



Declaration of Conformity

As Legal Manufacturer
We, 3M Company, 3M Health Care,
3M Center, 2510 Conway Ave, Bldg. 275-5W-06
Saint Paul, MN 55144 USA
hereby declare under our sole responsibility
that the CE marked products to which this declaration relates,

- 3M™ Steri-Strip™ Skin Closures (Reinforced)
R1540, R1540-01, R1540-02, R1540-12, R1541, R1541-01, R1541-02, R1541-12, R1542, R1542-01, R1542-12, R1546,
R1546-01, R1546-12, R1547, R1547-01, R1547-12, R1548, R1549
- 1540P-1, 1540P-2, 1540P-12, 1541P-1, 1541P-2, 1541P-12, 1542P-1, 1542P-2, 1546P-1, 1547P-1, 1547P-12
1540IP-1, 1540IP-2, 1540IP-12, 1541IP-1, 1541IP-12, 1542IP-1, 1546IP-1, 1546IP-12, 1547IP-1, 1547IP-12
1540NP-2, 1540NP-12, 1541NP-2, 1541NP-12, 1542NP-12, 1546NP-1, 1546NP-12, 1547NP-1, 1547NP-12
1541SP-2
- 3M™ Steri-Strip™ Blend Tone Skin Closures
B1550, B1551, B1553, B1557, B1559, B1551-02, B1551-12, 1551NP-2, 1551NP-12, 1551SP-2
- M™ Steri-Strip™ Elastic Skin Closures
E4540, E4541, E4541-12, E4542, E4546, E4547, E4548, E4549, 4541-12, 4541NP-12
- 3M™ Steri-Strip™ Wound Closure System
W8512, W8514, W8516
- 3M™ Steri-Strip™ Skin Closure Rack, 1555
- 3M™ Steri-Strip™ Compound Benzoin Tincture, C1544
- 3M™ Steri-Strip™ Skin Closures for Consumer
R150C (Nexcare), 5203.531 (M-Plast)
- 3M™ Nexcare™ Steri-Strip™ First Aid Skin Closures
SS08
- Viscoplast Steri-Strip™ Closures
1540R, 1541R, 1542R, R150C, R1551V

are classified,
per Rule 4 of Annex IX of the Medical Device Directive 93/42/EEC, as amended per 2007/47/EC
as a Class Is device
and

is in accordance with Annex(es) V (sterilization) and VII of Directive 93/42/EEC, as amended per 2007/47/EC,
on the approximation of the laws of the European Union Member States concerning medical devices.

This declaration is made on the basis of the quality assurance certificate CE 00493 delivered by BSI, 0086

EU Representative Address
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Carl-Schurz-Str. 1, 41453 Neuss, Germany

Signature:

Date: 12 MAY 2017

Karen Rittle
3M, 3M Health Care
Division Regulatory Manager
Critical & Chronic Care Solutions Division