

EUROPEAN MEDICAL DEVICE REGULATION

Declaration of Conformity

As Legal Manufacturer, we

3M Deutschland GmbH
Health Care Business
Single Registration Number: DE-MF-000012859
Carl-Schurz-Str. 1
41453 Neuss
Germany

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

Trade Name	Ketac Cem radiopaque			
Intended Purpose	e dental luting agent for cementation of indirect restoration			
	and orthodontic bands			
Reference	37200, 37201, 37210, 37211, 37220, 37221, 37230, 37231,			
	37251			
	(mixing block partially included as part of the device)			
Basic UDI-DI	06082232761020000000044DW			

is/are classified per rules 8 and 19 of Annex VIII of the Medical Device Regulation (EU) 2017/745, as Class IIa device(s) 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.

We hereby declare under our sole responsibility that the following CE marked accessory(ies) which are intended to be used together with the above medical device(s)

Accessory(ies)	Reference	Basic UDI-DI	Rule(s) of Annex VIII	Class
Dosing spoon	Not applicable	06082232761020000000087EG	1	ı



is/are classified according to Annex VIII of the Medical Device Regulation (EU) 2017/745 in accordance with all 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 QMS Certificate (MDR): G10 078535 0040 Issued by: TÜV SÜD Product Service GmbH (identification no. 0123)

Dr. Desi W. Soegiarto

Den W. Soegrant

Manager Regulatory Medical Devices

3M Deutschland GmbH

Seefeld/November 11, 2022 Location/Date