



EU DECLARATION OF CONFORMITY according to Regulation (EU) 2017/745

EN

Manufacturer: FIAB SpA

Registered address: Via Costoli 4, 50039 Vicchio (FI), Italia

Single Registration Number: IT-MF-000005988

Basic UDI-DI: 80330032610800004NW

Product name/ Intended Purpose: Synthetic buckskin electrodes holding bags

Models: See list in Attachment

Technical Documentation File: TDF 108

Risk Class (MDR Annex VIII): I

Conformity assessment procedure performed: Annex IV (EU Declaration of Conformity)

Technical standards and/or
Common Specifications applied:

EN 1041 [2008/A1:2013] - EN ISO 10993-1 [2018] - EN
ISO 13485 [2016] - EN ISO 14971 [2019] - EN ISO
15223-1 [2016]

With this Declaration of Conformity, issued under the sole responsibility of FIAB SpA as the Manufacturer, we hereby declare

- that the medical devices specified meet the provision of the Regulation (EU) 2017/745 for medical devices
- that the procedures of FIAB quality management system according to ISO 13485 have been followed, Certificate of Registration no.MD77846 issued by BSI
- that the products do not contain medicinal substances, elements of animal origin or their derivatives, human blood derivatives, and are latex free

Signature:

Vicchio, 08/10/2021

Alberto Calabrò
Managing Director

Declaration Code EU-00000127-108

First issued: 08/10/2021

Cod 99500038MD4B

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Attachment of EU Declaration of Conformity – List of models

PG912 - PG916 - PG917 - PG917/MTL - PG917/MTS3 - PG970 - PG980
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