


Document Title	<b>Declaration of Conformity</b>	 <b>CMS Dental</b> Our Innovation Your Success	Template	Page
ID	TD-219-05-01-01		F-41 v.14	1 of 2

The products presented by this declaration are in conformity with the listed regulations. The manufacturer is exclusively responsible for this declaration of conformity.

Manufacturer	
Name, registered trade name or registered trademark	CMS Dental A/S
Address	Elmevej 8, 7870 Roslev, Denmark
Single registration number (SRN)	DK-MF-000020260


Product Information	
Product description	Curing Lights
Product and accessories trade names	See appendix List of products
Intended purpose	The Curing Lights is used to light cure dental composites with the activator camphor quinone and/or activator BAPO.
Basis UDI-DI	5713223CUL1XY

Region	Classification (Risk Class & Rule)	Regulation Conformity
EU	Medical Device: Class I, rule 5	European Medical Device Regulation 2017/745
USA	Medical Device: Class I	Code of Federal Regulations, CFR 21 800-898 Medical Devices

Standard compliance	
ISO 13485:2016	Medical devices – Quality management systems – Requirements for regulatory purposes
ISO 14971:2019	Medical devices – Application of risk management to medical devices
EN 60601-1:2006 /A1:2013 /A2:2021	Medical electrical equipment - Part 1: General requirements for basic safety and essential performance
EN 60601-1-2:2015	Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - Collateral Standard: Electromagnetic disturbances - Requirements and tests
ISO 10993-1:2018	Biological evaluation of medical devices – Part 1: Evaluation and testing within a risk management process
ISO 2768-1:2002	General tolerances – Part 1: Tolerances for linear and angular dimensions without individual tolerance indications
ISO 15223-1:2021	Medical devices – Symbols to be used with information to be supplied by the manufacturer – Part 1: General requirements

Competent Authority					
Name	Danish Medicines Agency, Pharmacovigilance & Medical Devices				
Address	Axel Heides Gade 1, 2300 Copenhagen S, Denmark				
Phone	+45 4488 9595				
EUDAMED Actor ID	DK-CA-042				
Conformity assessment procedure	Medical Device Regulation 2017/745				
	Annex II (TD)	Annex III (PMS)	Annex IX, Chapter I (QMS)	Article 19/Annex IV (DoC)	
	Quality Management System – Medical devices				
	ISO 13485:2016 & EN ISO 13485:2016				
Identification of certificate(s)	Quality Management System – Medical devices				
	Certificate	Certificate no.	Issuer (NB ID)	Issued	Valid until
	ISO 13485:2016 & EN ISO 13485:2016	MD 810095	BSI (2797)	2025-01-22	2028-01-22

CMS Dental, the manufacturer, hereby declares that the listed products manufactured after the release date of this declaration of conformity are covered by this declaration of conformity and comply with the above-mentioned regulations, relevant product standards and their transposition into national laws of the member states into which we place the device.

Approval, on behalf of CMS Dental A/S		
Name / Title / Signature	Lisbeth Rose / Quality Assurance Director	
Place / Date of issue	Copenhagen / 2025-01-27	

Appendix – List of products

Products

Product and trade name	Item no.	GTIN/UDI	EUDAMED DI (Basis UDI-DI)
FlashMax P3 Cure and Ortho	100400	5713223000102	5713223CUL1XY
FlashMax P3 Cure and Ortho Wide Spectrum	100403	5713223000201	5713223CUL1XY
FlashMax P7 LAD	100420	5713223000096	5713223CUL1XY

Accessories

Product and trade name	Risk Class/Rule	Item no.	GTIN/UDI
Tips BLUNT d4mm	Class I / rule 5	100501	5713223000652
Tips BLUNT d8mm	Class I / rule 5	100502	5713223000669
Covers	Class I / rule 5	100510	5713223000881
Dental Sleeve Open - 250pcs	Class I / rule 5	100512	5713223000379
Dental Sleeve Closed - 250pcs	Class I / rule 5	100513	5713223000386
Protection Shield	Class I / rule 5	100500	5713223001376
Protection Shield Click-on	Class I / rule 5	100511	5713223001383