Version: 1 Status: Release Release Date: 05/25/2020

## 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	Scotchcast™Quick Step Double Sided Felt Roll Splint
Intended	Scotchcast™ Quick Step Double Sided Felt Roll
Purpose	Splint is intended for use in the construction of
	common orthopedic/trauma splints. Specific splinting
	application suitability should be the responsibility of a
	qualified, on-site medical professional
Reference	74002Q, 74003Q, 74004Q, 74005Q
Basic UDI-DI	06082232761010000000025CT

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

Margaret Bessenbach Manager Regulatory Affairs and Quality Health Care Business EMEA

Margaret Bessenbach

3M Deutschland GmbH

3M is a trademark of 3M.

May 25, 2020

Date