REVIEW

Safety and Efficacy of Dexamethasone Intravitreal Implant Given Either First-Line or Second-Line in Diabetic Macular Edema

Andrea Taloni^{1,*}, Giulia Coco ^{2,*}, Davide Rastelli³, Giacinta Buffon², Vincenzo Scorcia ¹, Giuseppe Giannaccare⁴

¹Department of Ophthalmology, University "Magna Graecia" of Catanzaro, Catanzaro, Italy; ²Department of Clinical Sciences and Translational Medicine, University of Rome Tor Vergata, Rome, Italy; ³Department of Ophthalmology, Policlinico Casilino, Rome, Italy; ⁴Eye Clinic, Department of Surgical Sciences, University of Cagliari, Cagliari, Italy

*These authors contributed equally to this work

Correspondence: Giuseppe Giannaccare, Full Professor of Ophthalmology, Eye Clinic, Department of Surgical Sciences, University of Cagliari, Via Università 40, Cagliari, 09124, Italy, Tel +0039 09613647041, Fax +0039 09613647094, Email giuseppe.giannaccare@gmail.com

Abstract: Diabetic macular edema (DME) is a common sight-threatening complication of diabetic retinopathy (DR) and the leading cause of severe visual impairment among the working-age population. Several therapeutic options are available for the management of DME, including intravitreal corticosteroids. They have been traditionally used as second-line treatment, due to the risk of intraocular pressure increase and cataract-related adverse events. However, attention has recently been focused on the primary or early use of intravitreal corticosteroids, due to growing evidence of the crucial role of inflammation in the pathogenesis of DME. Furthermore, intravitreal steroid implants offer the additional advantage of a longer duration of action compared to anti-vascular endothelial growth factor agents (anti-VEGF). This review aims to summarize the available evidence on the efficacy and safety profile of dexamethasone (DEX) intravitreal implant, with a specific focus on clinical scenarios in which it might be considered or even preferred as first-line treatment option by adequate selection of patients, considering both advantages and possible adverse events. Patients with contraindications to anti-VEGF, DME with high inflammatory OCT biomarkers, pseudophakic patients and phakic patients' candidates to cataract surgery as well as vitrectomized eyes may all benefit from first-line DEX implant. Additionally, DME not responders to anti-VEGF should be considered for a switch to DEX implant and a combination therapy of DEX implant and anti-VEGF could be a valid option in severe and persistent DME. **Keywords:** diabetic macular edema, DME, dexamethasone intravitreal implant, DEX implant, corticosteroids, Ozurdex

Introduction

Diabetic retinopathy (DR) affects about one-third of diabetic population and represents the main cause of visual disability and blindness among the working-age population in developed countries.^{1,2} Among sight-threatening complications of DR, diabetic macular edema (DME) is the most frequent cause of vision loss, with an estimated global prevalence among diabetic patients varying between 1.4% and 12.8%, influenced by the type of diabetes, race, ethnicity, and disease duration.³ Diabetic macular edema is characterized by the accumulation of exudative fluid intraretinally and, less commonly, subretinally, with consequent retinal thickening within the macular area.⁴ This condition can cause decreased vision, metamorphopsia, impaired color perception and, if prolonged, visual loss due to irreversible retinal damage.

Several therapeutic options are currently available for the management of DME, including intravitreal anti-vascular endothelial growth factor (VEGF) agents, intravitreal corticosteroids, laser photocoagulation and surgery. In 2017, the European Society of Retina Specialists (EURETINA) guidelines recommended intravitreal anti-VEGF agents as first-line treatment in patients with DME, and intravitreal steroids as second-choice treatment.⁵ Indeed, although many studies demonstrated the non-inferiority of intravitreal dexamethasone (DEX) implant (Ozurdex[®]; Allergan, Inc., Irvine, CA, United States) compared to anti-VEGF agents in terms of efficacy, the former is considered a second-line choice for

© 2023 Taloni et al. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/terms. work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission form Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, pisse see paragraphs 4.2 and 5 of our Terms (https://www.dovepress.com/terms.php). DME, mainly due to the relatively higher risk of ocular side effects, particularly intraocular pressure (IOP)-related and cataract-related adverse events (AEs).⁵ However, it was also pointed out that intravitreal steroids may be the first-line option in patients with a history of recent cardiovascular events, pregnant or breast-feeding women as well as those not keen on undergoing monthly injections.⁵ In addition, recent literature focused on specific subgroups of patients aiming to identify those who might benefit more from early or first-line treatment with steroids, for instance vitrectomized eyes, pseudophakic eyes and/or non-compliant patients.^{6–8}

Currently, DEX and fluocinolone acetonide (FA) implants are available and approved for intravitreal administration in DME. DEX intravitreal implant has been developed first and its use in patients with DME was approved by FDA and EMA in 2014. To date, it represents the first option when an intravitreal steroid is needed.⁶

The purpose of this review is to describe the efficacy and safety profile of DEX intravitreal implant, highlighting clinical scenarios in which it might be considered or even preferred as first-line treatment option.

Methods

Literature search was conducted on PubMed, Web of Science, Scopus, and Google Scholar using the following keywords: diabetic macular edema/oedema or DME/DMO; dexamethasone implant; DEX implant; Ozurdex; intravitreal steroids; intravitreal steroid implant; switching; poor response; cataract surgery; vitrectomized eyes; and combinations of them.

Randomized controlled trials (RCTs), large real-life studies, meta-analyses and systematic reviews assessing efficacy and safety of intravitreal DEX implant were included in this review. With regard to specific topics for which RCT was not available, for instance the comparison of DEX implant in vitrectomized versus non-vitrectomized eyes, prospective and retrospective studies were also included. The references of the examined articles were further investigated for additional publications. Only articles published in English were considered for our review (Figure 1). All authors had the

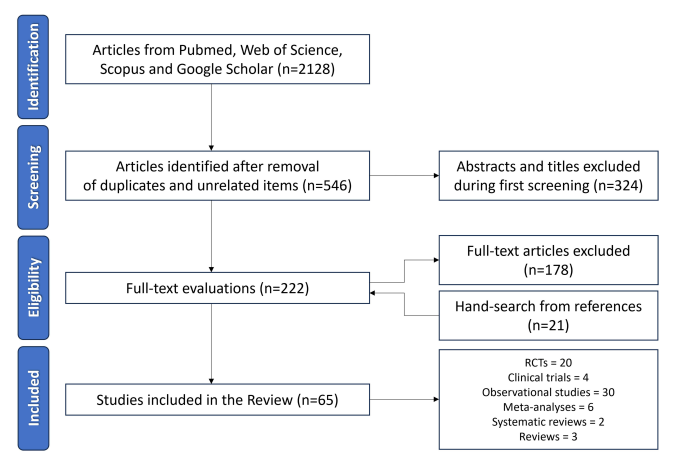


Figure I Preferred Reporting Items for a Systematic Review and Meta-Analyses (PRISMA) flowchart for the article selection process.

opportunity to make suggestions and comments to achieve a consensus on the collected data, in agreement with the literature evidence. A summary of results from clinical trials involving patients affected by DME who received DEX implant has been reported in Table 1. The risk of bias assessment has been summarized in Figure 2.

