
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 device

Trade Name	3M™ Actisorb™ Silver 220 Activated Charcoal Dressing with Silver
Intended Purpose	Management of all chronic wounds including fungating carcinomas, traumatic and surgical wounds where bacterial contamination, infection or odour occurs
Reference	MAS065, MAS105, MAS105I, MAS190
Basic UDI-DI	06082232761010000000041CR

is classified per rule 14 of Annex VIII of the Medical Device Regulation (EU) 2017/745, as Class III device 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 EU Quality Management Certificate and EU Technical Documentation Assessment Certificate:

EU Quality Management Certificate: 003626MDR2017Q
EU Technical Documentation Assessment Certificate: 31619999MDR2017P

Issued by: DQS Medizinprodukte GmbH, August-Schanz-Straße 21, 60433 Frankfurt am Main, Germany, No. 0297



Margaret Bessenbach
Director Regulatory Affairs and Quality
Health Care Business EMEA
3M Deutschland GmbH

Neuss, January 09, 2024
Location/Date

3M is a trademark of 3M.