

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	3M [™] Medipore [™] + Pad
Intended	Adhesive Wound Dressing
Purpose	
Reference	3562E, 3564E, 3566E, 3569E, 3570E, 3571E, 3573E
	3562NP, 3566NP, 3569NP, 3570NP, 3562SP, 3566SP,
	3562P-10, 3566P-10, 3569P-10, 3570P-10
Basic UDI-DI	0608223276101000000032CQ

are classified per rule 4 of Annex VIII of the Medical Device Regulation (EU) 2017/745, as Class 1 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 EU Certificate Number: 003626 MDR2017Q

Issued by: DQS Medizinprodukte GmbH, No. 0297

Margaret Bessenbach

August 24, 2022

Date

Manager Regulatory Affairs and Quality
Health Care Business EMEA
3M Deutschland GmbH

3M is a trademark of 3M.

Related to REG-STED-MDR-05-742932