EC Declaration of Conformity

We, the

P.L. Superior Dental Materials GmbH Stiller Weg 15a 22607 Hamburg

SRN. DE-MF-000005251

hereby declare under our sole responsibility that the medical device with the intended purpose: creating of dental impression.

belonging to the

UMDNS-group 16-679, dental impression materials, silicone rubber, GMDN-code 35866, Silicone dental impression material, EMDN-code Q010201, dental impression materials, alginates and others,

namely

Implant Sil,

REF: MF.002618. basic UDI-DI: ++EPLSNAS069C9

is in compliance with the requirements of the Regulation (EU) 2017/745 of the European Parlament and of the Council.

Conformity Assessment

Procedure:

According to Annex IX; Chapter I Section 2 of the Regulation (EU) 2017/745.

Classification:

According to Annex VIII, Chapter III, Rule 5 & 19 of the Regulation (EU) 2017/745: Class IIa.

Reference Standards:

EN ISO 13485:2016 + AC:2018 + A11:2021 - Medical devices - Quality management systems - Requirements for regulatory purposes

EN ISO 14971:2019 + A11:2021 - Medical devices - Application of risk management to medical devices

EN ISO 15223-1:2021 - Medical devices - Symbols to be used with medical device labels,

labelling and information to be supplied - Part 1: General regirements

EN ISO 20417:2021 - Information to be supplied by the manufacturer

ISO 10993 applicable parts - Biological evaluation of medical devices

EN ISO 7405:2018 - Dentistry - Evaluation of biocompatibility of medical devices used in dentistry

EN 1641:2009 - Dentistry - Medical devices for dentistry - Materials

EN ISO 4823:2021 - Dentistry - Elastomeric impression and bite registration materials

This declaration is effective for products placed on the market as of the date of issue. Any modifications of the device not authorized by above declarant will invalidate this declaration.

Notified Body: DNV Medcert GmbH Pilatuspool 2

20355 Hamburg

Identification-Number: 0482

Certificate No.: 1649GB448231211

Validity: 08 December 2027

Hamburg, 2024-06-04

Place, date

Dr. Marian Zasny

Authorised Person for Regulatory Compliance