

Hiermit wird bestätigt, dass die Produkte der
It is hereby confirmed that the products of

Ossenberg GmbH
Kanalstraße 79
48432 Rheine

identisch mit denen der von
are identical with those of

HEBEI Healthplus Medical Device Co., Ltd.
No. 1, Chuangye Street,
Southwest Industrial District,
Matou Ecological Industry Park,
Handan city, Hebei, 056046,
China

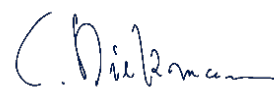
gelieferten Artikel sind.
delivers articles.

<u>Artikel-Nr.</u> <u>HEBEI Healthplus</u> <u>Medical Device Co.,</u> <u>Ltd.</u> <u>article no.</u> <u>HEBEI Healthplus</u> <u>Medical Device Co.,</u> <u>Ltd.</u>	<u>Produktbezeichnung</u> <u>HEBEI Healthplus Medical</u> <u>Device Co., Ltd.</u> <u>Product name</u> <u>HEBEI Healthplus Medical</u> <u>Device Co., Ltd.</u>	<u>Artikel-Nr.</u> <u>Ossenberg GmbH</u> <u>article no.</u> <u>Ossenberg GmbH</u>	<u>Produktbezeichnung</u> <u>Ossenberg GmbH</u> <u>Product name</u> <u>Ossenberg GmbH</u>
- -	Achselstütze Axillary Crutch	12K 12K	Achselstütze 120 kg - klein Axillary Crutch 120 kg - small
- -	Achselstütze Axillary Crutch	12G 12G	Achselstütze 120 kg - groß Axillary Crutch 120 kg - large

Rheine, den 28.06.2021

Ort, Datum / place, date


Carsten Diekmann



Name und Unterschrift Geschäftsführer / Name and signature General Manager

EU Declaration of Conformity

The following description for the medical device,

Device information	Description
Registered trade name and address	<i>HEBEI Healthplus Medical Device Co., Ltd. No. 1, Chuangye Street, Southwest Industrial District, Matou Ecological Industry Park, Handan City, Hebei, 056046, China Tel:+86-0310-2111888</i>
Authorized representative	<i>Y. Sung Handelsvertretung Duesselthaler Str. 24, 40211 Duesseldorf Germany</i>
Common device name	<i>Crutches</i>
Product and trade name	
UMDNS code	<i>11063, Crutches</i>
GMDN code	<i>31115: Axillary Crutch 34977: Walking Aid, One Arm, Crutch</i>
Single Registration Number (SRN)	<i>DIMDI register # 00299449</i>
Basic UDI-DI	<i>697322200MXCRUTCHD7</i>
Risk class of the device	<i>Class I</i>
Common Specification (CS) references	<i>Crutches, Underarm Crutches, Axillary Crutches, Height Adjustable Crutches</i>
Intended purpose (GMDN definition)	<i>A staff-like mobility aid used to assist a person with a disability to support their weight while walking. The device has one leg, a padded hand support at the level of the user's wrist, and an under-arm (the armpit or axilla) padded platform. The pads are typically made of non-slip material (e.g., rubber, glycerin gel, sheepskin). The device may be adjustable in length and is typically made of wood or metal [e.g., aluminum (Al), titanium (Ti)].</i>
Conformity assessment procedure performed and identification of the certificates issued by notified body, if applicable	<i>Quality Management System ISO 13485:2016 by NQA ISO 9001:2015 by NQA</i>
Name and identification number of the notified body, if applicable	<i>ISO 13485, NQA Certificate Number: 35429 ISO 9001, NQA Certificate Number: 46296</i>

that is covered by the present declaration is in conformity with the Medical Device **Regulation 2017/745/EU** and, if applicable, with any other relevant Union legislation that provides for the issuing of this EU declaration of conformity. The device is in conformity with conformity assessment procedure for **Class I devices** that should be carried out, as a general rule, under the sole responsibility of manufacturers in view of the low level of vulnerability associated with such devices. Thus, manufacturers of class I devices, other than custom-made or investigational devices, shall declare the conformity of their products by issuing an EU declaration of conformity referred to in Article 19 “EC declaration of conformity” after drawing up the technical documentation set out in Annexes II and III of the **Regulation**.

For the evaluation regarding Class I device (Risk class in accordance with the Rule 1 set out in Annex VIII of the **Regulation**), the following harmonized standards are applied:

- *EN ISO 13485:2016 Medical devices - Quality management systems - Requirements for regulatory purposes*
- *EN ISO 14971:2012 Medical devices –Application of Risk Management*
- *EN ISO 15223-1:2016 Medical device – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements*
- *EN 62366-1:2015 Medical devices Part 1-- Application of usability engineering to medical devices*

The following Union authorized representative is stated to the declaration:

Y. Sung Handelsvertretung
Duesselthaler Str. 24, 40211 Duesseldorf Germany

(Company name / Registered place of business)

The following person is exclusively responsible for the compliance of declaration:

HEBEI Healthplus Medical Device Co., Ltd.
No. 1, Chuangye Street, Southwest Industrial District, Matou Ecological Industry Park,
Handan city, Hebei, 056046, China

(Manufacturer’s name/ Registered address)

Dickson Su / General Manager

(Name/Function)



(Legal Signature)

March 5, 2020

(Date of issue)