



3M Deutschland GmbH

Carl-Schurz-Straße 1
41453 Neuss
GERMANY

+49 (0)2131/140
+49 (0)2131/142649
Internet: www.3M.com/de
E-Mail: innovation.de@mmm.com
WEEE-Reg.-Nr. DE 36963167
VAT-ID: DE 120679179

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Declaration of Conformity

We

**3M Deutschland GmbH
Health Care Business
Carl-Schurz-Str. 1
41453 Neuss
Germany**

hereby declare under our sole responsibility
that the CE marked products, to which this declaration relates,

Transpore™ Surgical Tape

**1527-0, 1527-1, 1527-2, 1527-3,
1527-0M, 1527-1M, 1527-2M, 1527-3M**

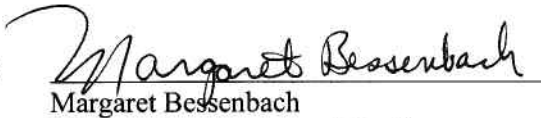
Plastic Tape

1527-0G, 1527-1G, 1527-2G, 1527-3G

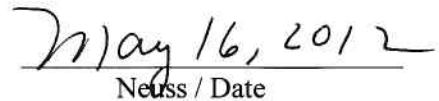
are classified per rule 1 of Annex IX of the Medical Device Directive 93/42/EEC,
as **Class I** devices
and

are in accordance with
Annex VII and all other applicable provisions of the Directive 93/42/EEC
on the approximation of the laws of the European Union Member States concerning medical devices.

Signature:


Margaret Bessenbach

Manager Regulatory and Quality
Europe, Middle East & Africa (EMEA) IP, S&W and FP
3M Deutschland GmbH, Health Care Business


Neuss / Date

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