Table I Design and Results of Clinical Studies Involving Patients Affected by Diabetic Macular Edema Who Received Dexamethasone
Intravitreal Implant

Study (First Author, RCT Name, Year)	Patients	Study Design and Study Groups	Follow-Up	Results Numerical Values Refer to the End of the Study Follow-Up Unless Stated Otherwise				
RCTs supporting	RCTs supporting the use of DEX in DME							
Kuppermann et al (2007) ⁹	315 ME pretreated with laser or medical therapy	Study Design: Phase 2 RCT Multicenter Double-masked Groups: 105 DEX 0.7 mg 105 DEX 0.35 mg 105 observation	6 months	BCVA improvement \geq 15 letters 18.1% DEX 0.7 mg (p<0.05 vs observation) 14.6% DEX 0.35 mg (p=0.02 vs observation) 7.6% observation Mean change in CRT -106.7 µm DEX 0.7 mg (p=0.02 vs observation) -53.2 µm DEX 0.35 mg (p<0.001 vs observation) +20.7 µm observation				
Haller et al (2010) ¹⁰	171 DME pretreated with laser or medical therapy	Study Design: Phase 2 RCT Multicenter Single-masked Sham-controlled Groups: 57 DEX 0.7 mg 57 DEX 0.35 mg 57 Observation	6 months	BCVA improvement \geq 10 letters 30% DEX 0.7 mg 19% DEX 0.35 mg 23% observation (p=0.4 vs DEX groups) Mean change in CRT -132.3 µm DEX 0.7 mg (p<0.001 vs observation) -42.6 µm DEX 0.35 mg (p=0.07 vs observation) +30.2 µm observation				
Boyer et al, MEAD (2014) ¹¹	1048 DME treatment naïve, or pretreated with laser or medical therapy	Study Design: Two Phase 3 RCT Multicenter Single-masked Sham-controlled Groups: 351 DEX 0.7 mg 347 DEX 0.35 mg 350 Sham	3 years	BCVA improvement ≥ 15 letters 22.2% DEX 0.7 mg 18.4% DEX 0.35 mg 12.0% Sham (p≤0.018 vs DEX groups) Mean change in CRT -111.6 μm DEX 0.7 mg -107.9 μm DEX 0.35 mg -41.9 μm Sham (p<0.001 vs DEX groups)				
Callanan et al, PLACID (2013) ¹²	253 DME pretreated with laser or medical therapy	Study Design: Phase 2 RCT Multicenter Double-masked Sham-controlled Groups: 126 DEX + laser 127 Sham + laser	12 months	$\begin{array}{l} \textbf{BCVA improvement} \geq \textbf{10 letters} \\ 27.8\% \ \text{DEX + laser} \\ 23.6\% \ \text{Sham + laser} \\ (p=0.453) \\ \textbf{Mean change in BCVA} \\ \text{DEX + laser > Sham + laser (p<0.001)} \\ \textbf{Mean change in area of vascular leakage} \\ \text{DEX + laser > Sham + laser (p<0.0041)} \end{array}$				

Study (First Author, RCT Name, Year)	Patients	Study Design and Study Groups	Follow-Up	Results Numerical Values Refer to the End of the Study Follow-Up Unless Stated Otherwise
Heng et al, OZLASE (2016) ¹³	80 DME Treatment naïve	Study Design: RCT Single-site Groups: 40 DEX + laser 40 laser	56 weeks	BCVA improvement \geq 10 letters 18.4% DEX + laser 20.5% laser (effect estimate 1.15) Mean change in CST -113 μ m DEX + laser -17 μ m laser (p<0.001)
Wei et al (2021) ¹⁴	284 DME no laser treatment or anti-VEGF injections in the previous 3 months	Study Design: RCT Single-site Open-label Patients: Groups: 145 DEX 0.7 mg 127 laser	12 months	Mean change in BCVA +4.3 letters DEX +1.4 letters laser (p<0.001) Mean change in CRT -209.5 μm DEX -120.3 μm laser (p<0.001) Mean change in total leakage area -8.367 mm ² DEX -0.637 mm ² laser (p<0.001)
Gillies et al, BEVORDEX (2014) ¹⁵	anti-VEGF injections 88 DME pretreated with laser	Study Design: Phase 2 RCT Single-masked Multicenter Sham-controlled Groups: 42 BEV 1.25 mg 46 DEX 0.7 mg	12 months	$\begin{tabular}{ c c c } \hline \mbox{Mean number of injections} \\ \hline \mbox{8.6 BEV} \\ \hline \mbox{2.7 DEX} \\ \hline \mbox{BCVA improvement} \geq 10 \ \mbox{letters} \\ \hline \mbox{40\% BEV} \\ \hline \mbox{41\% DEX} \\ \hline \mbox{(p=0.83)} \\ \hline \mbox{Mean change in CMT} \\ \hline \mbox{-122 } \mbox{μm BEV} \\ \hline \mbox{-187 } \mbox{μm DEX} \\ \hline \mbox{(p=0.015)} \\ \hline \end{tabular}$
Callanan et al (2017) ¹⁶	363 DME no laser treatment or anti-VEGF injections in the previous 3 months	Study Design: Phase 2 RCT Multicenter Groups: 181 DEX 0.7 mg 182 RAN 0.5 mg	I2 months	Mean number of injections 2.85 DEX 8.70 RAN Mean change in BCVA +4.34 letters DEX +7.60 letters RAN (noninferiority of DEX) Mean change in CRT -173.9 μm DEX -163.5 μm RAN (p>0.05)

Table I (Continued).

Study (First Author, RCT Name, Year)	Patients	Study Design and Study Groups	Follow-Up	Results Numerical Values Refer to the End of the Study Follow-Up Unless Stated Otherwise
Sharma et al (2020) ¹⁷	40 DME no anti-VEGF injections in the previous 4 months	Study Design: RCT Groups: 20 BEV 1.25 mg or RAN 0.5 mg 20 DEX 0.7 mg	6 months	$\label{eq:second} \begin{array}{l} \mbox{Mean number of injections} \\ 4.04 & \mbox{BEV/RAN} \\ 1.15 & \mbox{DEX} \\ \mbox{BCVA improvement} \geq 10 & \mbox{letters} \\ 50\% & \mbox{BEV/RAN} \\ 60\% & \mbox{DEX} \\ \mbox{Mean change in BCVA (month 3)} \\ -0.31 & \mbox{LogMAR BEV/RAN} \\ -0.42 & \mbox{LogMAR DEX} \\ (p=0.27) \\ \mbox{Mean change in CFT (month 3)} \\ -177 & \mbox{BEV/RAN} \\ -227 & \mbox{m DEX} \\ (p=0.02) \end{array}$
Ozsaygili et al (2020) ¹⁸	98 DME treatment naïve	Study Design: RCT Single-site Single-masked Groups: 48 DEX 0.7 mg 50 AFL 2.0 mg	12 months	Mean number of injections 2.6 DEX 7.2 AFL BCVA improvement \geq 10 letters 25% DEX 42% AFL (p=0.058) Mean change in BCVA +6.4 letters DEX +9.3 letters AFL (p<0.01) Mean change in CRT -318 µm DEX -209 µm AFL (p<0.001)
Comet et al, INVICTUS (2020) ¹⁹	60 DME treatment naïve	Study Design: Prospective Single-site Open-label Groups: 19 RAN 0.5 mg 20 AFL 2.0 mg 21 DEX 0.7 mg	12 months	Mean number of injections 7.1 RAN 6.9 AFL 2.6 DEX BCVA improvement \geq 15 letters 32% RAN 33% AFL 30% DEX (p>0.05) Mean change in BCVA +7.2 letters RAN +8.5 letters AFL +9.4 letters DEX (p=0.89) Mean change in CRT -99.4 µm RAN -101.7 µm AFL -162.7 µm DEX (p=0.07)

Table I (Continued).

Study (First Author, RCT Name, Year)	Patients	Study Design and Study Groups	Follow-Up	Results Numerical Values Refer to the End of the Study Follow-Up Unless Stated Otherwise				
DEX implant in	DEX implant in DME refractory to anti-VEGF							
Totan et al (2016) ²⁰	30 DME refractory to BEV	Study Design: Prospective Single-site Open-label Groups: 30 DEX 0.7 mg	6 months	Mean change in BCVA +0.03 LogMAR DEX (p=0.39) Mean change in CFT -106 µm DEX (p=0.01)				
Kim et al (2016) ²¹	35 DME refractory to BEV	Study Design: Retrospective Single-site Groups: 35 DEX 0.7 mg	6 months	BCVA improvement ≥ 10 letters 26% DEX Mean change in BCVA -0.03 LogMAR DEX (p=0.652) Mean change in CFT -69.2 µm DEX (p<0.01)				
Shah et al (2016) ²²	50 DME refractory to anti-VEGF injections	Study Design: RCT Single-site Single-masked Groups: 23 BEV 1.25 mg 27 DEX 0.7 mg	7 months	$\label{eq:statestarding} \begin{array}{ c c c } \hline \mbox{Mean number of injections} \\ \hline \mbox{7.0 BEV} \\ \hline \mbox{2.7 DEX} \\ \hline \mbox{BCVA improvement} \geq 10 \mbox{ letters} \\ \hline \mbox{26\% BEV} \\ \hline \mbox{33\% DEX} \\ \hline \mbox{(p=0.758)} \\ \hline \mbox{Mean change in BCVA} \\ \hline \mbox{+5.6 letters DEX} \\ \hline \mbox{(p=0.785)} \\ \hline \mbox{Mean change in CST} \\ \hline \mbox{-13 } \mu m \mbox{BEV} \\ \hline \mbox{-122 } \mu m \mbox{DEX} \\ \hline \mbox{(p=0.001)} \\ \hline \end{array}$				
Thomas et al (2016) ²³	22 DME refractory to RAN	Study Design: Retrospective Single-site Groups: II DEX 0.7 mg II RAN 0.5 mg	3 months	Mean change in BCVA -0.153 LogMAR DEX -0.125 LogMAR RAN (p=0.624) Mean change in CMT -105.8 μm DEX -47.9 μm RAN (p=0.332)				
Busch et al (2018) ²⁴	110 DME refractory to AFL, RAN or BEV	Study Design: Retrospective Multicenter Groups: 72 AFL/RAN/BEV 38 DEX	12 months	Mean change in BCVA -0.4 letters AFL/RAN/BEV +6.1 letters DEX (p=0.004) Mean change in CST +18.3 µm AFL/RAN/BEV -92.8 µm DEX (p<0.001)				

