



CERTIFICATE



This is to certify that the company

Polident d.o.o.

Volčja Draga 42 5293 Volčja Draga Slovenia

with the organizational units/sites as listed in the annex

has implemented and maintains a Quality Management System.

Scope of certification: Design and Development, Manufacturing and Distribution of Artificial teeth and CAD/CAM discs -AUS (a), CND, USA (a,b,c,d)

Through an audit, documented in a report, performed by DQS Medizinprodukte GmbH, it was verified that the management system fulfills the requirements of the following standard:

ISO 13485 : 2016

including applicable country-specific regulatory requirements, as indicated below the scope (full references of abbreviations are listed in the annex)

Certificate registration no.	092246 MDSAP16
Certificate unique ID	1000129515
Effective date	2023-10-30
Expiry date	2026-10-29
Frankfurt am Main	2023-10-11

DQS Medizinprodukte GmbH

Mb luna

Sigrid Uhlemann Managing Director



Marc Goedecke

Product Manager



August-Schanz-Straße 21, 60433 Frankfurt am Main, Tel. +49 (0) 69 95427-300, info-med@dqs.de DQS Medizinprodukte GmbH is recognized under the Medical Devices Single Audit Program. Visit https://www.dqs.de/en/customer-database/ to validate this certificate. The validity of this certificate can only be verified by the QR-code.





Annex to certificate Certificate registration No.: 092246 MDSAP16 Certificate unique ID: 1000129515 Effective date: 2023-10-30

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Volčja Draga 42 5293 Volčja Draga Slovenia

Audited site

REPs FEI No.: site scope and country-specific requirements

092246

Polident d.o.o. Volčja Draga 42 5293 Volčja Draga Slovenia Design and Development, Manufacturing and Distribution of Artificial teeth and CAD/CAM discs -AUS (a), CND, USA (a,b,c,d) REPs FEI No.: F004982





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Full references of country-specific requirements of MDSAP participating Regulatory Authorities

Abbreviation	Jurisdiction	Reference
AUS	Australia	 (a) Therapeutic Goods (Medical Devices) Regulations 2002, Schedule 3, Part 1 – Full Quality Assurance Procedure (b) Therapeutic Goods (Medical Devices) Regulations 2002, Schedule 3, Part 4 – Production Quality Assurance Procedure
BRA	Brazil	RDC ANVISA n. 665/2022 RDC ANVISA n. 551/2021 RDC ANVISA n. 67/2009
CND	Canada	Medical Device Regulations SOR/98-282, Part 1
JPN	Japan	MHLW Ministerial Ordinance No. 169 (2004) as amended by MHLW Ordinance No. 128 (2014), Articles 4 to 68 Japan PMD Act (as applicable)
USA	United States	 (a) 21 CFR Part 803 (b) 21 CFR Part 806 (c) 21 CFR Part 807 (d) 21 CFR Part 820 (e) 21 CFR Part 821