

EU Certificate

Technical Documentation Assessment REGULATION (EU) 2017/746 on In Vitro Diagnostic Medical Devices Annex IX Chapter II

Registration No.: IX 1804147-24

Manufacturer: Immucor, Inc.
3130 Gateway Drive
Norcross GA 30071
USA

EUDAMED Single
Registration No.: US-MF-000011568

Classification: D

General product group name: 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)]
IVR 0103 Devices intended to determine markers of the Kell
system [Kel1 (K)]
IVR 0104 Devices intended to determine markers of the Kidd
system [JK1 (Jka), JK2 (Jkb)]
IVR 0105 Devices intended to determine markers of the Duffy
system [FY1 (Fya), FY2 (Fyb)]
IVR 0106 Other devices intended to be used for blood
grouping
W01030303 - Antibody Detection (Immunohaematology)

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.: 1194556-20

Effective date: 2026-07-28

Expiry date: 2031-07-27

Issue date: 2026-07-28



Dr. Volker Schlueter
TÜV Rheinland LGA Products GmbH
Tillystraße 2 · 90431 Nürnberg · Germany

This certificate can be validated on <https://www.certipedia.com>

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.

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Precisely Right.

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Product name: Capture-R® Ready Indicator Red Cells

Models and types: 0006427, 0006428

Basic UDI-DI: 08882340006KDP

Intended use: Capture-R® Ready Indicator Red Cells is intended for use with Capture-R® Ready-Screen®, Capture-R® Ready-ID® or Capture-R® Select for the qualitative detection of unexpected IgG antibodies to red blood cells by manual, semiautomated or automated solid phase red blood cell adherence methods for the purpose of blood transfusion to determine safety and compatibility with potential. For laboratory professional use in testing blood specimens from patients and donors.

Authorized representative(s): Immucor Medizinische Diagnostik GmbH
Robert-Bosch-Straße 32, D-63303 Dreieich
SRN: DE-AR-000007083

Certificate history		
Revision:	Description:	Issue date:
0	Initial certification	2026-07-28

