EU Certificate

Technical Documentation Assessment REGULATION (EU) 2017/746 on in Vitro Diagnostic Medical Devices, Annex IX



Registration No.:

IX 1191616-3

Manufacturer:

IMMUCOR

hanulacturer, ilvilviocor

Medizinische Diagnostik GmbH

Robert-Bosch-Strasse 32

63303 Dreieich Germany

EUDAMED Single

Registration No.:

DE-MF-000006494

General product group

name:

Products class D

HAEMATOLOGY / HAEMOSTASIS / IMMUNOHAEMATOLOGY / HISTOLOGY /

CYTOLOGY

IVR 0102. Devices intended to determine markers of the Rhesus system [RH1 (D),

RHW1, RH2 (C), RH3 (E), RH4 (c), RH5 (e)] W0103030202 - RHESUS PHENOTYPES

Product name: Models and types: immuClone (1) Anti-C IgM - REF: 0007206, 0007204, 0007216, 0007214

immuClone (1) Anti-C Galileo IgM - REF: 0066011

Basic UDI-DI:

88823405W0103030202D22

The Notified Body hereby declares that the requirements of Annex IX, Chapter II, Section 4 and 5.1 / 5.2 of the REGULATION (EU) 2017/746 have been met for the listed products. The above named manufacturer has established and maintains a technical documentation defined by Annex IX, Chapter II, Section 4 and 5.1 / 5.2 of the aforementioned regulation. In addition to this certificate an EU quality management system certificate and for class D devices a batch verification is required before placing the listed products on the market.

Report No.:

1120777-20

Effective date:

2023-09-28

Expiry date:

2028-09-27

Issue date:

2023-09-28





Katja Mierisch TÜV Rheinland LGA Products GmbH Tillystraße 2 · 90431 Nürnberg · Germany

TÜV Rheinland LGA Products GmbH is a Notified Body according to REGULATION (EU) 2017/746 concerning in vitro diagnostic medical devices with the identification number 0197.

EU Certificate

Technical Documentation Assessment REGULATION (EU) 2017/746 on in Vitro Diagnostic Medical Devices, Annex IX



Registration No.:

IX 1191616-3

Manufacturer:

IMMUCOR

Medizinische Diagnostik GmbH

Robert-Bosch-Strasse 32

63303 Dreieich Germany

Intended use:

immuClone® (1) Anti-C IgM and immuClone® (1) Anti-C Galileo IgM are Blood Group Reagents used to detect the C erythrocyte antigen from donors and recipients by direct hemagglutination test for the purpose of a blood transfusion to ensure the safety and compatibility between the patient and the blood component selected for transfusion. For Manual Tube, Slide, Microplate and Automated Microplate Tests (qualitative), immuClone® (1) Anti-C IgM is intended for Manual Tube. Slide and Microplate Tests (qualitative). immuClone® (1) Anti-C Galileo IgM

is intended for Automated Microplate Tests (qualitative).

Authorised

representative(s):

N/A

Certificate history		
Revision:	Description:	Issue date:
0	Initial certification	2023-09-28

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