

DECLARATION OF CONFORMITY

ACCORDING TO (EU) 2017/745 MEDICAL DEVICE REGULATION

EU Representative

SUNGO Europe B.V.
Olympisch Stadion 24, 1076DE
Amsterdam, Netherlands

SRN: NL-AR-000000247

Conformity Assessment

Conformity Assessment Procedure
Annex II+III of Regulation (EU) 2017/745

Applicable Standards

EN ISO 14971: 2019, EN 1041:2008+A1:2013, EN ISO 15223-1: 2016, ISO 10993-1:2018, EN ISO 10993-5:2009, EN ISO 10993-10:2013, EN ISO 11199-2:2005

Remark

The declaration of conformity is valid in connection with the release technical document ** MER-TCF01, 1

All the supporting documentation is retained at the premises of the manufacturer.

The Declaration of Conformity is exclusively under the sole responsibility of the manufacturer.

Manufacturer

Name: ZHENJIANG RAPID REHAB & INSTRUMENT

CO., LTD.

Address: NO.35 HUANCHENG SOUTH ROAD, PICHENG, DANBEI TOWN, DANYANG CITY,

ZHENJIANG, JIANGSU, CHINA

Product Information

Name: Rollator

Model: See Annex GMDN: See Annex

Basic UDI-DI: -

Classification: Class I, according to Rule 1, Annex

VIII, Regulation (EU) 2017/745

Declaration

We herewith declare that the above-mentioned products meet the requirements of Medical Device Regulation (EU) 2017/745 and the applicable standards above.

Signatur

Name: Wilson Position: GM Place: Zhenjiang / China

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Annex

Product Name	Model	GMDN	Basic UDI-DI
Rollator	RL016, RL016Qe, RL016L, RL016B, RL016F, RL016BF, RL016QF (4*8"), RL016QF (10"+8"), RL008, RL008S, RL008M, RL001, RL001F, RL002, RL003, RL003F, RL004, RL005, RL006, RL009, RL015, RL025, RL025F	37951	-

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Signature:

Position: GM