

EU Certificate

Quality Management System

REGULATION (EU) 2017/746 on In Vitro Diagnostic Medical Devices

Annex IX Chapters I and III

Registration No.: HX 1483000-1

Manufacturer:
EUROIMMUN
Medizinische Labordiagnostika AG
Seekamp 31
23560 Lübeck
Germany

EUDAMED Single
Registration No.: DE-MF-000005296

Products:
Products of class B:

IMMUNOCHEMISTRY (IMMUNOLOGY)

IVR 0602: Devices intended to be used for screening, determination or monitoring of physiological markers for a specific disease

W01021001 - AUTOIMMUNE CONNECTIVE TISSUE
DISEASES

W01021002 - NEURO-AUTO-IMMUNE DISEASE

W01021004 - VASCULITIS

W01021007 - AUTOIMMUNE HEPATITIS

The Notified Body hereby declares that the requirements of Annex IX, Chapter I of the REGULATION (EU) 2017/746 have been met for the listed products. The above named manufacturer has established and applies a quality management system, which is subject to periodic surveillance, defined by Annex IX, Chapter I, Section 3 of the aforementioned regulation. The requirements of Annex IX, Chapter III are fulfilled.

If class D devices are covered by this certificate an EU technical documentation assessment certificate according to Chapter II, Section 4 is required before placing them on the market.

If class B, C or D devices for self-testing or near-patient testing are covered by this certificate an EU technical documentation assessment certificate according to Chapter II, Section 5.1 is required before placing them on the market.

If companion diagnostics are covered by this certificate an EU technical documentation assessment certificate according to Chapter II, Section 5.2 is required before placing them on the market.

Report No.: 1202331-20

Effective date: 2026-01-21

Expiry date: 2028-05-09

Issue date: 2026-01-21

Katja Mierisch

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 medical devices with the identification number 0197.



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Zentrale der Länder
für Gesundheitsschutz
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Products:
W01021090 - VARIOUS AUTO-IMMUNE DISEASE
W01021112 - ANTI-CYCLIC CITRULLINATED PEPTIDE
W01021199 - RHEUMATOID / INFLAMMATORY DISEASE
MARKERS – OTHER
W01021520 - CONTROLS – IMMUNOCHEMISTRY

IVR 0603: Devices intended to be used for screening,
confirmation/determination, or monitoring of allergies and
intolerances
W01020201 - IMMUNOGLOBULIN E – TOTAL
W01020204 - IMMUNOGLOBULIN E –
MONOTEST/PLURIRESULT-MULTI AG

IVR 0608: Devices intended to be used for screening,
determination or monitoring of physiological markers
W01020190 - OTHER SPECIFIC PROTEINS
W01020702 - VITAMINES
W01021520 - CONTROLS – IMMUNOCHEMISTRY
W01029099 - IMMUNOCHEMISTRY REAGENTS - OTHER

INFECTIOUS DISEASES

IVR 0503: Devices intended to be used to detect the presence
of, or exposure to an infectious agent including sexually
transmitted agents
W01050101 - CHLAMYDIA
W01050106 - LYME BORRELIOSIS
W01050117 - OTHER BACTERIOLOGY IMMUNOASSAYS
W01050404 - EPSTEIN BARR VIRUS
W01050405 - OTHER VIROLOGY - NA REAGENTS



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Products:
W01050406 - OTHER VIROLOGY ANTIGEN/ANTIBODY
DETECTION
W01050407 - MEASLES VIRUS
W01050408 - MUMPS / PAROTITIS VIRUS
W01050502 - MISCELLANEOUS PARASITOLOGY
W01050808 - CONTROLS - INFECT. IMMUNOLOGY

IVR 0504: Devices intended to be used to determine the
infectious load, to determine infective disease status or
immune status and devices used for infectious disease staging
W01050117 - OTHER BACTERIOLOGY IMMUNOASSAYS

