vibro sonic

CE DECLARATION OF CONFORMITY

Name of Manufacturer:

Vibrosonic GmbH

Address of Manufacturer:

Franz-Volhard-Straße 3

68167 Mannheim

Germany

Single registration number (SRN):

DE-MF-000022006

Product	Reference	Basic UDI-DI	Class	Classification Rule(s)
Vibrosonic harmonize	VFTS002	426070005VFTS0029X	lla	chapter II, section 3.3

Intended purpose:

The fitting software Vibrosonic harmonize is used to configure the hearing aids distributed by Vibrosonic and to adapt them to customer-specific hearing loss in terms

of their acoustic amplification properties.

The undersigned hereby declares that the medical devices (according to article 2 (1)) specified above meet the provisions of the Medical Device Regulation (EU) 2017/745, Annex I. This declaration of conformity is issued under the sole responsibility of the manufacturer.

The conformity assessment procedure used is "CONFORMITY ASSESSMENT BASED ON A QUALITY MANAGEMENT SYSTEM AND ON ASSESSMENT OF TECHNICAL DOCUMENTATION" according to the Medical Device Regulation (EU) 2017/745, Annex IX.

The product is classified in class IIa according to MDR Annex VIII and subject to the quality management system of Vibrosonic GmbH.

Certificate Registration No.

D1445600006

Date:

2023-07-24

Issued by:

mdc medical device certification GmbH

Kriegerstraße 6, 70191 Stuttgart, GERMANY

NotifiedBodyNo.:

0483

Expiring date of the CE Declaration of Conformity: 2028-07-23

The conformity is declared according to the Regulation (EU) 2017/745, Annex IV and is valid from date of signature.

Mannheim, 2023-08-31

Dr. Jonathan Schächtele

Person responsible for regulatory compliance (PRRC)

Dok-ID: P03Z05; rev. 001