

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

Registration No.: LL 60033608 0027

Report No.: 21111179 048

Manufacturer:

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

Product Identification:

Date: 2022-11-24

Erytype® S Rh-Pheno double Lot No.: 3244010 Article No.: 806150

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

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

Katja Mierisch

LGA

UVRheint

erungs

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.