Table I (Continued).

Study (First Author, RCT Name, Year)	Patients	Study Design and Study Groups	Follow-Up	Results Numerical Values Refer to the End of the Study Follow-Up Unless Stated Otherwise
Koc et al (2023) ²⁵	94 DME refractory to AFL or RAN	Study Design: Retrospective Multicenter Groups: 3AFL + 3AFL 3RAN + 3RAN 3AFL + DEX 3RAN + DEX	6 months	Mean BCVA 0.49 LogMAR 3AFL + 3AFL 0.57 LogMAR 3RAN + 3RAN 0.62 LogMAR 3AFL + DEX 0.80 LogMAR 3RAN + DEX (p=0.159) Mean CMT 291.6 µm 3AFL + 3AFL 293.1 µm 3RAN + 3RAN 320.0 µm 3AFL + DEX 332.5 µm 3RAN + DEX (p=0.295)
Maturi et al, DRCR.net prot U (2018) ²⁶	129 DME refractory to anti-VEGF injections	Study Design: Phase 2 RCT Multicenter Single-masked Sham-controlled Groups: 65 DEX 0.7 mg + RAN 0.3 mg 64 Sham + RAN 0.3 mg	24 weeks	Mean change in BCVA +2.7 letters DEX + RAN +3.0 letters RAN (p=0.73) Mean change in CST -110 µm DEX + RAN -62 µm RAN (p<0.001)
Simultaneous DE	X implant and anti-VEGF injection	ons		
Kaya et al (2021) ²⁷	68 DME treatment naïve or refractory to anti-VEGF	Study Design: RCT Single-site Open-label Groups: 34 DEX 0.7 mg + RAN 0.5 mg 34 RAN 0.5 mg	24 months	BCVA improvement \geq 15 letters 65.4% DEX + RAN 26.2% RAN (p<0.0001) Mean change in BCVA +20.2 letters DEX + RAN +9.1 letters RAN (p<0.001) Mean change in CFT -394 µm DEX + RAN -275 µm RAN (p<0.05)

Abbreviations: RCT, randomized clinical trial; ME, macular edema; DME, diabetic macular edema; DEX, dexamethasone; BCVA, best-corrected visual acuity; CRT, central retinal thickness; BEV, bevacizumab; CMT, central macular thickness; CST, central subfield thickness; RAN, ranibizumab; AFL, aflibercept; CFT, central foveal thickness.

Rationale for DEX Implant Use in DME

Inflammation is a pivotal pathogenetic factor in DME development and maintenance.²⁸ Several pro-inflammatory mediators are over-expressed in the vitreous and humor aqueous of patients with DR: angiogenic factors, cytokines, chemokines, complement and coagulation cascade proteins, proteins involved in ROS production, etc.^{29,30}

Intravitreal corticosteroids, due to their anti-inflammatory and anti-edematous effects, can well counteract the multifaceted pathways involved in the pathogenesis of DME, and may therefore be a high-performance treatment option in the therapeutic armamentarium of DME. In particular, corticosteroids inhibit phospholipase A2 and arachidonic acid pathways, reduce the VEGF expression and suppress the activity of inflammatory cytokines, thereby stabilizing the blood-retinal barrier, blocking

Study (first author, RCT name, year)	D1	D2	D3	D4	D5	Overall
RCTs supporting the use of DEX in DME				h		
Kuppermann et al. (2007) ⁹	+	+	!	+	!	+
Haller et al. (2010) ¹⁰		+			+	!
Boyer et al., MEAD (2014) ¹¹	+	+	+	+	+	+
Callanan et al., PLACID (2013) ¹²	+	+	+	+	!	+
Heng et al., OZLASE (2016) ¹³	+	+				!
Wei et al. (2021) ¹⁴	-	+	+	+	+	!
DEX implant vs anti-VEGF injections						
Gillies et al., BEVORDEX (2014) ¹⁵	+	+	+	+	+	+
Fraser-Bell et al., BEVORDEX (2016) 50	+	+	+	+		+
Callanan et al. (2017) ¹⁶	+	+	+	+	+	+
Sharma et al. (2020) ¹⁷		+	!	+		!
Ozsaygili et al. (2020) ¹⁸		+	+	+	!	!
Comet et al., INVICTUS (2020) ¹⁹	-	+		+		!
DEX implant in DME refractory to anti-V	ÆGF					
Totan et al. (2016) ²⁰	-	+	+	+	!	!
Kim et al. (2016) ²¹	-	+	+			!
Shah et al. (2016) 22	+	+	+	+	+	+
Thomas et al. (2016) ²³	+	+	+	+	-	!
Busch et al. (2018) ²⁴	-	+	!	+		!
Maturi et al., DRCR.net prot U (2018) ²⁶	+	+	+	+	+	+
Simultaneous DEX implant and anti-VEGF injections						
Kaya et al. (2021) 27		+	+	+	!	

Figure 2 Risk of bias assessment for clinical studies involving patients affected by diabetic macular edema (DME) who received dexamethasone (DEX) intravitreal implant. Notes: D1, randomization process; D2, deviations from the intended interventions; D3, missing outcome data; D4, measurement of the outcome; D5, selection of the reported result; "+", low risk; "!", some concerns; "--", high risk.

the leukostasis, enhancing density and activity of tight junctions in the retinal capillary endothelium, ameliorating retinal oxygenation and preventing endothelial cell injury and apoptosis.^{31,32}

Several optical coherence tomography (OCT) imaging biomarkers have been identified for DME and used to make diagnosis, monitor, and provide prognostic information on both disease progression and potential responses to different

treatments.^{30,33} In addition, some of these biomarkers are considered to correlate with intraocular inflammation levels and responsiveness to anti-VEGF and steroid therapy.³⁴

In summary, the OCT-biomarkers identified so far are the following:

- Increased retinal thickness: result of the intraretinal and/or sub-retinal accumulation of fluid, but also intracellular edema. This OCT parameter is commonly used in clinical studies to define cutoff for eligibility to a certain treatment, evaluate the activity as well as the course of the disease, and treatment response.³⁵ However, central macular thickness (CMT) poorly correlates with BCVA, especially in long-standing DME.³⁶

- Intraretinal cystoid spaces: due to intraretinal fluid accumulation; size, location, and chronicity have been associated with functional prognosis in diabetic eyes.³⁷

- Hyperreflective retinal foci (HRF): intraretinal dots in the inner or outer retinal layer showing similar reflectivity to the retinal nerve fiber layer, diameter <30 μ m and absence of back-shadowing. It has been hypothesized that these dots may represent extravasated lipoproteins or aggregates of activated microglial cells,³⁸ and that they are strictly linked to inflammation.³⁹

- Hyperreflective cystoid walls: consisting of collagen-like material or gliotic tissues; usually detected in association with HRF, hyperreflective cystoid walls are considered a sign of increased inflammatory activity.⁴⁰

- Exudates: located within the outer retinal layers, showing similar reflectivity to the RPE–Bruch's membrane complex, characterized by a size >30 μ m and by the presence of back-shadowing. They are associated with lipid levels and other systemic risk factors.⁴¹

