

Manufacturer's Declaration of Conformity

Lopital Nederland B.V. Laarakkerweg 9, 5061 JR, Oisterwijk

C∈ marking in accordance with the Medical Device Regulation (EU) 2017/745

The Netherlands

NL-MF-000004372

Manufacturer's name: Manufacturer's Adress:

Manufacturer's SRN

(Single Registration Number):

Document Number: DOC2050-2024-02

Brand Name:	Lopital	
Medical device:	Marina basic / Marina deluxe	
Model number(s): Device Description:	61002050 & 61002060 Mobile Shower Trolley with electrically powered high/ low adjustment	
Basic UDI-DI:	872025610349361002050P3 & 87202561035161002060K2	
Classification:	Class I	
Conforms to regulation:	Medical Device Regulation (EU) 2017/745	
Standards applied:	NEN-EN-ISO 14971:2019	
The product has been tested and	ISO 17966:2016 IEC 60529:2019	
evaluated in accordance with the following standards:		
ronowing standards.	IEC 60601-1:2005+A1:2012 IEC 60601-1-2:2014	
	IEC 60601-1-6:2010+A1:2013	
	IEC 62366:2007	042 / CAN /CCA COO 2 No (0404 4 4 4
	CSA CAN/ CSA-C22.2 No.60601-1-6	012/ CAN/CSA-C22.2 No. 60601-1:14 5:11 AMD 1/IEC 60601-1-6
The product has been designed and manufactured under a certified quality management system in accordance with:	UNI-CEI-EN-ISO 13485:2016	
This declaration of conformity is issued a device(s) specified above meet the pro-	under the sole responsibility of Lopital. Wision of the Regulation (EU) MDR 2017/7	e hereby declare that the medical 45 for medical devices.
Signature:	Date:	Place:
	12-12-2024	Oisterwijk
Jan Van Megen, CEO	dd-mm-yyyy	