

Manufacturer's Declaration of Conformity

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

| Manufacturer's name: Manufacturer's Adress: | Lopital Nederland B.V. Laarakkerweg 9, 5061 JR, Oisterwijk The Netherlands | |
|---|---|------------------------|
| Manufacturer's SRN (Single Registration Number): | NL-MF-000004372 | |
| Brand Name: | Lopital | |
| Medical device: Model number(s): Device Description: Basic UDI-DI: | Tango, Tango XL & Tango XXL 51005700, 51005705 & 51005800 Mobile Shower-Toiletchair 872025610305951005700LK, 8720 872025610315851005800LY |)25610314151005705F7 & |
| Classification: | Class I | |
| Conforms to regulation: | Medical Device Regulation (EU) 20 | 17/745 |
| Standards applied: | NEN-EN-ISO 14971:2019 | |
| The product has been tested and evaluated in accordance with the following standards: | ISO 17966:2016 IEC 62366: 2007 | |
| 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 under the sole responsibility of Lopital. We hereby declare that the medical device(s) specified above meet the provision of the Regulation (EU) MDR 2017/745 for medical devices. | | |
| Signature: | Date: | Place: |
| 1100 | 12-12-2024 | Oisterwiik |

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Jan Van Megen, CEO