

# Sterimedix Ltd

Quality System Document QD 2040 Issue P  
EC Declaration of Conformity

## Sterimedix Ltd. Declaration of Conformity for Hydrodissection Cannulae

Ref: DOC05

16/11/2018 - Issue 9

### 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.

<b>General Product Name</b>	Hydrodissection Cannulae	
<b>Product Reference</b>	<b>Hydrodissection Cannulae</b> M2273C; M2273D; M2273E; M2273F; M2273H; M2274; M3900; M3900A; M3901; M3902; M3904; M3906A; M3908; M3911; M3911A; M3912; M3916; M3919; M3920; M3992; SD5295; SD5158; SD4803; SD4802; SD3902; SD5003; SD5037; SD5099; SD5155; AU0003; AU3424 <b>Hydrolineation Cannulae</b> M3907; M3921 <b>Viscoexpression Cannulae</b> M3899; M3899A; M3899B; AU5090	
<b>Manufacturer</b>	Sterimedix Ltd 1 Madeley Road, North Moons Moat, Redditch Worcestershire, B98 9NB, UK	
<b>Notified Body</b>	Notified Body 0120 SGS United Kingdom Ltd 202b Worle Parkway, Weston-super-Mare, BS22 6WA	
<b>Intended Use</b>	Surgically invasive device for delivering fluids and viscoelastic to facilitate the separation of the cortex from the nucleus, and for insertion under the nucleus and expressing it using viscoelastic. They are intended for transient use.	
<b>Sterile</b>	Sterile – EO & Non-Sterile. Approved by SGS United Kingdom Limited	
<b>Measuring Function</b>	No	
<b>Conforming to Current Standards</b>	EN 556-1 ISO 594-1, 2 BS EN 980 EN 1041 EN 1707 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
<b>Classification</b>	Class IIa in accordance with Annex IX Rule 6	
<b>CE Marking</b>	Notified Body and Number: Initial Date of CE Marking: Certified to: Certificate Number: Certificate Issue Date: Certificate Expiry Date:	SGS 0120 21/05/1997 93/42/EEC GB97/9964 12/08/2015 06/02/2020
<b>Assessment Route</b>	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 13485:2016.	

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

Signed



Date

16/11/18

Name: Andrew Ellis

Position: QA/RA Manager