

MICROPURE

Monofocal Hydrophobic

BICONVEX ASPHERIC MONOFOCAL SURFACE

| S | | 2 | ש |
|---|----|---|---|
| | * | ~ | |
| | | | |
| | 51 | |) |

Description

۲

| Model | | MICROPURE | |
|-----------------------------------|--|----------------|--|
| Material | GFY Hydrophobic Acrylic ¹ | | |
| Overall diameter | -10D to 24.5D: 11.00mm 25D to 35D: 10.75mm | | |
| Optic diameter | -10D to 24.5D: 6.00mm 25D to 35D: 5.75mm | | |
| Optic | Biconvex Aspheric Monofocal | | |
| Haptic design | MICRO (closed loop quadripode) & Posterior Angulated Haptic | | |
| Filtration | UV & Blue Light | | |
| Refractive index | 1.53 | | |
| Abbe number | 42 | | |
| Injection system | Medicel Accuject 1.8 up to 24.5D Medicel Accuject 2.0/2.1/2.2 up to 35D | | |
| Spherical power | -10D to +9D (1D steps) +10D to +30D (0.5D steps) +31D to +35D (1D steps) | | |
| Suggested A constant ² | | Interferometry | |
| | Hoffer Q: pACD | 5.85 | |
| | Holladay 1: Sf | 2.06 | |
| | Barrett: LF | 2.09 | |
| | SRK/T: A | 119.40 | |
| | Haigis³: a0; a1; a2 | 1.70; 0.4; 0.1 | |

¹ The PhysIOL GFY[®] is patented since 2010.

² Values estimated only: surgeons are recommended to personalize their A-constant based on their surgical techniques and equipment, experience with the lens model and postoperative results.

³ Not optimized.

Note: The MICROPURE intraocular lens is not FDA approved.

Contact Information: www.bvimedical.com/customer-support/

BVI and all other trademarks (unless noted otherwise) are property of BVI. BVI ©2022

59<u>0642-02</u>

bvimedical.com

۲

Product Information

| Manufacturer | PhysIOL s.a Liège Science Park Allée des Noisetiers 4 B-4031 Belgium +32 4 361 05 49 physiol@bvimedical.com | |
|--|--|--|
| Certificate information | cate information CE (EU) 2017/745, Annex IX Chapter II : MDR 735732 R000 QMS (EU) 2017/745, Annex IX Chapter I and III : MDR 735719 R000 ISO 13485:2016 & EN ISO 13485:2016 : MD 658518 ISO 13485:2016 : MDSAP 691544 | |
| Shelf life | Five (5) years from manufacturing date for MICROPURE | |
| Intended Use The posterior chamber intraocular lens is intended to be placed into the capsular bag w capsulorhexis for the replacement of the human lens to achieve the visual correction of a patients in whom the cataractous lens has been removed. | | |
| Indication for use | The lens should be used as intended in adult patients surgically treated for cataract, who desire improved uncorrected far vision. | |
| Product Composition | No products of animal or human origin are present in the implant. The intraocular lens is 100% composed of the covalently crosslinked proprietary material of medical quality (GFY), which is a (2-hydroxyethylmethacrylate; phenoxy ethylacrylate; polypropylene glycol dimethacrylate) copolymer, including a UV and a blue light-filtering chromophores covalently bound to the material. | |
| For sterile product | All IOLs from PhysIOL are steam sterilized | |
| Packaging Material | Holder (Polypropylene) Container (Polypropylene) Storage liquid (0.9% NaCl solution) Aluminium lid (Aluminium Gold) Container label (paper) Blister PP (Polypropylene) Tyvek lid | |
| Product Class | Classified as Class IIb implantable long-term surgically invasive medical devices under Rule 8 of Anne VIII of the MDR 2017/745. Not available in the United States | |
| | | |

۲

۲

Contact Information: www.bvimedical.com/customer-support/

BVI and all other trademarks (unless noted otherwise) are property of BVI. BVI ©2022



۲