- Disorganization of retinal inner layers (DRIL): disorganization or destruction of cells within the inner retinal layers; it is a negative prognostic factor for visual outcome in eyes with DME.⁴²

- Disorganization of Retinal Outer Layer (DROL): damage of the outer retinal layers, particularly of the photoreceptor outer segment, correlates with poor visual prognosis and macular ischemia.⁴³

- Subfoveal subretinal fluid (SRF): due to the disruption of the external limiting membrane that allows fluid and protein to migrate to the subretinal space. It is a sign of inflammation and a biomarker for functional and anatomical treatment response.⁴⁴

- Central choroidal thickness (CCT): a consequence of both increased luminal and stromal structures, it also can be correlated with the presence of subretinal fluid. CCT usually increases during DR progression and its reduction after treatment can be used as a predictive biomarker of treatment response.⁴⁵

- Vitreomacular interface: mechanical traction on the fovea is correlated to a worse prognosis and may suggest a surgical add-on therapy.⁴⁶

As anticipated above, large intraretinal cystoid spaces and HRF have been associated with a significant inflammatory component that, therefore, may suggest favorable treatment response to corticosteroids.⁴⁷ It has been recently suggested that large intraretinal cysts (\geq 250 µm), DRIL, HRF and SRF in chronic DME may be OCT biomarkers to select candidates for intravitreal steroids as first-line treatment.^{33,47}

Randomized Controlled Trials Supporting the Use of DEX in DME

The first RCT evaluating the efficacy and safety of DEX implant in patients with macular edema was conducted in 2007: 315 patients with persistent macular edema due to different etiologies were randomized to either receive a single 0.7 mg or 0.35 mg DEX implant injection or to be observed for 6 months.⁹ The analysis of the subgroup of 171 patients with DME confirmed the superiority of DEX implant 0.7 mg in terms of proportion of patients achieving \geq 10-letters and \geq 15-letters BCVA improvement as well as anatomical improvement, suggesting a dose–response relationship.¹⁰ Similarly, the proportion of patients showing a significant reduction in both fluorescein leakage and central retinal thickness (CRT) was higher in the DEX implant 0.7 mg group.¹⁰

In 2014, the MEAD studies represented the registration trials for DEX implant in patients with DME, and were designed as two randomized, multicenter, masked, sham-controlled, Phase III trials with identical protocol.¹¹ A total of 1048 patients with center involving DME were randomized to receive either 0.7 mg or 0.35 mg of DEX implant or a sham procedure and followed up for 3 years. Percentage of patients with \geq 15-letter improvement in BCVA and average CRT reduction were significantly higher in the DEX implant groups (always p < 0.02).

RCTs also compared the efficacy of DEX implant with laser photocoagulation therapy, which was considered the standard of care for DME for almost 30 years. Specifically, the PLACID trial demonstrated the superiority of 0.7 mg DEX implant plus laser photocoagulation versus laser photocoagulation alone in terms of BVCA gain at 1- and 9-month follow-up (FU) (31.7% vs 11%, p < 0.001, and 31.7% vs 17.3%, p = 0.007, respectively), but not at 12-month FU. Central macular thickness (CMT) reduction and decrease of the area of vascular leakage were also greater in the DEX implant plus laser group.¹² Later on, the OZLASE clinical trial compared the efficacy and safety of combined repeated DEX implant plus macular laser therapy (MLT) and MLT only in 80 patients with center involving DME over a 1-year period. Although the CST decrease was higher in the combination group (p < 0.001), no difference in BCVA improvement was noted between groups (p > 0.05).

More recently, DEX implant was compared to laser photocoagulation by Wei et al, in a phase III RCT including 284 Chinese and Filipino patients with DME.¹⁴ The BCVA gain was significantly higher for DEX implant (p = 0.001); however, the proportion of patients with BCVA improvement ≥ 15 letters did not differ significantly between groups (p > 0.05). Both CRT reduction and decrease in the leakage area on FA were significantly higher in the DEX implant group at 12 months.¹⁴

DEX Intravitreal Implant and Anti-VEGF Injections for DME Dexamethasone vs Anti-VEGF

The BEVORDEX study was the first RCT comparing anti-VEGF injections and DEX implant for DME treatment;^{15,48} 88 eyes were randomized to receive either bevacizumab or DEX implant for 24 months. No difference was found in terms of BCVA gain at 2 years in pseudophakic eyes, and results remained similar at 5 years.⁴⁹ Although significantly higher CRT reduction in the DEX implant group was noted at 12 months (187 μ m vs 122 μ m in the bevacizumab group; p = 0.015), this became not significant at both 2 and 5 years.^{48,49} At 5 years, eyes initially randomized to receive DEX implant had significantly fewer treatments but were more likely to develop proliferative diabetic retinopathy (PDR) compared to eyes initially randomized to bevacizumab.⁴⁹

Subsequently, a multicenter, randomized study by Callanan et al evaluated the non-inferiority of DEX implant compared to ranibizumab in 363 eyes with DME.¹⁶ DEX implant matched the prespecified noninferiority criteria of BCVA gain at 12 months, which was defined as a mean BCVA change within 5 ETDRS letters compared to ranibizumab, even if BCVA gain was significantly higher in the ranibizumab group (7.6 vs 4.3 ETDRS letters, p < 0.05). In addition, Sharma et al did not find statistically significant differences between DEX implant and anti-VEGF (bevacizumab or ranibizumab) in terms of BCVA change in 40 naïve eyes with DME;¹⁷ while better anatomical results were observed in the DEX implant group in a short follow-up of 3 months. Similar better anatomical results were found by Ozsaygili et al comparing DEX implant with aflibercept in treatment-naive patients with DME and inflammatory biomarkers over a 12-month FU, showing a significantly higher mean CRT reduction in the DEX group (p < 0.05). In their study functional results favored the aflibercept group, however, this superiority ceased to exist when considering pseudophakic eyes only, with the advantage of a lower number of injections in the DEX implant group.¹⁸

The INVICTUS study, a real-life study comparing intravitreal ranibizumab (RAN), aflibercept (AFL) and dexamethasone on naïve patients with DME, did not show significant differences among treatments both in BCVA gain (DEX = +9.4 letters; RAN = +7.2; AFL = +8.5) and CRT reduction (DEX = $-162.7 \mu m$; RAN = $-99.3 \mu m$; AFL = $-101.5 \mu m$).¹⁹

A few meta-analyses also compared the effectiveness and safety of DEX implant and VEGF-inhibitors in DME treatment. He et al found comparable visual acuity improvements and CMT reduction at 12 months, with fewer injections in the DEX implant group. However, authors stated that visual acuity benefit could not be fully reflected in the DEX implant group due to cataract progression, and therefore recommended DEX implant as first-line therapy only in selected cases.⁶ Kodjikian et al reviewed only observational studies evaluating the efficacy of anti-VEGF and DEX implant. Although reaching similar final BCVA (62 and 61.2 for anti-VEGF and DEX implant respectively), the reported BCVA gain from baseline was + 4.7 letters in the anti-VEGF group and + 9.6 letters in DEX implant group. However, only the naïve status appeared to confer the best functional response in the DEX-implant studies. The authors suggested this

difference could be partially explained by the lower baseline BCVA in patients receiving the DEX implant or the less frequency of anti-VEGF injections in real-life observational studies compared to interventional ones.⁵⁰

More recently, two additional meta-analyses were conducted. Chi et al⁵¹ included 2409 eyes and showed no significant differences between anti-VEGF and DEX implant in terms of BCVA improvement from the pooled analysis of non-resistant eyes (mean difference = 0.00). Similarly, Patil et al included fourteen RCTs for a total of 827 eyes, reporting no significant difference in BCVA gain at 3, 6 and 12 months (p = 0.11, p = 0.21 and p = 0.24, respectively) as well as at final follow-up (p = 0.91). Significantly lower retinal thickness at 3 and 6 months (p = 0.04 and p < 0.00001 respectively) and final follow-up (weighted mean difference of 39.99 µm, p = 0.002) were noted in the DEX implant group; however, no significant difference was reported at 12 months (p = 0.18).⁵²

