

## Declaration of Conformity According to Medical Device Directive (MDD) 93/42/EEC

## Manufacturer

Auditdata A/S
Dalbergstrøget 5-7
2630 Taastrup
Denmark

## **Conformity Assessment Procedure**

Annex II.3 excluding (4) of the Medical Device Directive

MDD 93/42/EEC

**Notified Body** 

TÜV SÜD Product Service GmbH

Ridlerstr. 65 80339 München CE

**Product Identification** 

Category: Audiometers and Associated

Accessories

Brand: OTOPod Model: M1, M2

Lot/Batches/Serial number: All issued serial numbers from

M1: 28000001 M2: 29000001

**MDD Directive** 

Class IIa, Rule 10, MDD

93/42/EEC

The medical device compliance with the essential requirements in accordance with Annex I of the Medical Device Directive 93/42/EEC.

We declare under our sole responsibility that the products, to which this declaration relates, are in conformity with the Essential Requirements Annex I of the above directive.

This DOC is Valid until May 26<sup>th</sup> 2024 - EC certificate validity date.

. Taastrup, December 2<sup>nd</sup> 2019

Dan Haugbøl, Director QA/RA & IT Information security

Signature