

EC Design-Examination Certificate Directive 98/79/EC Annex IV, Section 4 In Vitro Diagnostic Medical Devices

Registration No.: IL 60139592 0001

Report No.: 60239191 001

Manufacturer:

IMMUCOR

Medizinische Diagnostik GmbH Robert-Bosch-Strasse 32

63303 Dreieich Deutschland

Product

Identification:

In vitro diagnostic reagents, including control material,

TÜVRheinland

Wisiorungsste

for determining blood groups: Kell (K)

(see attachments for products included)

The Notified Body hereby declares that an examination of the design dossier relating to the listed products has been performed according to Annex IV, section 4 of the directive 98/79/EC and that the design of the devices conforms to the requirements of the abovementioned directive.

Expiry Date: 2023-12-07

Effective Date: 2019-05-29

Date: 2019-05-28

Notified Body

∕Dr. H. Lüdemann

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.



TÜV Rheinland LGA Products GmbH Tillystraße 2, 90431 Nürnberg

Doc. 1/1, Rev. 0

Attachment to Certificate

Registration No.: Report No.: IL 60139592 0001 60239191 001

Manufacturer:

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Medizinische Diagnostik GmbH

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Products:

- immuClone (1) Anti-K (Kell) IgM and Galileo
- immuClone (2) Anti-K (Kell) IgM and Galileo
- Automated immuClone Anti-K (Kell) Galileo IgM
- Anti-K (Kell) quick

Date: 2019-05-29

10/020 h 04.08 @ TÜT, TUEN and TUY are registered trademarks. Utilisation and application requires prior approval.

TÜVRheinland Lüdemann

H. Lüdemann