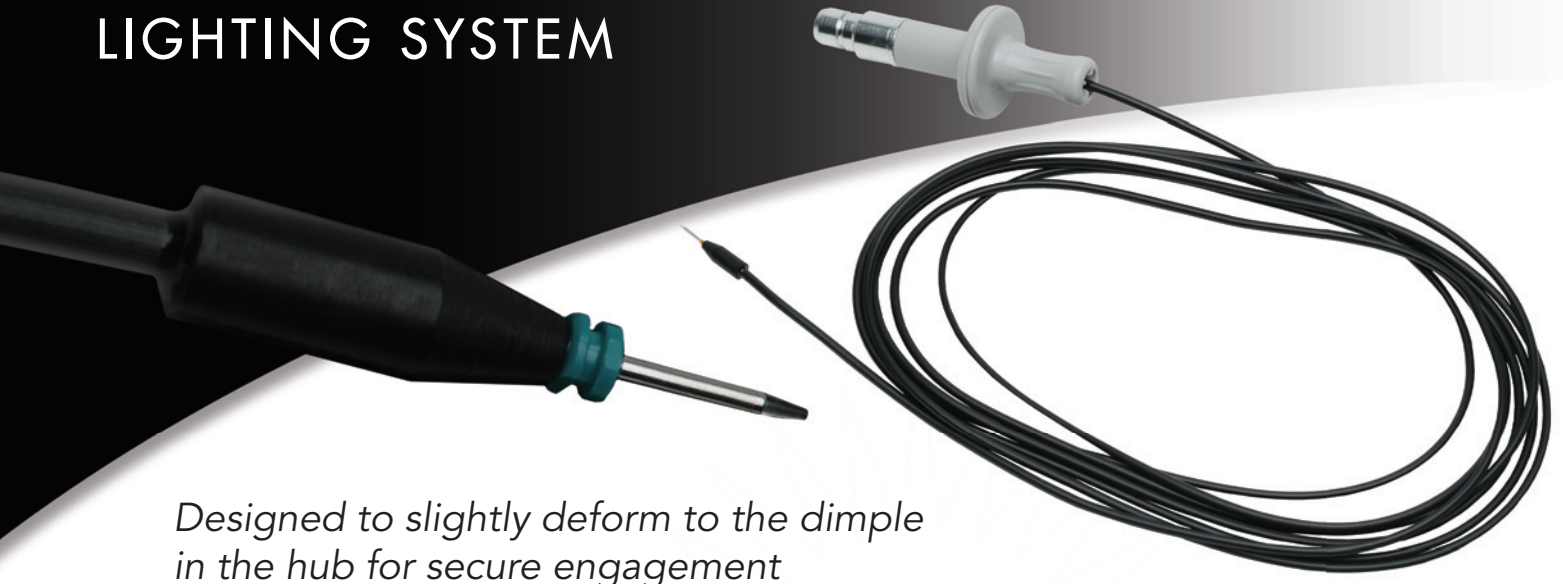


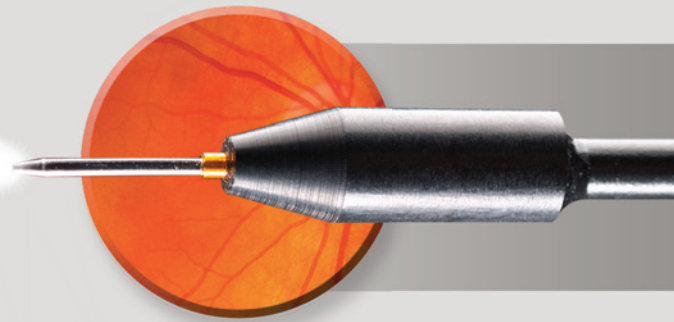
ALCON[®] CHANDELIER LIGHTING SYSTEM

Now Available!



Designed to slightly deform to the dimple in the hub for secure engagement

- Able to engage in either the 23G or 25G cannula hubs
- 25G EDGEPLUS[®] trocar/cannula, 1 each, set in pak
- Wider coverage area than standard or wide angle
- Designed to be shaped into a stable service loop
- Illumination angle is 106°



ALCON[®] CHANDELIER LIGHTING SYSTEM*

Chandelier, ACCURUS[®] Surgical System, Non RFID 8065751574

Chandelier, CONSTELLATION[®] Vision System, RFID 8065751577

*12/box

MIVS Indications:

INDICATIONS FOR USE: The CONSTELLATION[®] Vision System is an ophthalmic microsurgical system that is indicated for both anterior segment (i.e., phacoemulsification and removal of cataracts) and posterior segment (i.e., vitreoretinal) ophthalmic surgery. Caution: Federal (USA) law restricts this device to sale by, or on the order of, a physician. **WARNINGS AND PRECAUTIONS:** Attach only ALCON[®] supplied products to console and cassette luer fittings. Improper usage or assembly could result in a potentially hazardous condition for the patient. Mismatch of surgical components and use of settings not specifically adjusted for a particular combination of surgical components may affect system performance and create a patient hazard. Do not connect surgical components to the patient's intravenous connections. Each surgical equipment/component combination may require specific surgical setting adjustments. Ensure that appropriate system settings are used with each product combination. Prior to initial use, contact your ALCON[®] sales representative for in-service information. Care should be taken when inserting sharp instruments through the valve of the Valved Trocar Cannula. Cutting instrument such as vitreous cutters should not be actuated during insertion or removal to avoid cutting the valve membrane. Use the Valved Cannula Vent to vent fluids or gases as needed during injection of viscous oils or heavy liquids. Visually confirm that adequate air and liquid infusion flow occurs prior to attachment of infusion cannula to the eye. Ensure proper placement of trocar cannulas to prevent sub-retinal infusion. Leaking sclerotomies may lead to post operative hypotony. Vitreous traction has been known to create retinal tears and retinal detachments. Minimize light intensity and duration of exposure to the retina to reduce the risk of retinal photic injury. **ATTENTION:** Reference the Directions for Use labeling for a complete listing of indications, warnings, precautions, complications and adverse events.

Illuminated Indications:

CAUTION: Federal (USA) law restricts this device to sale by, or on the order of, a physician. **CAUTION:** This document is not intended to substitute for the necessity of reading and understanding the light source Operator's Manual. **INDICATIONS AND USAGE:** Fiber Optic Instruments with ENGAUGE[®] Radio Frequency Identification Device (RFID) for use with the CONSTELLATION[®] System. These instruments can be used on the ACCURUS[®] System or ACCURUS[®] High Brightness Illuminator (AHBI) using the RFID Adapter Model Number 8065751140. **WARNINGS AND PRECAUTIONS:** Minimize the light intensity and duration of exposure to the retina to reduce risk of retinal photic injury. Avoid operation of a fiber in air on consoles capable of illumination levels and settings higher than 10 lumens. This may result in fiber probe deformation and/or high surface temperatures that may cause patient injury. **ATTENTION:** Reference the Directions for Use labeling for a complete listing of indications, warnings, precautions, complications and adverse events.

Contact your local representative

alconretina.com

Alcon[®]

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