

EU Declaration of Conformity for Class_Ila_195395372


| Document Information (OTCS) | |
|-----------------------------|-----------|
| OTCS – Doc.-ID | 195395372 |
| Version | 3.0 |

| Signatures | |
|-------------|---|
| Author(s) | The signatures of all involved signatories are added on the last page of this document. |
| Reviewer(s) | |
| Approver(s) | |

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 | Preference anterior, Preference posterior |
|--------------|---|
| Basic-UDI-DI | 763081342APROS002VE |

| Legal manufacturer | | |
|---|---|--|
|  | Candulor AG Boulevard Lilienthal 8 8152 Glattpark/Schweiz www.candulor.com | Phone: +41 (0) 78 694 3311 www.candulor.com Legal Form: Joint Stock Corporation Corporate Headquarters: 8152 Glattpark Registration No.: CHE-107.821.754 VAT No.: CHE-107.821.754 |

| EU Declaration of Conformity Information | |
|--|---|
| EC-REP | SRN: DE-AR-000005472 Candulor Dental GmbH Am Riederngraben 6 78238 Rielasingen-Worblingen Germany |
| SRN (Legal Manufacturer) | CH-MF-000015795 |
| Intended Purpose | Replacement of missing teeth with removable dentures |
| Category (MDCG 2019-14) | MDN 1209 Non-active non-implantable dental materials |
| EMDN Code + term | Q010699 Materials for the preparation of custom-made dental devices - other |
| MDS Code | MDS 1007 |
| MDT Code | MDT 2002 MDT 2006 MDT 2011 |
| EU Classification | <input checked="" type="checkbox"/> Medical Device |
| EU Risk Class (MDR Annex VIII) | Class IIa CE 0123 |
| Conformity Assessment Procedure (MDR Annex IX) | Quality Management System |
| Notified Body Address | TÜV SÜD Product Service GmbH Ridlerstrasse 65 80339 Munich Germany |
| EC Certificate No. | G20 090341 0017 Rev. 00 |
| Valid until | 2028-11-01 |

Attachment to EU Declaration of Conformity

| Article No. | Description | MDR Classification (EU) | Rule MDR (EU) |
|-------------|----------------|-------------------------|---------------|
| 576697 | Preference 8er | Ila | 5 |
| 576818 | Preference 6er | Ila | 5 |

| Revision History | | | |
|------------------|------------|---------------------|-------------------|
| Version | Date | Author | Remark |
| 1.0 | 2024-05-15 | Miriam Stange | First MDR Version |
| 2.0 | 2024-10-21 | Alexander Schwaszta | APM 110593 |
| 3.0 | 2025-02-11 | Alexander Schwaszta | APM 111149 |

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Document Approval (OTCS)

UserName: Sara Marconcini (CHMAS)
Title: Communication Manager
Date: Wednesday, 12 February 2025, 07:45 W. Europe Daylight Time
Meaning: I have reviewed and hereby APPROVE the content
and properties of this document(s) in my role as Creator
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UserName: Alexander Schwaszta (LISCHA)
Title: Director of QM/RA
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UserName: Claudia Schenkel-Thiel (SCHECLA)
Title: Managing Director Candulor
Date: Monday, 17 February 2025, 10:41 W. Europe Daylight Time
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