## **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: light cured dental composite filling

belonging to the

UMDNS-group 16-736, composite restorative material kits, dental, light cured, GMDN-code 35870, Dental composite resin, EMDN-group Q010101, dental restorative devices,

namely

## NanoFill X,

REF: MF.000759

basic UDI-DI: ++EPLSNAS182C6

is in compliance with the requirements of the Directive 93/42/EEC and in accordance with Article 120 of the Regulation (EU) 2017/745, including the amendments made by Regulation (EU) 2023/607, Article 1.

## Conformity Assessment

Procedure:

According to Annex II excluding section 4 of the Directive 93/42/EEC.

Classification:

According to Annex IX Rule 8 of the Directive 93/42/EEC: 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 4049:2019 - Polymer-based restorative materials

Notified Body:

Medcert Zertifizierungs- und Prüfungsgesellschaft für Medizin GmbH

Pilatuspool 2

20355 Hamburg

Identification-Number: 0482

Certificate No.: 1649GB410190708

Validity: 31 December 2028

Hamburg, 2024-06-04

Place, date

Dr. Marian Casny

Authorised Person for Regulatory Compliance

PRR