

Declaration No.:	2031	
Deciaration No.:	201	

## EC/MDD DECLARATION OF CONFORMITY

This is a declaration made in accordance with the requirements of Council Directive 93/42/EEC of 14 June 1993 concerning medical devices (MDD 93/42/EEC).



Manufacturer's Name:

**Business Address:** 

NIHON KOHDEN CORPORATION

1-31-4 Nishiochiai, Shinjuku-ku

Tokyo 161-8560, Japan

**European Representative:** 

NIHON KOHDEN EUROPE GmbH

St. September 1984 in Landon

Raiffeisenstrasse 10, D - 61191 Rosbach, Germany

**Product Name and Model Name:** 

Finger Probe

TL-201T

Classification:

Address:

IIa

Each kind of medical device to which the Full Quality Assurance Procedures (Annex II) have been applied complies with the applicable provisions of the essential requirements, the classification rules, at each stage, from the design of the device until its final inspection before being supplied.

**Notified Body:** 

BSI Group The Netherlands B.V.

**EC** Certificate:

CE 01342

Standard Applied:

EN ISO 13485: 2016 EN ISO 14971: 2012 IEC 60601-1: 2005

IEC 60601-1 Amendment 1: 2012

IEC 60601-1-6: 2010

IEC 60601-1-6 Amendment 1:2013

IEC 62366: 2007

IEC 62366 Amendment 1: 2014

ISO 9919: 2005 ISO 10993-1: 2009 ISO 80601-2-61:2017

EN 1041: 2008

EN 1041 Amendment 1: 2013 EN ISO 15223-1: 2016

Note:

The underlined standard above is updated in this Declaration of Conformity.

Although the Declaration is issued after 26 May 2021, the Date of Application of MDR, the Declaration is made for the changes which do not correspond to significant changes specified in the MDCG 2020-3, and maintains compliance with the old standards listed on the Declaration dated 15 March 2019.

Consequently, this Declaration does not deviate from the MDR Article 120, and does not invalidate the Declaration of 15 March 2019.

**Authorized Signatory:** 

Tokyo, Japan / 18 February 2022

Place and date of issue

Hiroko Hagiwara General Manager

Clinical Development & Regulatory Affairs Division