

## **EU - Declaration of Conformity for Medical Devices**

according to Annex IX Chapter I of Regulation (EU) 2017/745 on medical devices

Registration number (SRN):

Manufacturer:

DETAX GmbH
Street:
Carl-Zeiss-Straße 4
City:
76275 Ettlingen
Country:
Germany

We hereby declare under our sole responsibility that the products in the Annex with the Basic UDI-DI

++EDET0866A

and the classification according to Annex VIII of Regulation (EU) 2017/745

Class IIa, rule 5, sentence 1, 3rd indent

and the intended use:

Surface finish for earmolds or silicone impressions

comply with the relevant provisions of the following EU legislation:

Regulation (EU) 2017/745 on medical devices

as well as the standards and common specifications:

DIN EN ISO 15223-1:2022 DIN EN ISO 20417:2022 DIN EN ISO 10993-1:2021

DIN EN 62366-1:2021 DIN EN ISO 14971:2022

with the participation of the following management systems

**DIN EN ISO 13485:2021** 

and the conformity assessment procedure referred to in the header

by the notified body: 0483

mdc medical device certification GmbH

Kriegerstr. 6

D-70191 STUTTGART

This declaration is valid until a significant change in the specification or manufacture of the product or until a change in the above-mentioned standards, common specifications, management systems and legal regulations, maximally until the expiry of the certificate with the No.:

D1002700058 in force until 31.07.2028

Ettlingen, 18.07.2024

Thorsten Preiss Managing Director Dr. Sönke Rössing Managing Director



Annex to

**EU-Declaration of Conformity for Medical Devices** 

Date: 18.07.24

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**Basic UDI-DI:** ++EDET0866A

## Distributed by: Egger Otoplastik +, DEUTSCHLAND

<b>UDI-DI</b>	Article
+EDET033291	Lack L, 30 ml
	Egger ArtNr. 375030
+EDET033441	Lack L, 100 ml
	Egger Nr. 37501
+EDET033461	Lack L, 500 ml
	Egger ArtNr. 37503
+EDET045111	Lack L, 250 ml
	Egger ArtNr. 37502
+EDET045151	Lack L, 1000 ml
	Egger ArtNr. 37506



