

EU Declaration of Conformity

The manufacturer:

EEC

Address: Via Verdi, 39 - 42043 Gattatico (RE) - Italy

SRN: IT-MF-000027951

Declares, under its own and exclusive responsibility, that the device(s)

| Code | Model | ID BD/RDM | BASIC UDI-DI |
|---|--|-----------|---------------------|
| 72850047 | OXYGEN CYLINDER HOLDER | 2322219/R | 80557742072850047KZ |
| Intended use: | The device is intended to be installed on Pam Mobility stretchers to support the oxygen cylinder. Environment of use: within healthcare and health facilities. The device cannot be used in a potentially explosive or flammable atmosphere. Personnel intended for use of the product: specialist operators and doctors. | | |
| Risk class: | Class I (in accordance with Rule 1, Annex VIII of Regulation (EU) 2017/745) | | |
| It complies with the following European Union legislative acts: | | | |
| (EU) 2017/745 | Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) no. 178/2002 and Regulation (EC) no 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/ | | |

The device is subject to the conformity assessment procedure provided for in article 52, section 7 of Regulation (EU) 2017/745

Gattatico, 27 april 2023 Managing Director Andrea Muzzini

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