Dexamethasone in DME Refractory to Anti-VEGF

Resistance to anti-VEGF therapy is a major concern in DME management, and DEX implant has been used as alternative or adjuvant therapy in eyes with no satisfactory DME regression after at least three anti-VEGF injections.⁵³

In 2016, Totan et al evaluated the efficacy of DEX implant in 30 eyes with chronic DME resistant to at least 3 bevacizumab injections. Significant improvements in BCVA and CRT were noted at 1 and 3 months; however, such improvements were not sustained over time and tended to decrease between the third and sixth months after the injection.²⁰ Similar results were found in other studies, with BCVA improvements at 6 months in about one-fourth of the patients.²¹ These results were consistent with other reports: Shah et al reported comparable visual acuity improvement at 7 months comparing intravitreal bevacizumab monotherapy and DEX implant monotherapy for persistent DME; however, reduced CRT and fewer injections in the DEX implant group were noted;²² Thomas et al in a contralateral eye-to-eye comparison showed no differences in BCVA change at 3 months between the eye that continued with ranibizumab injections and the eye that switched to DEX implant; CRT reduction, although similar in both eyes, reached statistical significance compared to baseline in the DEX implant treated eye only.²³

Interestingly, Busch et al evaluated 110 eyes with refractory DME in a real-world setting, showing that eyes that were switched to DEX implant had better visual and anatomical outcomes at 12 months compared to those that continued treatment with anti-VEGF therapy.²⁴

More recently, a report by Koc et al confirmed the absence of any difference in BCVA and CMT in 94 eyes with DME resistant to ranibizumab or aflibercept injections, which either continued with the anti-VEGF or switched to DEX implant.²⁵

Lately, a systematic review and meta-analysis revealed the DEX implant to be associated with significantly greater BCVA improvement and CRT decrease compared to anti-VEGF therapy in resistant DME (mean difference = 0.12 [95% CI, 0.02-0.21] and 65.37 [95% CI, 3.62-127.13] respectively) with, however, high heterogeneity between included studies.⁵¹

DEX implant has also been used as adjuvant therapy in refractory DME. A Diabetic Retinopathy Clinical Research Network (DRCR.net) study Protocol U, a Phase II multicenter randomized clinical trial, explored the role of adjunctive therapy with Ozurdex in 129 eyes with persistent DME after at least 3 ranibizumab injections.²⁶ DEX implant (65 eyes) or sham injection (64 eyes) was added to monthly ranibizumab protocol. No differences in BCVA were found between groups, as mean (SD) improvement at 24 weeks was +2.7 (9.8) letters in the DEX implant plus ranibizumab group, and +3.0 (7.1) in the ranibizumab only group (p = 0.73). However, the anatomical outcome favored the combination group ($-110 \mu m vs -62 \mu m$; p < 0.001) and the percentage of DME resolution was 52% in the DEX implant plus ranibizumab group vs 31% in the ranibizumab only group.

Considering DME refractory to a large number of intravitreal injections, a recent non-randomized interventional study explored the potential role of DEX implant in a small sample of such eyes.⁵⁴ Six eyes with failed clinical response after a mean of 43.5 (range 15–58) prior anti-VEGF injections, and all with failed affibercept, underwent DEX implant plus a breakthrough anti-VEGF injection when required. At 1 year, the mean BCVA improved from 0.475 to 0.375 logMAR and central subfield thickness (CST) reduced from 386.5 to 315 μ m. The authors concluded that DEX implant appeared effective in eyes with DME refractory to multiple intravitreal injections, with excellent anatomic improvement and modest visual improvement.

Simultaneous DEX Implant and Anti-VEGF for DME

Besides studies evaluating the efficacy of DEX implant in comparison or sequential to anti-VEGF treatments, a few studies explored the efficacy of the simultaneous administration of DEX implant and anti-VEGF for the treatment of DME. Although being small-sized studies, results seem to suggest the superiority of such simultaneous combined treatment compared to VEGF inhibitors alone, with a favorable visual and anatomic outcome and an acceptable safety profile.^{27,55–57}

Kaya et al assessed the efficacy and safety of simultaneously administered DEX implant and ranibizumab in comparison with ranibizumab only as the first-line treatment of DME in 68 eyes (34 in each group) with follow-up at 12 and 24 months.^{27,57} At both time-points, the simultaneous double protocol therapy significantly improved the visual outcomes (+21.6 vs +9.6 letters in the ranibizumab only group at 12 months, p < 0.001; and +20.2 vs +9.1 at 24 months), with 65.4% of patients in the simultaneous double protocol group gaining \geq 15 ETDRS letters from baseline (vs 26.2% in ranibizumab only group; p < 0.001). Moreover, FT reduced more in the simultaneous double protocol group at 12 months (-413 µm vs -282 µm; p = 0.001) and at 24 months (-394 µm vs -275 µm; p < 0.05); and inflammatory OCT biomarkers improved more in the simultaneous double protocol group (38% vs 18%; p = 0.012), as well as cataract progression (27% vs 12.5%; p = 0.032). Overall, the simultaneous double protocol resulted an effective first-line treatment option for DME with inflammatory OCT biomarkers or/and decreased visual acuity.^{27,57}

Economic Considerations on DEX Implant and Anti-VEGF for DME

A relevant concern regarding DME involves treatment costs. Simultaneous therapy with DEX implant and anti-VEGF may be justified by outcome improvements; however, health authorities have not yet approved such combination protocols. The current trend is to minimize the frequency of intravitreal injections. Monthly protocols for anti-VEGF injections have significantly reduced in popularity, and the new standard of care is now represented by treat-and-extend protocols, which can reduce economic burden, while maintaining similar degree of efficacy.⁵⁸

Treatments with intravitreal corticosteroids require even fewer injections, and a recent study suggested that switching to DEX implant after the first 3-monthly anti-VEGF injections may be more cost-effective than extending the treatment to 6-monthly injections.⁵⁹ However, longer-acting anti-VEGF drugs, such as brolucizumab or faricimab, have been recently produced, which still need to be compared to DEX implant for the purpose of cost-efficiency.^{60–62}

Ocular Adverse Events in DEX Implant vs Anti-VEGF

Beyond their therapeutic effect, steroids are able to bind tissues, inducing various effects on cell proliferation, transmembrane transport and reactive oxygen species activity.⁶³ Water solubility intensely impacts the wash-out time of intravitreal drugs and the exposure time is considered a key factor in determining the steroid-binding properties to intraocular tissues.⁶⁴ Among steroids approved for intravitreal use, dexamethasone is the most soluble in water, due to its reduced lipophilicity, thus showing faster clearance compared to other intravitreal steroids;²⁸ thus, it is not surprising that DEX implant showed a lower incidence of intraocular adverse events compared to triamcinolone acetonide and FA.^{65,66}

Intravitreal steroids showed particular affinity for lens and trabecular meshwork cells and this may explain the high incidence of cataract worsening and IOP increase.⁶⁵ For instance, it has been demonstrated that dexamethasone may interfere with pivotal pathways which regulate lens cell homeostasis, migration and differentiation.⁶⁷ Moreover, dexamethasone showed to increase the stiffness of trabecular meshwork cells in vitro, by inducing over-expression of contractile proteins and synthesis of extracellular matrix.⁶⁸ In addition, the accumulation of steroid molecules in the trabecular meshwork may determine increase in intraocular volume and obstruction due to precipitation which also play a role in IOP elevation.⁶⁵

The incidence of adverse events is largely variable among studies (Table 2).^{11,31,65,69–72}

In the MEAD registration trial for Ozurdex,^{11,69} the percentages of IOP-related adverse events were 36.0%, 34.1% and 5.1% in the DEX implant 0.7 mg, 0.35mg, and sham groups, respectively; 6.6% of patients in the DEX implant 0.7 mg (the approved dose) group experienced a measured IOP \geq 35 mmHg, compared to 0.9% in the sham group. IOP

 Table 2 Adverse Events of Clinical Studies Involving Patients Affected by Diabetic Macular Edema Who Received Dexamethasone

