EU Declaration of Conformity according to Annex IV of Regulation (EU) 2017 /745 on medical devices



We, the company

Semeda GmbH

Am Petersberg 36 29389 Bad Bodenteich Germany SRN: DE-MF-000008334

declare in sole responsibility,

that the short bar listed below and described and specified in the technical documentation "TD 018-01 Semeda Mini Steg, kurzer Schienensteg"

Semeda Mini Steg, kurzer Schienensteg – Semeda Mini Steg, Short Bar Semeda Mini Steg, kurzer Schienensteg – Semeda Mini Steg, Short Bar; AB 2800-000

fulfils the general safety and performance requirements of

Regulation (EU) 2017 /745 on medical devices (MDR), Annex 1.

The Semeda Mini Steg, Short Bar is used as an accessory for the foot abduction brace in the system foot abduction orthosis which is used for very small or premature babies diagnosed with idiopathic clubfoot according to the conservative therapy approach of Dr Ponseti.

The orthosis in which the accessory is used serves to maintain the correction of the foot that has been achieved by a plaster corrective. It serves to prevent a relapse and has no corrective effect.

The Semeda Mini Steg, Short Bar is used as a particularly short, lightweight middle piece in the foot abduction brace. This allows to reduce the width and weight of the splint to enable/ease therapy for particularly small patients.

Basic-UDI-DI

426020155ZBHSST6L

Risk class:

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(in accordance with MDR, Annex VIII)

(Rule 1)

Performed conformity assessment

procedure:

MDR, Article 52 (7)

After additions or changes to the product or to the technical documentation the declaration of conformity is reissued.

Bad Bodenteich, 30th August 2023

Place, Date

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