

EU DECLARATION OF CONFORMITY

This is a declaration made in accordance with the requirements of the following relevant Union harmonisation legislation. The manufacturer assures that the device that is covered by the present declaration is in conformity with this Regulation (EU) 2017/745 for Medical Devices and, if applicable, with any other relevant Union legislation that provides for the issuing of an EU declaration of conformity. The declaration of conformity is issued under the sole responsibility of the manufacturer.



Manufacturer's Name: NIHON KOHDEN CORPORATION

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Tokyo 161-8560, Japan

SRN: JP-MF-000019022

European

Representative: NIHON KOHDEN EUROPE GmbH

Address: Raiffeisenstrasse 10, 61191 Rosbach, Germany

SRN: DE-AR-000010740

Regulation (EU) 2017/745(MDR)

Classification/Risk Class: I

**Conformity assessment
procedure:**

Annex II and III

Directive 2011/65/EU and 2015/863/EU

Standard Applied: EN IEC 63000: 2018

Directive 2014/53/EU (RED)

Notified Body

Name and No. :

EU-Type Examination

Certificate No. :

Standard Applied:

Product Name, Model Number and Basic UDI-DI :

Product Name	Model Number	Basic UDI-DI	MDR	RoHS	RED
Disposable Pads	P-711	4931921P-711M7	×	×	—
Disposable Pads	P-713	4931921P-713MB	×	×	—
Disposable Pads	P-721	4931921P-721MA	×	×	—
Disposable Pads	P-730K	4931921P-730KDT	×	×	—
Disposable Pads	P-740K	4931921P-740KDY	×	×	—

Intended purpose: The products listed above are accessories for Defibrillator.

Additional Information: NA

Authorized Signatory:
Tokyo, Japan/ 31 August 2022
Place and date of issue


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