

Declaration No.:	2250	

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.



	2.45	200	5.00	Programme Company	
V	lanı	ıfactu	rer's	Name.	

NIHON KOHDEN CORPORATION

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JP-MF-000019022

European

Representative:

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Address:

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SRN:

DE-AR-000010740

☑ Regulation (EU) 2017/745(MDR)

Classification/Risk Class:

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Conformity assessment

procedure:

Annex II and III

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

Standard Applied:

☐ Directive 2014/53/EU (RED)

Notified Body

Name and No.:

EU-Type Examination

Certificate No.:

Standard Applied:



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Product Name, Model Number and Basic UDI-DI:

Product Name	Model Number	Basic UDI-DI	MDR	RoHS	RED
Disposable Cuff Infant	YP-840T	4931921YP-	×	_	
Disposable Cult Illiant		840TRK	^		
Disposable Cuff Child	YP-841T	4931921YP-	×	_	•
Disposable Cult Cliffd		841TRN			
Disposable Cuff Small Adult	YP-842T	4931921YP-	×	-	
Disposable Cult Shan Addit		842TRR			_
Disposable Cuff Adult	YP-843T	4931921YP-	×	_	-
Disposable Cult Adult		843TRU			
Disposable Cuff Large Adult	was Adult VD 944T	4931921YP-	×	_	
Disposable Cull Large Adult	YP-844T	844TRX			_
Disposable Cuff Thigh	YP-845T	4931921YP-	×	_	,
Disposable Cult Tiligli		845TS2			_

Intended purpose:

The products listed above are used to measure non-invasive blood pressure.

Additional Information:

NA

Authorized Signatory:

Tokyo, Japan/ 31 August 2022

Place and date of issue

Hiroko Hagiwara General Manager

Clinical Development & Regulatory Affairs Division