 Intravitreal Implant

Study (First Author, RCT Name, Year)	IOP ≥ 10 mmHg Than Baseline	IOP ≥ 25 mmHg at Any Timepoint	Cataract-Related Adverse Events
RCTs supporting the use of DEX i	n DME		
Kuppermann et al (2007) ⁹	17% DEX 0.7 mg 12% DEX 0.35 mg 3% observation	18% DEX 0.7 mg 15% DEX 0.35 mg 0% observation	N/A
Haller et al (2010) ¹⁰	15% DEX 0.7 mg 15% DEX 0.35 mg 2% observation	13.2% DEX 0.7 mg 16.4% DEX 0.35 mg 0% observation	N/A
Boyer et al, MEAD (2014) ¹¹	27.7% DEX 0.7 mg 24.8% DEX 0.35 mg 3.7% Sham	32.0% DEX 0.7 mg 27.4% DEX 0.35 mg 4.3% Sham	67.9% DEX 0.7 mg 64.1% DEX 0.35 mg 20.4% Sham
Callanan et al, PLACID (2013) ¹²	15% DEX + laser 9% Sham + laser	16.8% DEX + laser	22% DEX + laser 9% Sham + laser
Heng et al, OZLASE (2016) ¹³	N/A	N/A	77.8% DEX + laser 14.8% laser
Wei et al (2021) ¹⁴	33.1% DEX 1.6% laser	33.8% DEX 4.7% laser	15.2% DEX 6.2% laser
DEX implant vs anti-VEGF injection	ons		
Gillies et al, BEVORDEX (2014) ¹⁵	IOP ≥ 5 mmHg 19% BEV 1.25 mg 46% DEX 0.7 mg	0% BEV 1.25 mg 26% DEX 0.7 mg	4.8% BEV 1.25 mg 13% DEX 0.7 mg
Fraser-Bell et al, BEVORDEX (2016) ⁴⁸	IOP ≥ 5 mmHg 48% BEV 1.25 mg 74% DEX 0.7 mg	N/A	6% BEV 1.25 mg 37% DEX 0.7 mg
Callanan et al (2017) ¹⁶	34.3% DEX 3.8% RAN (p<0.001)	36.5% DEX 2.2% RAN (p<0.001)	22.2% DEX 9.5% RAN (p<0.001)
Sharma et al (2020) ¹⁷	N/A	0% BEV/RAN 20% DEX (p<0.001)	0% BEV/RAN 15% DEX (p<0.001)
Ozsaygili et al (2020) ¹⁸	IOP ≥ 5 mmHg I6.7% DEX 0% AFL (p=0.002)	N/A	8.3% DEX 0% AFL
Comet et al, INVICTUS (2020) ¹⁹	IOP increase ≥ 10 mmHg than IOP ≥ 25 mmHg at any timepo 0% RAN 0% AFL 19% DEX	N/A	

Study (First Author, RCT Name, Year)	IOP ≥ 10 mmHg Than Baseline	IOP ≥ 25 mmHg at Any Timepoint	Cataract-Related Adverse Events				
DEX implant in DME refractory to anti-VEGF							
Totan et al (2016) ²⁰	N/A	IOP ≥ 21 mmHg 13.3% DEX	N/A				
Kim et al (2016) ²¹	N/A	8.6% DEX	N/A				
Shah et al (2016) ²²	N/A	IOP ≥ 21 mmHg 0% BEV 52% DEX (p<0.001)	17% BEV 26% DEX (p=0.515)				
Thomas et al (2016) ²³	N/A	IOP > 30 mmHg 18.1% DEX 0% RAN	N/A				
Busch et al (2018) ²⁴	N/A	N/A	11.3% AFL/RAN/BEV 26.3% RAN				
Maturi et al, DRCR.net prot U (2018) ²⁶	23% DEX + RAN 0% RAN	N/A	5% DEX + RAN 0% RAN (p=0.24)				
Simultaneous DEX implant and a	nti-VEGF injections						
Kaya et al (2021) ²⁷	IOP ≥ 5 mmHg 35.3% DEX + RAN 18% RAN	N/A	35.3% DEX + RAN 29.4% RAN				

Table 2 (Continued).

Abbreviations: RCT, randomized clinical trial; DME, diabetic macular edema; DEX, dexamethasone; BEV, bevacizumab; RAN, ranibizumab; AFL, aflibercept.

rises were usually managed with IOP-lowering drops or no treatment, and only 0.6% of patients in the 0.7mg group required glaucoma surgery. There was no cumulative effect of repeated dexamethasone implants on IOP, as the incidence and level of IOP elevations did not increase over time, nor the proportion of patients using IOP-lowering medications over the 3-year observation time. Cataract-related adverse events in phakic eyes were 67.9%, 64.1%, and 20.4% in the DEX implant 0.7 mg, 0.35 mg, and sham groups, respectively. However, cataract surgery successfully restored vision without further complications.¹¹

Additionally, a more recent large study on the safety of over 6000 intravitreal dexamethasone implants reported a frequency of 26.5% of eyes developing IOP \ge 25 mmHg during follow-up, with only 0.5% requiring glaucoma surgery, and 32.5% requiring cataract surgery, with an additional 14.6% experiencing not-significant cataract worsening.⁷²

Compared to other steroids approved for intravitreal use, a systematic review and meta-analysis reported an incidence of increased IOP in 11–15% of patients treated with dexamethasone implants (0.35 and 0.7 mg doses, respectively), significantly lower compared to the 32% of 4mg triamcinolone acetonide, and 66–79% with FA implants (0.59 mg and 2.1 mg doses, respectively).⁶⁵

Other ocular side effects of dexamethasone implant are relatively rare and mostly related to the surgical procedure rather than to the drug itself. Endophthalmitis (sterile or not), retinal detachment, vitreous hemorrhage, implant dislocation in the anterior chamber, conjunctival hyperemia or hemorrhage have been reported in some studies, generally with an incidence lower than 1%.^{72,73}

Many studies reported higher number of eyes which underwent cataract surgery in DEX implant groups compared to anti-VEGF groups. However, literature reports conflicting results regarding cataract development rates, with some studies showing significant differences, while others do not.^{6,16,17,22,26,48,49,52} In the BEVORDEX study, 37% of patients treated

with DEX implant underwent cataract surgery at 24 months, compared to 6% of patients in the bevacizumab group, and similar rates were found by Callanan et al^{16,48} He et al reported higher incidence of cataract-related AEs in the DEX implant group compared to the anti-VEGF group (RR = 2.68, p = 0.0005)⁶ and the risk of developing cataract was even higher in a recent meta-analysis of RCTs (OR = 4.70, p = 0.0002).⁷⁴ Conversely, Patil et al did not find differences in cataract-related AEs between DEX implant and anti-VEGF in their systematic review and meta-analysis.⁵² Actual differences between DEX implant and anti-VEGF drugs in terms of cataract progression rates remain unclear, probably due to differences in follow-up times, protocols and proportions of pseudophakic patients at baseline among studies. Despite this, phakic status remains one of the most important factors to consider, and anti-VEGF still represents the first-line treatment in phakic patients with DME.⁵

A Cochrane meta-analysis of two clinical trials demonstrated that the need for IOP-lowering therapy increased more than 4 times with DEX implant compared to anti-VEGF, with moderate-certainty evidence,^{16,48} and these results were consistent with those of He et al (RR = 4.14; 95% CI, 1.89–8.65; p = 0.0002)⁶ and Patil et al.⁵² The INVICTUS study showed a significantly higher proportion of patients with IOP \geq 25 mmHg or an increase of \geq 10 mmHg in the DEX implant group compared to the aflibercept or ranibizumab group.¹⁹ In addition, 23% of eyes that switched to DEX implant plus ranibizumab experienced an increase \geq 10 mmHg compared to 0% of eyes in the ranibizumab only group in the DRCR.net protocol U,²⁶ and similar percentages were also reported by Sharma et al.¹⁷ A recent meta-analysis of RCTs also reported higher incidence of high IOP following DEX implant compared to anti-VEGF (OR = 13.77, p < 0.00001).⁷⁴ Surprisingly, recent meta-analysis by Chi et al did not reveal significant difference in IOP change between DEX implant group and anti-VEGF group (MD = -0.42).⁵¹

Overall, intravitreal dexamethasone has an excellent systemic safety profile. Its slow-release technology and the intravitreal route of administration allow to achieve the maximum therapeutic concentration into the vitreous body and the minimum systemic absorption, limiting the incidence of systemic adverse events.⁷⁵ In clinical trials, the incidence of systemic AEs after Ozurdex administration is extremely rare and usually with no differences compared to sham treatments.¹¹ Compared to anti-VEGF agents, DEX implant offers a better systemic safety profile and is indicated as first-line treatment in patients at high risk for cardiovascular events, in which anti-VEGF is contraindicated.⁵

