EU Declaration of Conformity according to Medical Device Regulation MDR 2017/745 Annex IV

We hereby declare that this EU Declaration of Conformity is issued under the sole responsibility of the manufacturer and the below mentioned products meet the provisions of the EU Regulation above. All supporting documentation is retained on the premises of the manufacturer and the Notified Body.

Products	Bite Fork Bite Fork Set
Project Number(s) / Change Number(s)	n/a
Basic-UDI-DI	763081341XBITE002VW

Legal manufacturer		
CANDULOR	Candulor AG Boulevard Lilienthal 8 8152 Glattpark/Schweiz www.candulor.com	

EU Declaration of Conformity Information		
SRN	CH-MF-000015795	
Intended Purpose	Bite registration	
EU Risk Classification (MDR Annex VIII)	Medical Device Class I	
Conformity Assessment Procedure (MDR Annex IX)	Quality Management System	
MDR Certificate No.	G20 090341 0017 Rev. 00	
Valid until	2025-05-25	

Attachment to EU Declaration of Conformity

Article No.	Description	MDR Classification (EU)	Rule MDR (EU)
662451	Bite Fork	Class I	5, bullet 2
662453	Bite Fork Set	Class I	5, bullet 2

Document Control			
Name	Place of issue	Date of issue	Signature
Approver (PRRC): Alexander Schwaszta	Glattpark, CH	2024-05-22	Sum
Approver (Managing Director): Claudia Schenkel-Thiel	Glattpark, CH	2024-05-22	C. Churchel-Thinz

Document Revision History			
Version	Date	Author	Remark
1.0	2021-05-20	Miriam Stange	First MDR Version
2.0	2021-12-02	Miriam Stange	Correction of validity date
3.0	2024-05-16	Miriam Stange	Removal of Basic UDI-DI HIBC / use of Basic UDI-DI GS1; Removal of DE SRN: Correction of validity date