

Sterimedix Ltd

Quality System Document QD 2040 Issue P
EC Declaration of Conformity

Sterimedix Ltd.

Declaration of Conformity for Irrigation / Aspiration Cannulae and Hand Pieces

Ref: DOC07

18/12/2018 - Issue 12

Council Directive 93/42/EEC Concerning Medical Devices

The undersigned declares that the product named in this document meets the applicable Council Directive provisions that apply and the CE Mark may be affixed.

General Product Name Irrigation / Aspiration Cannulae and Hand Pieces

Product Reference **Bimanual Cannulae and Hand Pieces**
M5405; M5501; M5502; M5505; M5510; M5511; M5512; M5513; M5514; M5515; M5518;
M5519; M5520; M5521; M5522; M5523; M5525; M5526; M6524
Bimanual I/A Systems
M5331; M5531; M5532; M5533; M5535; M5536; M5537; M5538; M5539; M5543
Coaxial Cannulae and Hand Pieces
M5560; M5561; M5562; M5563; M5570; M5571; M5572; M5573; M5580; M5581; M5582;
M5583; M5584; M5590; M5591; M5592; M5593; M5594
Irrigation / Aspiration Cannulae
M6600; M6601; M6602; M6605; M6611; M6614; M6615; M7900A; AU4600

Manufacturer Sterimedix Ltd
1 Madeley Road, North Moons Moat, Redditch
Worcestershire, B98 9NB, UK

Notified Body Notified Body 0120
SGS United Kingdom Ltd
202b Worle Parkway, Weston-super-Mare, BS22 6WA

Intended Use Surgically invasive device for introducing Balanced Salt Solution (BSS) into the anterior chamber to maintain the shape, and aspirate fluid and cortical debris. Some bimanual aspiration devices incorporate a micro etched tip so that they can be used for gently polishing the capsule. They are intended for transient use.

Sterile Sterile - EO. Approved by SGS United Kingdom Ltd

Measuring Function No


Conforming to Current Standards EN 556-1
ISO 594-1, 2
BS EN 980
EN 1041
EN 1707
EN ISO 9001
EN ISO 10993-1, 3, 4, 5, 6, 7, 11
EN ISO 11135-1
EN ISO 11607-1, 2
EN ISO 11737-1, 2
EN ISO 13485
EN ISO 14155-1
EN ISO 14937
EN ISO 14971
EN ISO 15223-1

Classification Class IIa in accordance with Annex IX Rule 6

CE Marking Notified Body and Number: SGS 0120
Initial Date of CE Marking: 21/05/1997
Certified to: Directive 93/42/EEC
Certificate Number: GB97/9964
Certificate Issue Date: 30/09/2017
Certificate Expiration Date: 06/02/2020

Assessment Route By Annex II excluding section 4 and Article 11 of the European Communities Council Directive 93/42/EEC amended by Directive 2007/47/EC, and the application of BS EN ISO 13485:2012.

This product does not contain human blood derivatives and is not manufactured utilising tissues of animal origin.

Signed  Date 19/12/18
Name: Andrew Ellis
Position: QA/RA Manager