



Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

No. CE 550877

Issued To: LifeGlobal Group, LLC

a.d.b.a LifeGlobal 393 Soundview Road Guilford, CT 06437

USA

In respect of:

The design, development and manufacture of sterile medical devices for Assisted Reproductive Technology (ART) procedures; Media with antibiotic and/or protein supplementation for gamete and embryo retrieval, micromanipulation, culture, maintenance, transfer and preservation. Oil for overlay of ART media during gamete and embryo culture and micromanipulation. Plastic dishes for gamete and embryo culture and micromanipulation.

on the basis of our examination of the quality assurance system under the requirements of Council Directive 93/42/EEC, Annex II excluding section 4. The quality assurance system meets the requirements of the directive. For the placing on the market of class III products an Annex II section 4 certificate is required.

For and on behalf of BSI, a Notified Body for the above Directive (Notified Body Number 2797):

Gary E Slack, Senior Vice President Medical Devices

Gary C Stade

First Issued: **2010-04-23** Date: **2020-05-11** Expiry Date: **2024-05-26**

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Validity of this certificate is conditional on the quality system being maintained to the requirements of the Directive as demonstrated through the required surveillance activities of the Notified Body. This approval excludes all products designed and/or manufactured by a third party on behalf of the company named on this certificate, unless specifically agreed with BSI.





Supplementary Information to CE 550877

Issued To:

LifeGlobal Group, LLC a.d.b.a LifeGlobal 393 Soundview Road Guilford, CT 06437 USA

Number	Device Name	Intended purpose per IFU		
Class III				
	LG Global			
	LG Global for Fertilization			
	LG Global w/ HEPES			
	LG Global Collect			
	Global Blastocyst Fast Freeze Thawing Kit See CE 549386			
	Global total LP			
	Global total LP w/HEPES			
	Global total LP for Fertilization			
	HSA			
Class IIa	70	2000		
SMD 0109	IVF Media (Oil)	-/		
SMD 0109	ART Dishes			

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This certificate was issued electronically and is bound by the conditions of the contract.





Directive 93/42/EEC on Medical Devices, Annex II excluding Section 4

List of Significant Subcontractors

Recognised as being involved in services relating to the product covered by:

Certificate No: **CE 550877**Date: **2020-05-11**

Issued To: LifeGlobal Group, LLC

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Subcontractor: Service(s) supplied

CooperSurgical Canada 24 Norwich St. E Guelph Ontario N1H 2G6 Canada

Manufacture

Manufacture

CooperSurgical, Inc. 95 Corporate Drive Trumbull

Trumbull Connecticut 06611 USA

Crucial Supplier

Fagron Belgium NV Venecoweg 20 A Nazareth 9810 Belgium

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Medicinal Substances

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List of Significant Subcontractors

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USA

Subcontractor: Service(s) supplied

FertiPro N.V. **Manufacture**Industriepark Noord 32

Fujian Fukang Pharmaceutical Co. ,Ltd No. 6 Gaogang Avenue Jiangyin Industrial Estate Fuqing

Fuzhou 350309 Fujian China

8730 Beernem Belgium

LifeGlobal Europe EU Representative

Rue de la Presse 4 1000 Brussels Belgium

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Subcontractor:

Octapharma AB Lars Forssells gata 23

112 75 Stockholm Sweden Service(s) supplied

Crucial Supplier

Thermo Fisher Scientific Inc. 75 Panorama Creek Drive Rochester New York 14625-2385 USA **Manufacture**

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EC Certificate - Full Quality Assurance System Certificate History

Certificate No:

CE 550877

Date:

2020-05-11

Issued To:

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Date	Reference Number	Action
23 April 2010	7374717	First issue.
02 December 2011	7374715	Extension of Certificate Scope to include devices containing ancillary antibiotic.
28 May 2013	7804078	Scope extension to include Pioneer Pro-Pump for Aspiration.
01 October 2013	8067356	Change of company name from 'genX International Inc' to 'LifeGlobal, LLC, a wholly owned subsidiary of LifeGlobal Group, LLC. Change of name of genX Scientific LLC, Ontario to Lifeglobal LLC.
27 March 2014	8123787	Change of company name from 'LifeGlobal, LLC, a wholly owned subsidiary of LifeGlobal Group, LLC' to 'LifeGlobal Group, LLC, a.d.b.a LifeGlobal'.
18 November 2014	8239163	Change of EU Representative name 'IVFonline Europe' to 'LifeGlobal Europe'. Removal of significant subcontractor 'Embryotech Laboratories, Inc.' and 'design' added to the activities of significant subcontractor 'LifeGlobal, LLC'.
07 April 2015	8292497	Certificate renewal. Addition of 'Fagron Industry, 8790 Waregem' as significant subcontractor.

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Information and Contact: BSI, Say Building, John M. Keynesplein 9, 1066 EP Amsterdam, The Netherlands Tel: + 31 20 346 0780 BSI Group The Netherlands B.V. registered in The Netherlands under 33264284.

This certificate was issued electronically and is bound by the conditions of the contract.





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Date	Reference Number	Action
07 June 2017	8728102	Addition of laboratory dishware and labware to scope. Addition of Thermo Fisher Scientific as a significant subcontractor.
29 August 2017	8789185	Addition of media with protein supplementation. Addition of Octapharma AB, Sweden as crucial supplier.
19 February 2019	9719698	Administrative Subcontractor Service wording update from 'Sterile Manufacture' to 'Manufacture' for Thermo Fisher Scientific Inc., New York 15625-2385, USA. Traceable to NB 0086.
09 May 2019	9658394	Addition of subcontractor CooperSurgical, Inc., Trumbull, Connecticut, USA for manufacture of vacuum pumps. Correction to LifeGlobal Guelph address.
11 December 2019	3095439	Name change from LifeGlobal, LLC, Guelph, Canada, to CooperSurgical Canada, Guelph, Canada. Removal of design activity from Guelph Canada facility.

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Date	Reference Number	Action
Current	9770430	Certificate renewal.
		Name and address updated for crucial supplier Fagron Industry.
		Address updated for crucial supplier Octapharma AB.
		Fujian Fukang Pharmaceutical Co.,Ltd added as significant subcontractor (Medicinal substances) as the manufacturer of gentamicin sulphate
		Removal of vacuum pumps for aspiration (Pioneer Pro Pump) from scope.
		Removal of class Is dishware and labware from scope.
		Removal of media without antibiotic and protein supplementation from scope.
		Added plastic dishes for gamete and embryo culture or micromanipulation to scope.
		The scope has been reformatted to better reflect the certified devices.

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