Continuous Range of High-Quality Vision at All Distances.

From the Leader in Presbyopia-Correcting IOLs

**Anterior Side**
- Posterior Achromatic Diffractive Surface and Echelette Feature

**Posterior Side**
- Haptics Offset for 3 Points of Fixation
- Tecnis® IOL Wavefront-Designed Anterior Aspheric Surface
- 13.0 mm Overall Diameter
- Frosted, Continuous 360° Posterior Square Edge
- 6.0 mm Optic Diameter

Johnson & Johnson Vision
OPTICAL CHARACTERISTICS

Model Number: ZXR00
Powers: +5.0 D to +34.0 D in 0.5 diopter increments
Diameter: 6.0 mm
Center Thickness: 0.7 mm (20.0 D)
Shape: Biconvex, wavefront-designed anterior aspheric surface, posterior achromatic diffractive surface designed to reduce chromatic aberration for improved image contrast and echelette feature to extend the range of vision.
Material: UV-blocking hydrophobic acrylic
Refractive Index: 1.47 at 35º C
Edge Design: ProTEC frosted, continuous 360° posterior square edge

BIOMETRY*

A-constant: 118.8
AC Depth: 5.4 mm
Surgeon Factor: 1.68 mm

CONTACT ULTRASOUND†

A-constant: 119.3
AC Depth: 5.7 mm
Surgeon Factor: 1.96 mm

OPTICAL††

HAPTIC CHARACTERISTICS

Overall Diameter: 13.0 mm
Thickness: 0.46 mm
Style: C
Material: Soft, Foldable, UV-blocking hydrophobic acrylic
Design: TRI-FIX, Haptics offset from optic; 1-piece lens

RECOMMENDED INSERTION INSTRUMENTS

UNFOLDER® Platinum 1 Series Screw-Style Inserter (DK7796)
UNFOLDER® Emerald-AR Inserter (EMERALDAR)
ONE SERIES® Ultra Syringe-Style Inserter (DK7786)
UNFOLDER® Platinum 1 Series Cartridge (IMTEC30)
UNFOLDER® Emerald-AR Cartridge (ICART30)
ONE SERIES® Ultra Cartridge (IVIPR30)

INDICATIONS and IMPORTANT SAFETY INFORMATION for TECNIS SYMFONY EXTENDED RANGE OF VISION IOLs

Rx Only

INDICATIONS FOR USE
The TECNIS Symfony Extended Range of Vision IOL, Model ZXR00, is indicated for primary implantation for the visual correction of aphakia, in adult patients with less than 1 diopter of pre-existing corneal astigmatism, in whom a cataractous lens has been removed. The lens mitigates the effects of presbyopia by providing an extended depth of focus. Compared to an aspheric monofocal IOL, the lens provides improved intermediate and near visual acuity, while maintaining comparable distance visual acuity. The Model ZXR00 IOL is intended for capsular bag placement only.

WARNINGS
Patients with any of the conditions described in the Directions for Use may not be suitable candidates for an intraocular lens because the lens may exacerbate an existing condition, may interfere with diagnosis or treatment of a condition, or may pose an unreasonable risk to the patient’s eyesight. Lenses should not be placed in the ciliary sulcus. May cause a reduction in contrast sensitivity under certain conditions, compared to an aspheric monofocal IOL; fully inform the patient of this risk before implanting the lens. Special consideration should be made in patients with macular disease, amblyopia, corneal irregularities, or other ocular disease. Inform patients to exercise special caution when driving at night or in poor visibility conditions. Some visual effects may be expected due to the lens design, including: a perception of halos, glare, or starbursts around lights under nighttime conditions. These will be bothersome or very bothersome in some people, particularly in low-illumination conditions, and on rare occasions, may be significant enough that the patient may request removal of the IOL.

PRECAUTIONS
Interpret results with caution when refactoring using autorefractors or wavefront aberrometers that utilize infrared light, or when performing a duochrome test. Confirmation of refraction with maximum plus manifest refraction technique is recommended. The ability to perform some eye treatments (e.g. retinal photocoagulation) may be affected by the optical design. Target emmetropia for optimum visual performance. Care should be taken to achieve IOL centration, as lens decentration may result in a patient experiencing visual disturbances under certain lighting conditions.

SERIOUS ADVERSE EVENTS
The most frequently reported serious adverse events that occurred during the clinical trial of the Tecnis Symfony lens were cystoid macular edema (2 eyes, 0.7%) and surgical reintervention (treatment injections for cystoid macular edema and endophthalmitis; 2 eyes, 0.7%). No lens-related adverse events occurred during the trial.

ATTENTION: Reference the Directions for Use for a complete listing of indications and important Safety Information.