

**EUROPEAN MEDICAL DEVICE REGULATION****Declaration of Conformity**

As Legal Manufacturer, we

3M Company
Single Registration Number (TBD)
2510 Conway Ave. St. Paul, MN 55144 USA

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

Trade Name	Nexcare™ Waterproof Bandages Nexcare™ Aqua Clear Waterproof Bandages Nexcare™ Aqua Clear MAXI Waterproof Bandages Spofaplast™ Tattoo™ Waterproof Viscoplast™ Waterproof Tattoo™ Sample Pack Tattoo™
Intended Purpose	Bandages are used to cover and protect minor wounds
Reference	582-10DN 586-20DN 588-30DN N0610NAKDMN N1205DMN N126ASDX02N N1214ASD01N 12100HD1N 12100HD2N 115N V5D28K10N 02N
Basic UDI-DI	0608223840105000000009E8

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

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

3M Deutschland GmbH
Health Care Business
Single Registration Number (TBD)
Carl-Schurz-Str. 1



41453 Neuss, Germany

Bryan Becker, Division Quality and Regulatory Leader
3M Company
2510 Conway Ave. St. Paul, MN 55144 USA

09 MARZO 21

Date

3M, Nexcare, Spofaplast, and Viscoplast are trademarks of 3M.