

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.	Prinsessegracht 20,2514 AP,The Hague,The Netherlands	NL-AR-00000011 6	+31.70.345.8570 EmergoEurope@ul.com	

PRODUCT IDENTIFICATION			
Product Name	Туре	Code / Catalog Number	
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	

RIS	K CLASS FOR	R MEDICAL DEVICES
Device Classifi	cation	Common Specifications / Standards
Class:	lla	Medical Devices Regulation (EU) 2017/745
Rule:	6	

NOTIFIE	D BODY		
Name of Company	ID Number	Conformity Assessment Procedure	Certificate Reference(s)
TÜV SÜD Product Service Gmbh	C € ₀₁₂₃	Medical Devices Regulation (EU) 2017/745 ,Annex IX	Certificate No.: G10 093119 0001 Rev.00
		Chapters I and III	Issue date: 2022-11-24
			Valid until: 2027-11-23

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

SIGNATURE:

Medical Devices Regulation (EU) 2017/745

COMPANY REPRESENTATIVE: Zhang Xuanchao

PLACE: Suzhou DATE:

Thong Luanchao Lon-12-29

TITLE: Quality Manager