

## **EUROPEAN MEDICAL DEVICE REGULATION**

## **Declaration of Conformity**

As Legal Manufacturer, we

3M Company Single Registration Number, US-MF-000014086 2510 Conway Ave. St. Paul, MN 55144 USA

hereby declare under our sole responsibility that the following CE marked device(s)

Trade Name	Steri-Strip <sup>TM</sup>
	Steri-Strip™ Elastic Skin Closures
Intended Purpose	Steri-Strip Skin Closures are intended for use as skin closure
	devices in the treatment of lacerations and surgical incisions. They
	may be used in conjunction with skin sutures and staples or after
	their removal for wound support.
Reference	4541-12
Basic UDI-DI	06082238401010000000071AH

is classified per rule 4 of Annex VIII of the Medical Device Regulation (EU) 2017/745, as Class Is devices in accordance with Annex IX and all other applicable provisions of the REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL concerning medical devices.

This declaration is made based on the quality assurance certificate EC Certificate Number: MDR 725202 Issued by: BSI, 2797

The Authorized European Representative for the concerned device(s) is

EU Representative Address
3M Deutschland GmbH
Health Care Business
Single Registration Number, DE-AR-000011642
Carl-Schurz-Str. 1
41453 Neuss, Germany

Granne Stils 22 November 221

Dianne Gibbs, RAC Regulatory Affairs Director 3M Medical Solutions Division Date

3M and Steri-strip are a trademark of 3M.