



3M Deutschland GmbH

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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™ White Surgical Tape**

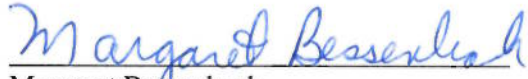
**1534-0, 1534-1, 1534-2, 1534-3**

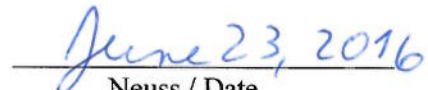
are classified per rule 1 of Annex IX of the Medical Device Directive 93/42/EEC,  
as **Class I** devices

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  
Health Care Business West Europe & CEE  
3M Deutschland GmbH, Health Care Business

  
Neuss / Date

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