

To our customers

Michael Eberhard

+49 7461 705 1318

Michael.Eberhard@hettichlab.com

www.hettichlab.com

Dear Customer

Tuttlingen, 22.07.2021

RoHS 2011/65 / EU conformity and compliance with the Delegated Directive (EU) 2015/863

We hereby inform you that Andreas Hettich GmbH & Co. KG manufactures and delivers the products of the product groups

- Medical devices according to Directive 93/42/EEC, as well as their accessories and spare parts from July 22nd, 2014
- In-Vitro diagnostic devices according to Directive 98/79/EC, as well as their accessories and spare parts from July 22nd, 2016

conformal to EU Directive 2011/65/EU of the European Parliament and the Council of June 8th, 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS II) as well as to the Delegated Directive (EU) 2015/863 of the Commission from March 31, 2015 amending Annex II of Directive 2011/65/EU.

The EC declarations of conformity for these products contain the 2011/65/EC and (EU) 2015/863 as applicable directives. They are part of the technical documentation of every product supplied.

Best regards,
Andreas Hettich GmbH & Co. KG



i.A. Michael Eberhard
Head of Regulatory & Quality Affairs