

## Declaration of Conformity

This European Declaration of Conformity is issued under the sole responsibility of the manufacturer.

### MANUFACTURER

Name of Company	Address	SRN
SteriLance Medical(Suzhou)Inc.	No.168 PuTuoShan Road,New District,215153 Suzhou,Jiangsu, PEOPLE'S REPUBLIC OF CHINA	CN-MF-000002860

### AUTHORIZED REPRESENTATIVE

Name of Company	Address	SRN	Phone/email
Emergo Europe B.V.	Westervoortsedijk 60 6827 AT Arnhem The Netherlands	NL-AR-000000116	+31.70.345.8570 EmergoEurope@ul.com


### PRODUCT IDENTIFICATION

Product Name	Type	EMDN Code
Heel Incision Safety Lancets	SteriHeel,SteriHeel Plus, Neoheel,OctaHeel	V010401
Intended Purpose		Basic UDI-DI
It is used for blood sampling form heels of infants during blood tests.		6945630130BG

### RISK CLASS FOR MEDICAL DEVICES

Device Classification	Common Specifications / Standards	
<b>Class:</b> Ila	Medical Devices Regulation (EU) 2017/745	
<b>Rule:</b> 6		

### NOTIFIED BODY

Name of Company	ID Number	Conformity Assessment Procedure	Certificate Reference(s)
TÜV SÜD Product Service GmbH	 0123	Medical Devices Regulation (EU) 2017/745 ,Annex IX Chapters I and III	Certificate No.: G10 093119 0001 Rev.01 Valid from: 2023-06-19 Valid until: 2027-11-23

The SteriLance Medical(Suzhou)Inc. declares that the above-mentioned products meet the provision of the following EU legislation:

Medical Devices Regulation (EU) 2017/745

**COMPANY REPRESENTATIVE: Cao Yueping**

**TITLE: PRRC**

**SIGNATURE:**

*Cao Yueping*

**PLACE: Suzhou**

**DATE:**

*2023-06-19*