Indications and Usage

**Retinal Vein Occlusion:** OZURDEX® (dexamethasone intravitreal implant) is a corticosteroid indicated for the treatment of macular edema following branch retinal vein occlusion (BRVO) or central retinal vein occlusion (CRVO).

**Posterior Segment Uveitis:** OZURDEX® is indicated for the treatment of noninfectious uveitis affecting the posterior segment of the eye.

**Diabetic Macular Edema:** OZURDEX® is a corticosteroid indicated for the treatment of diabetic macular edema.

**IMPORTANT SAFETY INFORMATION**

**Contraindications**

**Ocular or Periocular Infections:** OZURDEX® is contraindicated in patients with active or suspected ocular or periocular infections including most viral diseases of the cornea and conjunctiva, including active epithelial herpes simplex keratitis (dendritic keratitis), vaccinia, varicella, mycobacterial infections, and fungal diseases.

Please see additional Important Safety Information on page 3.
FDA-approved indications

- Diabetic macular edema
- Macular edema following branch or central retinal vein occlusion
- Noninfectious posterior segment uveitis

Delivers intravitreal dexamethasone via NOVADUR® technology

- A sustained-release, biodegradable steroid implant
  — Solid polymer matrix contains 0.7 mg of dexamethasone, a corticosteroid
  — Biodegrades to lactic acid and glycolic acid
  — Administered by injection as an in-office procedure

Dosage and Administration

FOR OPHTHALMIC INTRAVITREAL INJECTION. The intravitreal injection procedure should be carried out under controlled aseptic conditions. Following the intravitreal injection, patients should be monitored for elevation in intraocular pressure and for endophthalmitis. Patients should be instructed to report any symptoms suggestive of endophthalmitis without delay.

Important Safety Information (continued)

Contraindications (continued)

Glaucoma: OZURDEX® (dexamethasone intravitreal implant) is contraindicated in patients with glaucoma, who have cup to disc ratios of greater than 0.8.

Torn or Ruptured Posterior Lens Capsule: OZURDEX® is contraindicated in patients whose posterior lens capsule is torn or ruptured because of the risk of migration into the anterior chamber. Laser posterior capsulotomy in pseudophakic patients is not a contraindication for OZURDEX® use.

Please see additional Important Safety Information on page 4.
Congratulations (continued)

Hypersensitivity:
OZURDEX® (dexamethasone intravitreal implant) is contraindicated in patients with known hypersensitivity to any components of this product.

Warnings and Precautions
Intravitreal Injection-related Effects:
Intravitreal injections, including those with OZURDEX® (dexamethasone intravitreal implant), have been associated with endophthalmitis, eye inflammation, increased intraocular pressure, and retinal detachments. Patients should be monitored regularly following the injection.

Corticosteroid-related Effects:
Use of corticosteroids including OZURDEX® may produce posterior subcapsular cataracts, increased intraocular pressure, glaucoma, and may enhance the establishment of secondary ocular infections due to bacteria, fungi, or viruses.

Corticosteroids should be used cautiously in patients with a history of ocular herpes simplex because of the potential for reactivation of the viral infection.

Please see additional Important Safety Information on page 6.

Preparing for intravitreal injection

Recommendations for aseptic technique, antimicrobial prophylaxis, and anesthesia

1. Both preparation and the intravitreal injection procedure should be carried out under controlled aseptic conditions, which include the use of sterile gloves, a sterile drape, and a sterile eyelid speculum (or equivalent).

2. A broad-spectrum microbicide applied to the periocular skin, eyelid, and ocular surface is recommended to be given prior to the injection.

3. Subconjunctival anesthesia should also be administered before the intravitreal injection. Subconjunctival anesthesia was used in the phase 3 clinical trials of OZURDEX® (dexamethasone intravitreal implant).³

(continued on next page)

4. Remove the foil pouch from the carton and examine it for damage. Then, in a sterile field, open the foil pouch and gently drop the applicator on a sterile tray.

