## **Auditdata**

## DECLARATION OF CONFORMITY

## REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT

Manufacturer name and

address

Auditdata A/S Wildersgade 10B 1408 Copenhagen

Denmark

**Notified Body name and** 

address

TÜV SÜD Product Service

GmbH

Ridlerstrasse 65 80339 München  $\epsilon$ 

0123

**Product Identification** MD Category: Hearing Medical Diagnostic (Hardware

and Software)

Trademark: Primus PFU+

Type/Model: Primus Fitting Unit - PFU+ (PFU+)
CS (Common specification) N/A no common specification has been

published

SRN: DK-MF-000011415
Basic UDI/DI: 05711781DHF2000ZC

Risk class: Ila, rule 10

Lot/Batches/Serial All issued serial numbers from

number: 21000001

**Intended purpose** Audiometer is a device used for evaluating hearing acuity. The

audiometer records the subject's responses to produce an audiogram of threshold sensitivity, or speech understanding profile. Audiometer with stated accessories is indicated for non-continuous, noninvasive air and optionally bone conduction and speech audiometric testing. Audiometer is indicated for non-continuous real-ear measurements (REM) at the ear drum by means of noninvasive external ear canal

insertion of a probe tube.

**Conformity assessment** Annex IX (Quality system and technical documentation assessment)

**EC-Certificate No.:** G10 076081 0015

**DOC valid until** 2029-02-18

This declaration of conformity is issued under the sole responsibility of Auditdata A/S. We hereby declare that the medical device specified above Is in conformity with the European Regulation (EU) 2017/745 and Directive 2011/65/EU.

## Copenhagen, February 19th 2024

Denys Lebedev, QA/RA Manager