

Technical Data Sheet I-Stream Hybrid

Product code: ISP60H + diopter

Model: I-Stream H

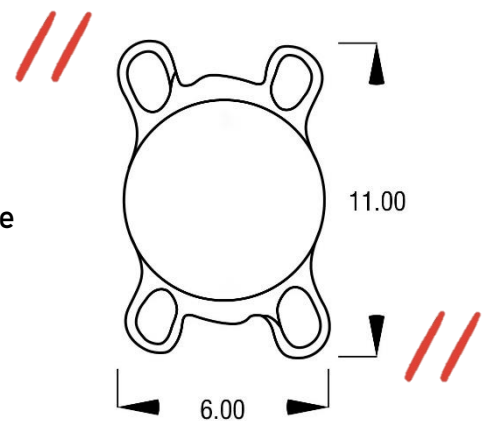
Manufacturer: Md Tech Srl - Via Fratelli Bandiera, Ncc - 80026 Casoria (Na), Campania Italia

Intended Use: Cataract surgery

Use: Intraoperative ophthalmic use

Italian Registration Number: 119903-119904-125908

Acrylic copolymer hydrophilic/hydrophobic preloaded single-piece iol aspheric. Quadrilateral shape with fenestrated haptics.



// General Features

Material	Hybrid (hydrophilic with hydrophobic surface), biocompatible with eye tissue
Lens Geometry	Biconvex optic, Monofocal, Aspheric, Square Edge 360°.
Dioptric Range	from -5.0D a + 10.0D with increment of 1D; from +10.5D to +31.5D with increment of 0.5 Dpt; from +32.0 to 40.0 with increment of 1D
Incision	from 1.8mm with preloaded injector

// Optical Features

Optic Diameter	6.0mm
Total Diameter	11.0mm
Sferical Aberration	-0,26 μ m
Color	Natural Yellow
Refractive Index	1,46
UV Filter	Natural Yellow UV CUT OFF < 2% @420nm
A Costant	118,2
Constant with Ulib System	SRK/T = 118,7; SRK/2 = 118,9; Holladay1 sf = 1,60; Holladay2 = 5,374; Barrett = 1,73 HofferQ pACD = 5,39; Haigis a0 = 1,25; a1 = 0,40; a2 = 0,10

Haptics Features

Type	Optical disc and haptics without profile discontinuity
Haptic Angulation	5° for a long term better stability in the capsular bag

// Other Specifications

Reference Standard	Directive 93/42 CEE, UNI EN ISO:11979
Class	II b Certiquality Srl, Notified Body n°0546, certif. N.18539/1/I
Control Performed	DIOPTER , MTF, Cosmetic
Packaging	Each lens is packed in a lens case in WFI contained in a blister of PETG and Tyvek (primary packaging) and in carton box (secondary packaging) in kit with MEDICEL injector.
Product Type	Medical device in compliance with the current legislation
Compatibility With Solutions	Product can be used with hyaluronic acid, balanced salt solution. It must not be used with acid or melted with other substances.
Shelf Life and Sterilization Method	Product has 36 months of validity from sterilization date and however until expiry date reported on labels. Steam sterilized in conformity to UNI EN ISO 17665. The injector is sterilized with EtO in accordance with UNI EN ISO: 11135-1 / 2. The product is NOT RESTERILISABLE.
Tests	In vitro cytotoxicity tests, allergic sensitization tests on guinea pigs, eye irritation tests on guinea pigs, product stability tests have been carried out on the product in the experimental phase (reports available in the company)
Presence of Latex	No
Storage	Store at ambience condition, in dry place without humidity. Temperature must be in the range +5°C and +45°.
Information Reported on the Packaging	Batch number, expiry date, serial number, sterilization, manufacturer, CE mark.