Each applicator can only be used for the treatment of a single eye

• If the contralateral eye requires treatment, a new applicator must be used, and the sterile field, syringe, gloves, drapes, and eyelid speculum should be changed before OZURDEX® (dexamethasone intravitreal implant) is administered to the other eye.

IMPORTANT SAFETY INFORMATION (continued)

Contraindications (continued)

Hypersensitivity: OZURDEX® (dexamethasone intravitreal implant) is contraindicated in patients with known hypersensitivity to any components of this product.
IMPORTANT SAFETY INFORMATION (continued)

Adverse Reactions
Retinal Vein Occlusion and Posterior Segment Uveitis (continued)
Adverse reactions reported by greater than or equal to 1% of patients in the two combined 3-year clinical trials following injection of OZURDEX® (dexamethasone intravitreal implant) for diabetic macular edema include: cataract (68%), conjunctival hemorrhage (23%), visual acuity reduced (9%), conjunctivitis (6%), vitreous floaters (5%), conjunctival edema (5%), dry eye (5%), vitreous detachment (4%), vitreous opacities (3%), retinal aneurysm (3%), foreign body sensation (2%), corneal erosion (2%), keratitis (2%), anterior chamber inflammation (2%), retinal tear (2%), eyelid ptosis (2%). Non-ocular adverse reactions reported by greater than or equal to 5% of patients include: hypertension (13%) and bronchitis (5%).

Please see additional Important Safety Information on page 8.
Patients should be monitored for elevation in intraocular pressure and for endophthalmitis. Monitoring may consist of the following:

- Perfusion check of the optic nerve head immediately after injection
- Tonometry within 30 minutes following the injection
- Biomicroscopy between 2 and 7 days following the injection

**Recommended follow-up**

**Patients should be counseled regarding the risk of potential complications including, but not limited to, endophthalmitis, elevated intraocular pressure, or cataract**

- Inform patients of the need to be vigilant for new symptoms in the days following intravitreal injection
- Stress the importance of seeking immediate care from an ophthalmologist if the eye becomes red, sensitive to light, painful, or develops a change in vision
- Advise patients that they may develop increased intraocular pressure with OZURDEX® (dexamethasone intravitreal implant) treatment, and the increased IOP will need to be managed with eye drops and, rarely, with surgery
- Advise patients that a cataract may occur after repeated treatment with OZURDEX®, which may decrease their vision and require an operation to remove the cataract and restore their vision
- Also advise patients that they may experience temporary visual blurring after receiving an intravitreal injection — They should not drive or use machines until the blurring has resolved

**IMPORTANT SAFETY INFORMATION (continued)**

**Adverse Reactions (continued)**

**Diabetic Macular Edema (continued)**

**Increased Intraocular Pressure:** IOP elevation greater than or equal to 10 mm Hg from baseline at any visit was seen in 28% of OZURDEX® (dexamethasone intravitreal implant) patients versus 4% of sham patients. 42% of the patients who received OZURDEX® were subsequently treated with IOP-lowering medications during the study versus 10% of sham patients.

The increase in mean IOP was seen with each treatment cycle, and the mean IOP generally returned to baseline between treatment cycles (at the end of the 6-month period).

**Cataracts and Cataract Surgery:** The incidence of cataract development in patients who had a phakic study eye was higher in the OZURDEX® group (68%) compared with Sham (21%). The median time of cataract being reported as an adverse event was approximately 15 months in the OZURDEX® group and 12 months in the Sham group. Among these patients, 61% of OZURDEX® subjects versus 8% of sham-controlled subjects underwent cataract surgery, generally between Month 18 and Month 39 (Median Month 21 for OZURDEX® group and 20 for Sham) of the studies.

Please see additional Important Safety Information on back cover.
IMPORTANT SAFETY INFORMATION (continued)

Contraindications

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Glaucoma: OZURDEX® is contraindicated in patients with glaucoma, who have cup to disc ratios of greater than 0.8.

Please see full Prescribing Information.