

## EUROPEAN MEDICAL DEVICE REGULATION

## **Declaration of Conformity**

As Legal Manufacturer, we

3M Company Single Registration Number: US-MF-000014086 2510 Conway Ave. St. Paul, MN 55144 USA

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

Trade Name	3M <sup>™</sup> Surgical Clipper with Pivoting Head, Charger Stands and
	Starter Kit
Intended	Surgical clipper
Purpose	
Reference	9661L (Surgical Clipper with Pivoting Head)
	9665L, 9668L (Charger Stands)
	9667L-E (Surgical Clipper with Pivoting Head Starter Kit)
Basic UDI-DI	0608223840101000000050A9

are classified per rule 13 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.

3M Company self-declares conformity with Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment, as amended per (EU) 2015/863, and compliance to the requirements of EN IEC 63000:2018.

EU Authorized Representative:

3M Deutschland GmbH
Health Care Business
Single Registration Number: DE-AR-000011642
Carl-Schurz-Str. 1
41453 Neuss, Germany

17 Julnian 272 Date

Dianne Gibbs, RAC

Regulatory Affairs Director 3M Medical Solutions Division

3M is a trademark of 3M.