

Declaration of Conformity Certificate

We

AHN Biotechnologie GmbH	
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Germany	
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Declare with sole responsibility, that our product/s:

EDMA Code	EDMA Description	Internal Product Name	Classification Rationale per IVDD
21-09	Pipette tips	Expell, ExpellPlus, myTip Pipette Tips, Sterile Tips, Non-Sterile Tips, Low Retention Tips and Filter Tips	Other IVD, Annexe III

meet, the essential requirements of Council Directive 98/79/EC pertaining to in vitro diagnostics. Pathway of conformity per Annex III.

Notified Body: -

The product(s) identified above meet requirements of the IVDD by meeting the following standards

Standard No. Council Directive 93/42/EEC as amended by 2007/47/EC or 98/79/EC

We hereby appoint mdi Europa GmbH, Langenhagener Str. 71, 30855 Langenhagen, Germany to act as European Authorized Representative as explicitly defined in Article 1, § 2(g) of Directive 98/79/EEC.

Signed this day: 05-03-2018, by Magdalena Babut-Carstensen, Compliance Manager

Expiry Date: -

mdi Europa use only!

mark on the above mentioned products and The necessary pre-requisites for placing the

Signed this day: 05-03-2018

mareuropa

THE MEDICAL DEVICE SERVICE—MANAGEMENT 7-0
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Commercial AG 6/5 Commerzbank AG 6/\$
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