

IC-8™ Aphera™

Small Aperture IOL



WAVEFRONT-FILTERING
OPTIC
TECHNOLOGY

The first approved small aperture IOL designed to provide extended depth of focus.

EXCELLENT RESULTS IN EYES
WITH UP TO **1.5D OF ASTIGMATISM**^{1,2}

HARNESS THE POWER OF FOCUSED LIGHT

Built on a high-quality monofocal IOL platform with the addition of a carbon black FilterRing™

Filters out peripheral low OQ light entering the eye, delivering only high OQ central light rays to the retina

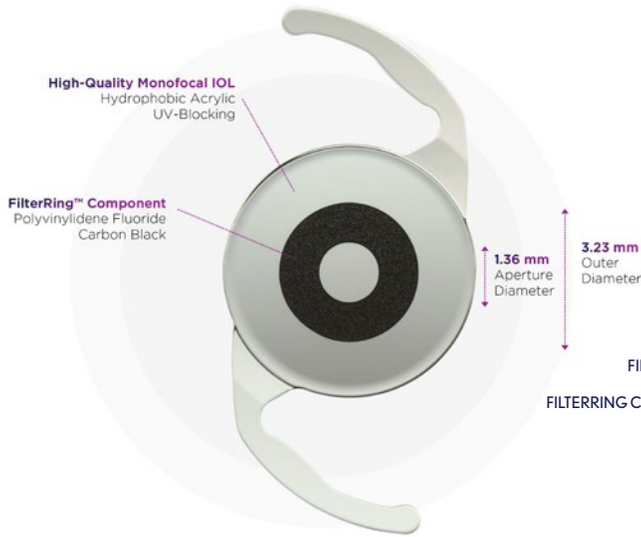
OQ = optical quality

Delivers extended depth of focus, free from “blurry zones”,¹ providing a continuous clear range of vision from near to far

BAUSCH + LOMB



IC8APT order number IC8APT-XXX



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| MODEL | IC-8™ Aphera™ IOL |
| MATERIAL (OPTIC AND HAPTIC) | Hydrophobic Acrylic UV-blocking |
| POWERS | +10.0 D through +30.0 D in 0.5 D increments |
| OPTIC TYPE | One-piece Biconvex, anterior aspheric surface |
| OPTIC DIAMETER (ØB) | 6.0 mm |
| OVERALL DIAMETER (ØT) | 12.5 mm |
| OPTIC EDGE DESIGN | 360° posterior square edge |
| HAPTIC DESIGN | Modified C-loop haptic with 5° angulation |
| REFRACTIVE INDEX | 1.483 at 35°C and 589 nm |
| FILTERRING™ COMPONENT MATERIAL | Polyvinylidene fluoride (PVDF) with carbon black |
| FILTERRING COMPONENT OUTER DIAMETER | 3.23 mm |
| FILTERRING COMPONENT INNER DIAMETER (APERTURE) | 1.36 mm |
| BIOMETRY | Optical |
| A-CONSTANT | 120.5 |
| SURGEON FACTOR | 2.64 |
| ANTERIOR CHAMBER DEPTH (ACD) | 6.42 |

Optical biometry A-constant values are presented as a guideline. Physicians should calculate the lens power based on their experience and preference.

bioli™ IOL Delivery System

bioli-A1

FOR INSERTING LENS MODEL IC-8 Aphera
RECOMMENDED INCISION SIZE 3.2mm
TYPE OF ACTION Push-type
COMMENTS Single-handed delivery. Disposable.



STORZ
Ophthalmic Instruments

Find B+L IOL surgical equipment
online at www.StorzEye.com

1. IC-8 Aphera Directions for Use, Bausch + Lomb, Inc.
2. Burkhard Dick et al. Prospective multicenter trial of a small-aperture intraocular lens. *Journal of Cataract and Refractive Surgery* 2017; Vol. 43, Issue 7; 956-968

Indications and Important Safety Information for IC-8 Aphera IOL

INDICATIONS: The IC-8 Aphera IOL is indicated for unilateral implantation for the visual correction of aphakia and to create monovision in patients of age 22 or older who have been diagnosed with bilateral operable cataract, who have up to 1.5 D of astigmatism in the implanted eye, and who do not have a history of retinal disease and who are not predisposed to experiencing retinal disease in the future. The device is intended for primary implantation in the capsular bag, in the non-dominant eye, after the fellow eye has already undergone successful implantation (uncorrected distance visual acuity 20/32 or better and best-corrected distance visual acuity 20/25 or better) of a monofocal or monofocal toric IOL that is targeted for emmetropia. The refractive target for the IC-8 Aphera IOL should be -0.75 D. The lens mitigates the effects of presbyopia by providing an extended depth of focus. Compared to an aspheric monofocal or monofocal toric IOL, the lens provides improved intermediate and near visual acuity, while maintaining comparable distance visual acuity.

CONTRAINDICATIONS: (1) Patients with dilated pupil size less than 7.0 mm. (2) Patients with a history of retinal disease including but not limited to, high myopia, diabetes, macular disease, sickle cell disease, retinal tear, retinal detachment, retinal vein occlusion, ocular tumor, uveitis, and patients who are predisposed to experiencing retinal disease in the future.

WARNINGS: The lens should not be implanted if appropriate intraocular support of the lens is not possible. Severe subjective visual disturbances (e.g., glare, halo, starburst, hazy vision) may occur after device implantation. There is a possibility that these visual disturbances may be significant enough that a patient may request removal of the lens. Contrast sensitivity in eyes implanted with this lens is significantly reduced when compared to the fellow eye implanted with a monofocal or monofocal toric IOL. Although there was no significant reduction in binocular contrast sensitivity in the IDE clinical study, it is essential that prospective patients be fully informed of this visual effect in the implanted eye before giving their consent for unilateral implantation of the lens. Patients should be informed that they may need to exercise caution when engaging in activities

that require good vision in dimly lit environments (such as driving at night or in poor visibility conditions). There is a possibility that visual symptoms due to reduced contrast sensitivity may be significant enough that a patient may request removal of the lens. This lens should not be implanted bilaterally because bilateral implantation is expected to cause significant reduction in contrast sensitivity under all lighting conditions. The use of this lens in patients with corneal astigmatism greater than 1.5 D is not recommended. Diagnostic tests in patients implanted with the lens may take longer and require some additional effort from the patient and the physician to perform. Use of some medical lasers to treat certain eye conditions may present potential risks of damaging the FilterRing component of the lens. Removal of the lens may be necessary prior to retinal or vitreal procedures. Surgeons should perform a careful benefit-risk assessment based on individual patient characteristics, weighing all the risks disclosed in the Directions for Use labeling against the benefit of extended depth of focus. Nd:YAG laser capsulotomy treatments may be more difficult to perform and may be less effective in an IC-8 Aphera IOL implanted eye. Specific training from Bausch + Lomb or an authorized representative of Bausch + Lomb related to YAG capsulotomy is required before a surgeon is authorized to implant the IC-8 Aphera IOL.

PRECAUTIONS: Prior to surgery, prospective patients should be informed of the possible risks and benefits associated with this lens and a Patient Information Brochure should be provided to the patient. Patients with a predicted postoperative astigmatism between 1.0 D and 1.5 D may not obtain as great an amount of improvement in intermediate vision compared to patients with lower amounts of astigmatism.

CAUTION: Federal law restricts this device to sale by or on the order of a licensed physician.

ATTENTION: Reference the Directions for Use labeling for a complete listing of important safety information.