

EUROPEAN MEDICAL DEVICE REGULATION

Declaration of Conformity

As Legal Manufacturer, we

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

hereby declare under our sole responsibility that the following CE marked device(s)

Trade Name	Micropore™
Intended	3M [™] Micropore [™] Surgical Tape is a general-purpose
Purpose	gentle tape used to secure dressings, lightweight tubing, and devices to skin.
Reference	1530-0, 1530-1, 1530-2, 1530-3
	1530-0S, 1530-1S, 1530-2S, 1530-3S
Basic UDI-DI	0608223276101000000006CP

are classified per rule 1 of Annex VIII of the Medical Device Regulation (EU) 2017/745, as Class I devices in accordance with all applicable provisions of the REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL concerning medical devices.

Harald Ceschinski Manager Regulatory Affairs and Quality Management System Health Care Business EMEA 3M Deutschland GmbH

3M is a trademark of 3M.

Brd Masch 2020 Date