

EU Declaration of Conformity Class I

Neuhausen, 2023-09-18

We herewith declare under our sole responsibility that the Class I medical devices listed below, first placed on the market by IVF HARTMANN AG, satisfy the applicable provisions, in particular the General Safety and Performance Requirements, of the Regulation (EU) 2017/745 of the European Parliament and of the Council of 5. April 2017 on medical devices.

The conformity assessment procedures according to Article 52 (7) have been performed and the Technical Documentation is kept available.

Single Registration Number of Manufacturer: CH-MF-000015962

Swiss Single Registration Number of Manufacturer: CHRN-MF-20000305

High Level Intended Purpose	Non-active, non-implantable devices for wound and skin care		
Product Name	Product Group Number	Classification Rule (according to Annex VIII of Regulation (EU) 2017/745)	Basic UDI-DI
DermaPlast Sparablanc Military (scope see Table 1)	3690	Rule 1	76116003690ND

IVF HARTMANN AG

i.V.



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Valid until: 2026-OCT-06

Table 1: Scope

REF	Description
988610	DermaPlast Sparablanc transp 1,25cmx9,2m
988620	DermaPlast Sparablanc transp 2,5cmx9,2m