

EU Technical Documentation Assessment Certificate

We hereby certify that the company

Ivory Graft Ltd.
Diamond Tower
Tuval St. 21, Floor 13, #1380
Ramat Gan 5252236
Israel

has submitted a technical documentation in accordance with Annexes II and III of Regulation (EU) 2017/745, which meets the following requirements:

Annex IX – Chapter II (Assessment of the Technical Documentation)

of Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices.

This certificate from mdc medical device certification GmbH (Notified Body 0483) consists of 2 pages. Details of the devices covered as well as further information and conditions are contained on the following pages.

Valid from 2026-01-21
Valid until 2027-08-29

Registration No. D1478900012
Report No. P23-00876-272079

Stuttgart, 2026-01-21



Notified Body



EU Authorized Representative:

MedEnvoy Global BV
Princes Margrietplantsoen 33, Suite 123,
2595 AM The Hague
Netherlands
NL-AR-000024028

Devices:

Ivory Dentin Graft

Particel size: 300 – 900 µm
Vials: 0.25 g, 0.5 g, 1.0 g, 2.0 g, 5.0 g
Syringes: 0.5 g, 1.0 g

Particel size: 300 – 500 µm
Vials: 0.25 g, 0.5 g

Intended purpose:
A sterile bone graft material for the repair or augmentation of bone defects in dental procedures

Risk class: III
Basic UDI-DI: 7290018472IDG001VG

Notes:

For the placing on the market of the devices an EU Quality Management System Certificate according to Annex IX, Chapter I of Regulation (EU) 2017/745 on medical devices is also required.

The certificate is based on the previous certificate

D1478900004 (2022-08-30)
D1478900007 (2024-04-22)
D1478900011 (2025-10-02)

with the following changes to D1478900011:
Supplemented: Including the new variants with smaller particle size