

EU - DECLARATION OF CONFORMITY

according to Regulation (EU) 2017/745 article 19 read in conjunction with annex IV

Hereby the manufacturer



anmed

Am Gewerbegebiet 5 09474 Crottendorf Federal Republic of Germany

SRN: DE-MF-000006213

declares under sole responsibility that the named product

ANABOX[®] 7×4

ALSO, THE BRAND NAMES CINIBOX AND SUPAIRBOX MEDICATION DISPENSER FOR 1 WEEK

BASIC UDI-DI: 4260041530475.7X42Z

complies with the relevant provisions of Regulation (EU) 2017/745 of April 5, 2017 on medical devices.

A conformity assessment procedure was carried out in accordance with article 52 (7) of Regulation (EU) 2017/745. In accordance with annex VIII rule one of Regulation (EU) 2017/745 the product mentioned above is classified as MEDICAL DEVICE RISK CLASS I

INTENDED PURPOSE:

Medication dispensers to alleviate or compensate for injuries or disabilities

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116

119 / HRB 18717 Chemnitz

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CROTTENDORF, 2024/01/01

place, date

SEBASTIAN RICHTER operating manager / proxy