

EU DECLARATION OF CONFORMITY

Division/Group: RAQA

Revision: 1

IH-500**REF** 001500

BUDI-DI : 361052A002437X



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Real Manufacturing Site

Bio-Rad Singapore
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#05-01 Techview
415941 Republic of Singapore

We, as the manufacturer of the device(s) take sole responsibility for and hereby declare that the above mentioned product(s) meet(s) the provisions of the following Regulation(s) / Directive(s):

- Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS)
- Regulation EU 2017/746 on *in vitro* Diagnostic medical devices

Risk CLASS:

A B C D

CONFORMITY ROUTE

ANNEX I & II+III

Common Specification (CS) : Not Applicable

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
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Date of the first issuance of the EU Declaration of Conformity: 04-05-2022

Place, Date:	Cressier, 03.10.2023
Signed by:	Jérémy Poropane
Function:	Associate Director, Regulatory Affairs, Bio-Rad CDG
Signature:	

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