EU- Declaration of conformity for medical devices

(According to Regulation (EU) 2017/745 for medical devices, Annex VIII of May 2017)

HAIDER BIOSWING GmbH
DE-MF-000008765

We hereby declare under our sole responsibility that the sensorimotor therapy device $\mathsf{PROPRIOMED}^{\, \otimes}$

Туре:	BIOSWING [®] PROPRIOMED [®] 100
Basic-UDI-DI:	4260742520051
Serial number:	2021: 2521000000 to 2521999999 2022: 2522000000 to 2522999999

complies with the relevant provisions of the following directive:

Regulation (EU) 2017/745 for medical devices

(further ongoing)

Classification according to Regulation (EU) 2017/745 for medical devices, Annex VIII:

Class I non-active medical device,

according to classification rule 1



95704 Pullenreuth, Germany, May 26th, 2021

Management

Eduard Haider

Technical Manager

Dipl.-Ing. Alexander Haider

BIOSWING Therapiesysteme



HAIDER BIOSWING GmbH | Dechantseeser Straße 4 | 95704 Pullenreuth, Germany Tel.: +49 (0) 92 34 / 99 22 - 0 | Fax: +49 (0) 92 34 / 99 22 - 166 | E-Mail: info@bioswing.de