatment with the FAc implant again led to improved VA and CRT, and responses that were similar to those achieved with the first FAc implant.

Keywords: fluocinolone acetonide, diabetic macular edema, microdosing

Background

The diagnosis of diabetes continues to grow, from 388 million sufferers in 2013 to a projected 592 million in 2030, with an expected corresponding increase in the complications of diabetes.¹ Diabetic retinopathy is a common visual complication in the diabetic population, estimated to affect 30% of patients¹ and around 10% of the prevalent population have vision-threatening states such as diabetic macular edema (DME) or proliferative diabetic retinopathy.¹ This therefore represents a significant socioeconomic cost for health care systems across the globe and highlights the need for developing new treatment strategies. On the counter side of this argument, however, it also means that health care providers need to be selecting the optimal therapy for treating their patients as well as using any therapy cost-effectively.²
In DME, the main treatment modalities include laser photocoagulation, intravitreal anti-VEGFs, and intravitreal corticosteroids or corticosteroid implants. In the last decade, the use of intravitreal anti-VEGFs or corticosteroids has revolutionized DME management with the emergence of randomized clinical trials showing these therapies led to a significant proportion of patients experiencing an improvement in both visual acuity (VA) and retinal anatomy.4,5

The ILUVIEN (fluocinolone acetonide [FAc]) implant received its first European medical license approval in 2012 and is therefore a relatively recent development for the treatment of DME, where it is indicated for the treatment of vision impairment associated with chronic DME considered insufficiently responsive to available therapies (ie, for the treatment of persistent or recurrent DME despite treatment).6 The evidence for the efficacy and safety of FAc implant comes from the Fluocinolone Acetonide for Diabetic Macular Edema (FAME) trials in which the FAc implant, which provides a daily microdose of fluocinolone acetonide, was shown to provide a substantial visual benefit that lasted for up to 3 years.7

Since its approval in 2012, there have been a growing number of patients completing 3 years of therapy, and this has been observed at recent conferences such as Association for Research in Vision and Ophthalmology (ARVO) in Baltimore between the 7th and 11th May, 2017.8 The current case was presented at ARVO 2017,9 and the current article presents the patient outcomes.

Case presentation
A 61-year-old male with type 2 diabetes mellitus was diagnosed with DME on January 5, 2009 (Table 1). Comorbidities included hypertension and dyslipidemia. Between May 2009 and March 2010, the patient underwent 3 core vitrectomy surgeries, and was concurrently being treated with anti-VEGF (Avastin) and short-acting steroids (DEX implant and intravitreal injection of triamcinolone [IVTA]). On 2 separate occasions in March and April 2010, the patient was treated with laser focal therapy. Phacoemulsification was performed in August 2010 before a fourth core vitrectomy surgery was performed in December 2010. Ranibizumab was administered twice (March 2011 and April 2011) before the patient was treated with his first DEX implant in October 2011 (Table 2 and Figure 1).

Core vitrectomy or 2 ports pars plana vitrectomy is used at the University Eye Clinic of Frankfurt not only to remove vitreous opacities, but also to inject a combination of different drugs into the vitreous cavity without any volume restrictions and without risk of vision-threatening intraocular pressure (IOP) rise. Furthermore, the core vitrectomy itself serves other purposes as it allows a posterior vitreous detachment to be achieved under visual control, which is prognostically important for

| Date (DD/MM/YYYY) | DME treatment | CRT and VA values* |
|-------------------|---------------|---------------------|
| 05/01/2009        | DME diagnosed |                     |
| 22/05/2009        | Core vitrectomy + avastin + DEX implant + triamcinolone |                     |
| 18/09/2009        | Core vitrectomy + avastin + DEX implant + triamcinolone |                     |
| 16/03/2010        | Core vitrectomy + avastin + DEX implant + triamcinolone |                     |
| 16/03/2010        | Focal laser   |                     |
| 26/04/2010        | Focal laser   |                     |
| 13/08/2010        | Phacoemulsification, implantation of PC-IOL + avastin |                     |
| 17/12/2010        | Core vitrectomy + avastin + DEX implant + triamcinolone + cryocoagulation |                     |
| 09/03/2011        | Ranibizumab   |                     |
| 04/04/2011        | Ranibizumab   |                     |
| 10/10/2011        | DEX implant   | 622 µm, 60 ETDRS letters |
| 22/05/2012        | DEX implant   | 676 µm, 50 ETDRS letters |
| 15/11/2012        | DEX implant   | 592 µm, 60 ETDRS letters |
| 07/01/2013        | Panretinal photocoagulation |                     |
| 29/01/2013        | Panretinal photocoagulation |                     |
| 01/03/2013        | Panretinal photocoagulation |                     |
| 09/08/2013        | DEX implant   | 760 µm, 60 ETDRS letters |
| 17/03/2014        | FAc implant   | 748 µm, 50 ETDRS letters |

Notes: *Please see Figure 1 where CRT and VA values for the DEX and FAc implants are plotted against time.
Abbreviations: CRT, central retinal thickness; DME, diabetic macular edema; DEX, dexamethasone; ETDRS, Early Treatment Diabetic Retinopathy Study; FAc, fluocinolone acetonide; IOL, intraocular lens; VA, visual acuity.
Intravitreal steroid injections

diabetic patients. Repeated core vitrectomy allows for the serial washout of proinflammatory molecular mediators from the vitreous cavity with the objective of achieving optimal effects with intravitreal injected pharmacological agents.

