

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

As Legal Manufacturer, we

3M Deutschland GmbH  
Health Care Business  
Single Registration Number DE-MF-000011641  
Carl-Schurz-Str. 1  
41453 Neuss  
Germany

hereby declare under our sole responsibility that the following CE marked devices

Trade Name	<b>Tegaderm™ Transparent</b> IV Transparent Film Dressing with Border <b>Tegaderm™ Film</b> Transparent Film Dressing Frame Style <b>Tegaderm™ Film</b> Transparent Film Dressing with Border
Intended Purpose	Dressing can be used to cover and protect catheter sites and wounds, to maintain a moist environment for wound healing or to facilitate autolytic debridement, as a secondary dressing, as a protective cover over at-risk skin, to secure devices to the skin, to cover first and second degree burns and as a protective eye covering.
Reference (Products made in Germany)	<b>1633, 1635, 1655</b> <b>1623W, 1624W, 1626W, 1626, 1630</b>
Basic UDI-DI (Products made in Germany)	0608223276101000000000CB
Reference (Products made in USA)	<b>1610, 1655</b> <b>1622W, 1624W, 1626, 1626W, 1627, 1628, 1629, 1630</b> <b>1614, 1616</b>
Basic UDI-DI (Products made in USA)	06082232761010000000051CU

are classified per rule 4 of Annex VIII of the Medical Device Regulation (EU) 2017/745, as Class 2a sterile 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: 003626 MDR2017Q

Issued by: DQS Medizinprodukte GmbH, No. 0297

Claudia Inden  
Manager Regulatory Affairs  
3M Deutschland GmbH

November 06, 2024

Date