DEX Implant vs FA Implant for DME

The FA implant presents different pharmacokinetics compared to the DEX implant: in its current formulation, it releases 0.2 μ g of FA/day, maintaining a steady-state concentration over a longer period of 36 months; however, peak functional and anatomical efficacy is achieved more slowly, usually 6 months after implantation.⁷⁶

Both RCTs and real-life studies have provided insights about the efficacy and safety profiles of the FA implant.^{77–79} In the phase 3 trials FAME A and B, Campochiaro et al compared the effect of 0.2 μ g/day (low dose) or 0.5 μ g/day (high dose) FA implants on subjects with persistent DME. After a period of 2 years, both the low and high dose FA implants significantly improved both BCVA (mean changes of +4.4 and +5.4 letters, respectively; p = 0.02 and p = 0.016 vs sham) and foveal thickness (mean changes of –168 μ m and –177 μ m, respectively; p = 0.005 and p < 0.001 vs sham); however, the safety profile was more favorable towards the low-dose insert, with significantly lower incidence of glaucoma and cataract surgery.⁷⁸

In the PALADIN study, a Phase 4, nonrandomized, open-label observational study,⁷⁷ Singer et al analyzed the functional and anatomical outcomes of 202 eyes during a follow-up period of 36 months, reporting a mean BCVA change of +3.61 letters (p = 0.02) and a mean CST change of -60.69 µm (p < 0.0001) compared with baseline.⁷⁷ These positive results were confirmed in a meta-analysis of real-world studies by Fallico et al.⁸⁰

Retrospective observational studies were conducted to evaluate the efficacy, safety and timing for switching FA implant to DEX implant.^{81,82} Baillif et al reported results of 113 eyes switched to FA implant after a mean time of 11.12 ± 22 weeks. BCVA improved from 54.1 to 59.8 letters at month 4 and remained stable during the observation period. A significant reduction was noted in CMT, with a minimum of 320.9 µm at month 3. Finally, a mean IOP remained below 19 mmHg at all evaluated timepoints. A longer time between the last DEX and the first FA implant was a risk factor for additional treatments; however, no significant outcome differences were reported between eyes that received their last DEX implant either less than or more than 8 weeks prior to switching to FA implant.⁸¹

More recently, in the ILUVI1MOIS study, 41 pseudophakic eyes with chronic DME (time to recurrence ≤ 6 months) were switched to FA implant 1 month after the last DEX implant. BCVA and CMT remained stable during a follow-up of at least 6 months (mean 13.1 ± 4.4 months), with 14% of patients requiring additional treatments at 12 months. The mean IOP at month 12 did not differ significantly from that measured at baseline.⁸²

According to a retrospective study by Cicinelli et al, the anatomical outcomes and IOP elevation of FA implant seem to be predicted by the previous response to DEX implant; however, no significant relationship was found between BCVA improvement after DEX and after FA.⁸³

To our knowledge, only one study by Coelho et al directly compared the effectiveness and safety of the FA and DEX implants for the treatment of DME. However, the sample size was limited, and the follow-up time between the two groups was different.⁸⁴ Comparing the outcomes of FA and DEX implants is challenging because of the different pharmacokinetics of the two drugs.

Overall, switching from DEX to FA implants is effective and safe, and can be justified in patients who might not respond adequately to DEX implant, who obtained only short-term improvements or who are willing to undergo a lower number of injections. Notably, FDA has approved the use of FA implant for DME patients who previously underwent DEX implant, as long as they did not experience a significant increase in IOP.⁸¹

DEX Implant in Diabetic Eyes Undergoing Cataract Surgery

Diabetic patients, especially those with type 2 diabetes, have a higher risk of developing early cataracts,⁸⁵ with an incidence almost two-fold higher compared to non-diabetic subjects (20.4 vs 10.8 per 1000 person/years).⁸⁶ In fact, chronic hyperglycemia causes increased oxidative stress and non-enzymatic glycation of eye lens proteins that may precipitate the formation of cataract.⁸⁷ As a result, cataract surgery is often necessary. Among complications after cataract surgery, post-cataract macular edema (PCME) has an incidence after uncomplicated surgeries ranging between 0.1% and 2.3% which, however, can be significantly higher in diabetic patients, reaching values around 16.3% in patients with DR and/or previous DME, with a risk proportional to the severity of DR, mainly due to a compromised blood-retinal barrier even before the surgery.^{88–91}

Chronic and persistent DME represents the major risk factor for edema worsening after cataract surgery, owing to the combination of chronic inflammation in DR and surgery-related inflammation. Therefore, performing intravitreal dexamethasone implant injection at the same time of cataract surgery would be a safe and effective option for treating both diabetic and post-cataract edema.⁹²

Several studies showed the beneficial effects of DEX implant in eyes with DME undergoing cataract surgery. In the MEAD study, the subgroup of eyes that underwent phacoemulsification during the follow-up period experienced an increase in CRT after cataract surgery in the sham group, but not in the DEX implant group, suggesting a protective role of DEX implant following cataract surgery.¹¹ Subsequently, Panozzo et al evaluated the role of DEX implant at the time of cataract surgery in reducing DME worsening. However, with no control group, their results showed that intraoperative DEX implant was effective in preventing DME worsening after phacoemulsification with maximum efficacy at 4–8 weeks after the injection and lasting for at least 3 months.⁹³ A control group was included in the retrospective-comparative study conducted by Furino et al on diabetic patients with DME and cataract,⁹⁴ which showed that phacoemulsification combined with DEX implant (phaco-DEX group) had better functional and tomographic outcomes compared to standard phacoemulsification (phaco-alone group). Specifically, BCVA improved significantly after surgery in the phaco-DEX group (p = 0.0005), while no significant changes were noted in the phaco-alone group. Similarly, CST was significantly lower in the phaco-DEX group compared to the phaco-alone group.⁹⁴

In this context, compared to anti-VEGF therapy, DEX implant seems to better antagonize the inflammatory storm that comes from DME pathogenesis and after cataract surgery. A recent meta-analysis conducted by Fallico et al found that cataract surgery combined with DEX implant showed a 3-month mean change in CMT of $-98.35 \mu m$, significantly higher compared to cataract surgery combined with anti-VEGF therapy ($-21.61 \mu m$; p < 0.001). However, no differences were found in visual gain between the two groups (p = 0.13).⁹⁵

The protective effect of DEX implant after cataract surgery is compatible with the well-known effects of the preoperative prophylaxis with topical steroids and nonsteroidal anti-inflammatory drugs. Laursen et al conducted a meta-

analysis of 6 RCTs with nearly 900 patients to compare prophylactic interventions with (1) nonsteroidal antiinflammatory drugs (NSAIDs) alone or in combination with topical steroids, (2) depot and topical steroids, and (3) anti-VEGF alone or in combination with topical steroids. Intravitreal corticosteroids showed similar protective effect against macular edema in comparison to both topical steroids alone and to topical steroids in combination with NSAIDs.⁹⁶

A series of studies were carried out to establish the optimal timing of DEX implant in diabetic patients undergoing cataract surgery. Barone et al compared DEX implant administered either 1 month before or at the time of cataract surgery in 40 patients with treatment naïve DME; in a twenty-week follow-up, no difference was found in terms of visual acuity gain or anatomical outcomes between the two groups.⁹⁷ A larger retrospective study with similar design was conducted on 221 patients with cataract and DME.⁹⁸ BCVA gain was comparable between groups, but better anatomical results were noticed in the group receiving DEX implant at the time of cataract surgery (p < 0.001). Lastly, Corbelli et al, in a sample of 40 eyes with DME and cataract, found that DEX implant injection performed at the time of cataract surgery achieved similar long-term anatomical and functional outcomes compared to a 1-month deferred injection.⁹⁹