On the date the first DEX implant was injected, VA was 60 letters converted from Snellen fraction to an ETDRS letter score and CRT was 622 microns (Table 1). A second and third implant were administered in May 2012 (50 letters and

Table 2 CRT, VA, and OCT images following the administration of steroid implants to the left eye

| Date       | CRT (microns) | VA (letters) | Steroid implant | OCT image |
|------------|---------------|--------------|-----------------|-----------|
| 10/10/2011 | 622           | 60           | DEX implant     |           |
| 17/01/2012 | 397           | 60           |                 |           |
| 22/05/2012 | 676           | 50           | DEX implant     |           |
| 22/06/2012 | 257           | 60           |                 |           |
| 15/11/2012 | 592           | 60           | DEX implant     |           |
| 09/08/2013 | 760           | 60           | DEX implant     |           |
| 29/11/2013 | 424           | 60           |                 |           |
| 16/01/2014 | 588           | 50           |                 |           |
| 17/03/2014 | 748           | 50           | FAc implant     |           |
| 14/10/2014 | 176           | 65           |                 |           |
| 17/03/2015 | 258           | 70           |                 |           |
| 09/09/2015 | 205           | 70           |                 |           |
| 05/04/2016 | 236           | 65           |                 |           |
| 11/10/2016 | 221           | 65           |                 |           |
| 11/04/2017 | 616           | 65           | FAc implant     |           |
| 14/09/2017 | 221           | 70           |                 |           |

Abbreviations: CRT, central retinal thickness; DEX, dexamethasone; FAc, fluocinolone acetonide; OCT, optical coherence tomography; VA, visual acuity.

Figure 1 VA and CRT over time following treatment with repeated injections of the DEX (n=4) and FAc implant (n=2).

Notes: Intravitreal injections of DEX and FAc implants are shown in green and blue, respectively. The patient was re-treated with a second FAc implant in April 2017.

Abbreviations: CRT, central retinal thickness; DEX, dexamethasone; ETDRS, Early Treatment Diabetic Retinopathy Study; FAc, fluocinolone acetonide; OCT, optical coherence tomography; VA, visual acuity.
676 microns) and November 2012 (60 letters and 592 microns) at which points VA remained relatively stable irrespective of treatment, and recurrence of edema was seen within 6–7 months of the DEX implant being injected (Table 2 and Figure 1). Between January 2013 and March 2013, panretinal photocoagulation was administered 3-times before a fourth DEX implant was injected in August 2013. CRT at this point was 760 microns and VA still remained unchanged (60 letters).

In March 2014, the first FAc implant was injected after CRT had again rebounded within 7 months of the fourth DEX implant being given (to 748 microns) and VA had started to worsen (50 letters). Figure 1 highlights the impact of FAc implant in this patient with CRT remaining below 300 microns until October 2016 (30 months) and VA being stable between 65 and 70 letters. After 3 years, edema had recurred (616 microns) and the patient was re-treated with a second FAc implant and the macula was dry in September 2017 and VA had improved further to 70 letters.

No additional therapies were given in combination with FAc implant. Side effects included the elevation of IOP, which was effectively managed with IOP-lowering drops, and remained below 21 mmHg.

**Conclusion**

DEX and FAc implants led to similar improvements in peak CRT and VA, with the main difference between therapies being the sustained duration of effect lasting up to 36 months with a single FAc implant. After 30 months, the first signs of recurrent edema were observed, but without change in VA, indicating that re-treatment with a second implant was required in this patient. After injection of a second FAc implant, CRT markedly improved along with a slight improvement in VA to 70 letters. This improvement has a direct benefit to the patient as this is the VA required to hold a driving license in Europe. It is also notable that the improvements in VA and CRT were achieved after extensive DME prior therapies, including repeated injections of the short-acting DEX implant. Indeed, the control achieved with the DEX implant was only maintained for up to 6 or 7 months (Figure 1) before CRT rebounded to values ≥600 microns. In stark contrast, FAc implant led to sustained drying of the macula and sustained/improved VA to levels not achieved with the previous treatments with the DEX implant. This observation needs to be confirmed in a larger group of vitrectomized patients and also in non-vitrectomized patients.

This case clearly shows that the patient had been heavily treated with therapies (laser, intravitreal injections, and vitrectomy) prior to being treated with the FAc implant (Table 1). The patient history, therefore, clearly shows they were not treatment naïve and had persistent or recurrent DME despite receiving prior treatments. It also raises questions as to the effect of these prior treatments on the outcomes with the FAc implant and whether these outcomes could be further optimized by earlier treatment in the disease process, as has been suggested in recent publications. In the current case, the patient received 4 injections of the DEX implant prior to FAc implant, and this was because the FAc implant was only launched in Europe in 2013 and was not commercially available. The current license means that FAc implant can be used once an insufficient response to a first-line therapy, predominantly the injection of anti-VEGF drugs, has been established. This means that the clinician can decide whether to switch directly from a first-line therapy to FAc implant or whether they want to confirm the patient is responding to a short-acting steroid before administering the long-acting steroid, FAc.

**Ethics**

This article does not contain any new studies with human or animal subjects performed by any of the authors. Written informed consent has been provided by the patient to have the case details published.

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**Disclosure**

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