

## Certificate of Verification of Manufactured Products Directive 98/79/EC Annex IV, Section 6 In Vitro Diagnostic Medical Devices

Registration No.: LL 60030810 0032

Report No.: 21110698 051

## Manufacturer:

Bio-Rad Medical Diagnostics GmbH Industriestr. 1 63303 Dreieich Deutschland

## Product Identification:

Erytype® S ABD Confirmation Lot No.: 3247050 Article No.: 806110

The Notified Body hereby declares that a verification relating to the listed products has been performed according to Annex IV, section 6 of the directive 98/79/EC and that the devices conform to the requirements of the abovementioned directive.

Notified Body

TÜVRheinland IGA Products

Date: 2022-11-29

TÜV, TUEV and TUV are registered trademarks. Utilisation and application requires prior appro

Katja Mierisch

## **TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg**

TÜV Rheinland LGA Products GmbH is a Notified Body according to Directive 98/79/EC concerning in vitro diagnostic medical devices with the identification number 0197.