Even if there is no general agreement about the optimal timing of DEX implant (before, during, or after cataract surgery), the combination of the two procedures may protect against the acute surgical-induced inflammation, potentially preventing short-term subclinical damages and may decrease the burden of the number of visits and examinations.⁹⁹ Therefore, consensus statements of experts recommend the use of DEX implant in patients with DME at the time of phaceenulsification to prevent the surgery-related inflammatory load and to protect from potential decompensation of DME.^{100,101}

DEX Implant in Vitrectomized Eyes with Macular Edema

Persistent vitreous hemorrhages and tractional retinal detachment are well-known complications of proliferative diabetic retinopathy (PDR), and pars plana vitrectomy (PPV) is the only available therapy to treat these sight-threatening conditions; therefore, it is not uncommon for patients with PDR to undergo PPV during their lifetime.^{102,103} Interestingly, in addition to its main mechanical effect, PPV has shown several secondary positive effects on the course of diabetic retinopathy: for instance, it reduces the concentration of pro-inflammatory cytokines and VEGF near the macular area,¹⁰⁴ improves the oxygen diffusion to the ischemic inner retinal layers,¹⁰⁵ and accelerates the washout from the vitreous cavity of signaling molecules involved in diabetic retinopathy progression.¹⁰⁶ Due to the chronic nature of PDR, as well as the potential pro-inflammatory effect of the surgery itself, even patients who undergo PPV may show DME development or worsening.¹⁰⁴

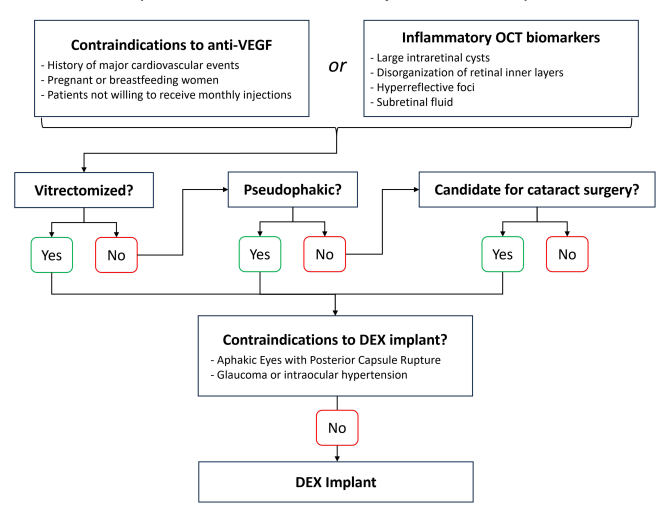
Management of DME in vitrectomized patients may be challenging; vitreous replacement with other media may alter the pharmacokinetic profile of intravitreal drugs, thus limiting their beneficial effects.¹⁰⁷ According to the law of Stokes-Einstein, molecular transport by diffusion or convection is inversely related to the viscosity of the medium,¹⁰⁵ and in saline solution diffusion results quicker than in vitreous humor.¹⁰⁸ Previous studies demonstrated that several molecules such as anti-VEGF agents, antibiotics and triamcinolone acetonide have faster clearance in vitrectomized eyes,^{109–111} which may result in a reduction of their efficacy in vivo.¹¹² DEX implant, due to its slow-release properties, has shown long-lasting clearance even in patients who underwent PPV, showing no differences in efficacy on DME treatment comparing vitrectomized and non-vitrectomized eyes.^{113–117}

The CHAMPLAIN study was the first designed to evaluate the efficacy of a single injection of 0.7 mg DEX implant in a cohort of 55 patients with a history of PPV and refractory DME.¹¹³ A significant reduction of CCT was noticed at week 8 (-156μ m; p < 0.001) and maintained at 26 weeks (39 µm; p=0.004); similarly, BCVA significantly improved at week 8 (+6.0 letters; p < 0.001) and maintained at 26 weeks (+3.0 letters; p = 0.046) with an acceptable safety profile. Maximum efficacy was seen at week 8 after injection.¹¹³

Many subsequent studies explored the differences in DME treatment efficacy between eyes with and without previous PPV. Medeiros et al compared BCVA and foveal thickness in 58 patients with refractory DME (24 vitrectomized and 34 non-vitrectomized) which received DEX implant with a follow-up of 6 months.¹¹⁴ Both groups showed significant improvements in BCVA and foveal thickness, with no significant difference between treatment groups at any time point (1, 3 and 6 months). Later, Bastakis et al evaluated the efficacy and duration of action of DEX implant in 18 eyes with persistent DME after anti-VEGF injection. Ten out of 18 eyes were vitrectomized.¹¹⁵ Each eye received 1 to 3 injections and was followed-up for 6 to 18 months. No difference in BCVA and CMT variations after dexamethasone injections

were observed between vitrectomized and non-vitrectomized eyes and DEX implant showed its maximum efficacy within the first 3 months of implantation. A further observational, retrospective, real-life study including 186 eyes (59 vitrectomized and 127 non-vitrectomized) showed no difference in BCVA gain and CMT decrease between groups, with a similar intraocular pressure profile. Although no difference was noted in the overall number of injections, the mean interval between injections was significantly shorter in vitrectomized eyes (5.2 vs 6.9 months in non-vitrectomized eyes; p = 0.001).¹¹⁶ The same group also analyzed a small sample of 15 eyes with macular edema secondary to different conditions, including diabetes, that received DEX implant both before and after PPV, with each eye acting as its own control.¹¹⁶ No significant differences in CMT reduction, BCVA improvement, number of injections required, interval between the injections and IOP profile were noticed before and after PPV.¹¹⁶

Recently, the VITDEX study, a multicenter, retrospective, interventional study, investigated the effect of DEX implant in 236 eyes with DME with or without previous PPV with a follow-up of 12 months; 130 eyes were vitrectomized and 106 were non-vitrectomized.¹¹⁷ Both groups showed improvements in BCVA and CST with no statistically significant difference between groups. In addition, no differences were found in the number of DEX implants, the need for additional treatments and the proportion of patients who underwent cataract surgery or started IOP-lowering medications during the follow-up period. Thus, the authors concluded that both efficacy and safety of DEX implant were not influenced by PPV.



DEX implant could be considered as a viable first-line treatment option in:

Figure 3 Flowchart summarizing possible determinants in patient selection for DEX implant in DME.

Finally, a systematic review and meta-analysis including 7 retrospective studies with a total of 582 eyes (374 nonvitrectomized and 208 vitrectomized) and a follow-up ranging from 4 to 26 months showed no significant difference between groups in terms of BCVA improvement, CMT reduction, number of injections and number or magnitude of adverse events at 1, 3, 6 and 12 months; however, the duration of action of DEX implant was significantly shorter in vitrectomized eyes, with a mean difference of 0.8 months (p = 0.005), confirming that vitreous removal may accelerate drug clearance after injections. However, this difference was not deemed clinically relevant.⁸

Conclusion

The EURETINA guidelines recognized the importance of corticosteroids in the treatment armamentarium of DME; however, they were largely defined on a second-choice level compared to anti-VEGF, mainly due to the relatively higher risk of ocular side effects.⁵

More recent literature and expert recommendations using a Delphi approach have partially moved beyond these guidelines, suggesting the use of DEX implant as a viable first-line treatment option in pseudophakic patients and recommending its use as first-choice in vitrectomized eyes, given the better pharmacokinetic profile in this setting compared to anti-VEGF.^{101,118,119}

Furthermore, DEX implant could also be the preferred choice in phakic patient candidates for cataract surgery, by protecting from potential decompensation of previous DME,^{101,118–120} and in patients with DME with high inflammatory component, evaluated through the presence of inflammatory OCT biomarkers (large intraretinal cysts, DRIL, HRF and SRF).^{33,101,118,119,121} Additionally, in patients not responding to anti-VEGF injections, clinicians should consider switching to DEX implant after the loading dose of 3–6 consecutive anti-VEGF injections, while patients with severe and persistent DME can benefit from a combination of DEX implant and anti-VEGF (Figure 3).^{101,118}

Conversely, exclusion criteria for the administration of the DEX implant include torn or ruptured posterior lens capsule, due to the risk of implant migration in the anterior chamber, as well as glaucoma and intraocular hypertension. In these cases, anti-VEGF injections should be preferred.

Overall, when appropriate, using DEX implant as first-line treatment option in the real-world setting can facilitate treatment compliance by reducing the number of injections and visits, potentially improving treatment outcomes.^{19,24,101}

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