

# CE

#### EU DECLARATION OF CONFORMITY No. DPDOC/12/FAB

Elaborated in accordance with Annex IV of the Medical Device Regulation 2017/745 of EU Parliament and the Council

MANUFACTURER: DEVOLPE s.r.o.

REGISTERED TRADE MARK: **DEVOLPE** 

REGISTERED ADDRESS: Nam. 14. rijna 1307/2, Prague 5, 150 00, Czech Republic

UDI-ID: 859420959BBZN (BACKBAG TYPE FIRST AID BAG), 859420959CPRTS (CPR TYPE FIRST AID

BAG)

NAME OF THE DEVICE: FIRST AID BAGS

MODELS: BACKBAG TYPE FIRST AID BAG, CPR TYPE FIRST AID BAG

CATALOG CODE: FAB/02/BBDP (BACKBAG TYPE FIRST AID BAG), FAB/01/CPRBDP (CPR TYPE FIRST

AID BAG)

LOT NO: CZR23001BB (BACKBAG TYPE FIRST AID BAG), CZR23001CPR (CPR TYPE FIRST AID BAG)

RISK CLASS: Class I

According to Rule 1 of the Chapter III (Classification rules) namely non-invasive device of Annex VIII of the Medical Device Regulation 2017/745 of EU Parliament and the Council

#### MANUFACTURER HEREBY DECLARES:

- that the above specified medical device represents **risk class I** device according to the requirements of Annex IX of the Medical Device Regulation 2017/745 of EU Parliament and the Council.
- that the product complies with the all applicable provisions of Annexes of the Medical Device Regulation 2017/745 of EU Parliament and the Council concerning medical devices and the product's conformity is applicable to conformity procedure of the abovementioned legislation.

### FURTHER IT IS DECLARED THAT THE PRODUCT HAS BEEN:

designed, developed, manufactured and tested in accordance with all applicable standards mentioned below and by using a quality management system according to DIN EN ISO 13485:2016 for medical devices.

The above mentioned products have met the requirements of the medical device for placing CE marking and ISO 13485:2016 for medical devices.

The current declaration of conformity has been drafted in accordance with the requirements of the Annexes IV and V of the Medical Device Regulation 2017/745 of EU Parliament and the Council.

## STANDARDS APPLIED TO THE PRODUCT:

- ISO 13485:2016 Quality management system for medical device
- EN 1041:2008 + A1: 2013 information supplied by manufacturer
- EN ISO 15223:2016 Medical devices Symbols to be used with medical device labels, labelling and information to be supplied
- EN 14971:2012 Medical devices Application of risk management to medical devices

AFFIRMED BY THE MANUFACTURER'S REPRESENTATIVE

In Prague, on February 3, 2023

Devoipe s.r.o.
nám. 14. října 1307/2
150 00 Prague 5, Czech Republic
10: 09776885, TIN: CZ09776885
info@devolpe.com | www.devolpe.com