

## **Declaration of Conformity**

## Manufacturer

Tyromotion GmbH Bahnhofgürtel 59, A-8020 Graz, Austria SRN (Single Registration Number): AT-MF-000016168

**Product Specification** 

Article Number and Product Name

UDI-DI

Intended Purpose

Basic UDI-DI Including Software

Components

Classification according to Medical Device Regulation (EU) 2017/745, Annex VIII

(11310) MYRO® 9120077180069

The MYRO® therapy system is generally used for neurological

rehabilitation of the upper extremities.

912007718MYROE4

TyroS Software Version 6.5.x

MYRO® Therapydesk

MYRO® Objects (Coin, Ball, Cup, Grip, Force, Blank)

Class I according to Rule 1 and 13

**Conformity Assessment** 

Conformity Assessment Procedure according to

Certificate

Medical Device Regulation (EU) 2017/745, Annex IX, Chap. I This certificates is valid for devices manufactured in the

manufacturing period given, only.

Valid for

12. Dec. 2024 - 05. Jan. 2028

This Declaration of Conformity is issued under the sole responsibility of the manufacturer.

The products described above, as delivered, comply with the requirements of Regulation (EU) 2017/745 (MDR), the Machinery Regulation (EU) 2023/1230, the Directive 2014/53/EU (Radio Equipment Directive) and the Austrian Medical Devices Act BGBl. I No. 122/2021 as amended.

This declaration is supported by the certification of the Quality Management System according to EN ISO 13485 by DQS Medizinprodukte GmbH. The products are provided with CE and UKCA marking.

CE

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Sabine Seereiner Head of QM & RA

Graz, 12. Dec. 2024