



ID-Nr. 047-MDD-001-007

EC Declaration of Conformity for Medical Devices

In accordance with Annex II excluding (4) of Council Directive 93/42/EEC

We

PARI GmbH Spezialisten für effektive Inhalation

**Moosstraße 3
82319 Starnberg
GERMANY**

hereby declare under sole responsibility that the medical device

**PARI BOY mobile S (Type 047)
(Compressor for Inhalation Therapy)**

GMDN Code: 31253

Risk Classification according to Council Directive 93/42/EEC, Annex IX: IIa

complies with the essential requirements of the

Council Directive 93/42/EEC (Medical Device Directive, MDD) dated 14th June 1993.

This Declaration remains valid at the latest until May 26, 2024).

Compliance has been achieved in conformity with Annex I of the above named Directive.

The applied harmonized standards are listed in the Technical Documentation.

Notified Body: TÜV SÜD Product Service GmbH, Ridlerstraße 65, D-80339 München, Germany

EC Quality System Certificate: Certificate No. G1 011861 0076 Rev. 02 (Expiry: May 26, 2024)



Starnberg, May 12, 2021

Dr. Davia Viellechner
- President -



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Attachment

This Declaration of Conformity is valid for the following configuration:

REF Number	Product Name
047B1010	PARI BOY mobile S

END OF DOCUMENT