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**EU DECLARATION OF CONFORMITY OF MEDICAL DEVICE  
DRAWN UP BY THE MANUFACTURER**

**Manufacturer:** Pro Medicare S.r.l. with registered office and operational headquarters  
in Mesagne (BR) Via Montagna

**General Device Name:** "HIPPO MB Modular Postural Pad for Hippotherapy"

**UDI-DI BASIC:** 805571347HIPPOEM

**SRN:** IT-MF-000022537

**Device Code:**

**D02-701-1** " HIPPO MB Modular Postural Pad with girth closure on the right "

**D02-702-1** " HIPPO MB Modular Postural Pad with girth closure on the left "

**D02-703-1** " Footrest strap for HIPPO MB Modular Postural Pad "

Pursuant to Article 19 of the EU Regulation 2017/745 of the European Parliament and Council, the undersigned, Dr. Franco Cariolo, Sole Director and Legal Representative of Pro Medicare S.r.l., manufacturer of standard class I medical devices according to the rules provided, having performed the conformity assessment as prescribed by the above Regulation and having fully complied with the procedures established in Chapter V Section 2 and Annex II and III,

*ensures and declares that*

- the devices comply with the requirements set out in article 5 and in Annex I, therefore can be lawfully placed on the market and into service after affixing the CE Mark as required in Article 20 of the EU Regulation 2017/745.
- Pro Medicare S.r.l. is responsible for the design, manufacture, packaging, and labeling.

Mesagne (BR), 08/08/2022

Pro Medicare S.r.l.  
Sole Director and Legal Representative

  
  
L'amministratore Unico