## PLANMECA

## **DECLARATION OF CONFORMITY**

We

Planmeca Oy, Asentajankatu 6, 00880 Helsinki Finland

declare under our sole responsibility that the product

## Planmeca ProSensor HD

to which this declaration relates is in conformity with following standards or other normative documents:

IEC 60601-1 + A1:2012	Medical electrical equipment – Part 1: General requirements for basic safety and essential performance
IEC 60601-1-2:2014	Medical electrical equipment. Part 1: General requirements for safety. 2. Collateral Standard: Electromagnetic compatibility - Requirements and tests.
IEC 60601-1-6 + A1:2013	Medical electrical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability.
IEC 62366 + A1:2014	Medical devices – Part 1: Application of usability engineering to medical devices.

following the provisions of **Council Directive 93/42/EEC** as set out in **Annex II**. Planmeca ProSensor HD is Class IIa device.

EC certificate: FI15/07006 The Notified Body is SGS Fimko Ltd. no 0598.

Helsinki, 2021-05-25

Niina Vuorikallas Director, Quality & Regulatory Affairs