CHEMISTRY / IMMUNOCHEMISTRY INSTRUMENTS
IVR 0503: Devices intended to be used to detect the presence
of, or exposure to an infectious agent including sexually
transmitted agents
W0201020192 - AUTOMATED IMMUNOCHEMISTRY
ANALYSERS – IVD MEDICAL DEVICE SOFTWARE

Products of class C:
IMMUNOCHEMISTRY (IMMUNOLOGY)
IVR 0602: Devices intended to be used for screening,
determination or monitoring of physiological markers for a
specific disease
W01021004 - VASCULITIS
W01021090 - VARIOUS AUTO-IMMUNE DISEASE



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Products:

INFECTIOUS DISEASES

IVR 0501: Devices intended to be used for pre-natal screening of women in order to determine their immune status towards transmissible agents
W01050401 - RUBELLA VIRUS
W01050402 - CYTOMEGALOVIRUS
W01050501 - TOXOPLASMA

IVR 0503: Devices intended to be used to detect the presence of, or exposure to an infectious agent including sexually transmitted agents

W01050101 - CHLAMYDIA
W01050103 - SYPHILIS (TREPONEMA PALLIDUM)
W01050107 - MYCOBACTERIA GENUS + SPECIES
W01050405 - OTHER VIROLOGY - NA REAGENTS
W01050406 - OTHER VIROLOGY ANTIGEN/ANTIBODY DETECTION
W01050409 - VARICELLA ZOSTER VIRUS
W01050501 - TOXOPLASMA
W01050604 - MYCOLOGY - NA REAGENTS

GENETIC TESTING

IVR 0402: Devices intended to be used to predict genetic disease/disorder risk and prognosis
W01060101 - MONOGENETIC DISORDERS
W01060104 - POLYMORPHISMS



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Products:

NUCLEIC ACID TESTING INSTRUMENTS
IVR 0402: Devices intended to be used to predict genetic
disease/disorder risk and prognosis
W02050292 - MICRO-ARRAY INSTRUMENTS – IVD
MEDICAL DEVICE SOFTWARE

IVR 0501: Devices intended to be used for prenatal screening
of women in order to determine their immune status towards
transmissible agents
W02050292 - MICRO-ARRAY INSTRUMENTS – IVD
MEDICAL DEVICE SOFTWARE

Authorized representative(s): N/A



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Certificate history		
Revision:	Description:	Issue date:
0	Initial certification	2023-05-10
1	Scope extension, EUROI_PDQ2_HX_2023-07-12_2_20230822_extsigned.pdf	2023-08-22
2	Scope extension: Products of class B (W01050405, W01021199, W01020204, W01020190, W01021520) Scope reduction: Products of class B (W01020299) and class C (W01050403, W01050705) EUROI_PDQ2_HX_2023-12-15_2024-03-26_extsigned.pdf	2024-03-26
3	Scope extension: Products of class B (W01050117, W01021001), Products of class C (W01021001), EUROI_PDQ2_HX_2024-03-06_extsigned.pdf	2024-05-17
4	Scope extension: Product of class D (W01050406) and class C and B EUROI_PDQ2_HX_2024-08-07	2024-08-19
5	Scope extension: Products of class B (W01050407, W01029099), EUROI_PDQ2_HX_2024-12-18.pdf	2024-12-18
6	Scope extension: Products of class C (W01050501) Scope reduction: Products of class B (W01050106) EUROI_PDQ2_HXIX_2025-03-26.pdf	2025-03-31
7	Scope extension: Products of class B (W01050106, W01021002) and class C (W01050101, W01050401) Scope reduction: Products of class D (W01050406)	2025-06-13
8	Scope extension: Products of class B (W01020190) and class C (W01050103, W01050409)	2025-08-20
9	Scope extension: Products of class B (W01021090, W01021112, W01021001, W01050404, W01050101, W01050408) and class C (W01050406, W01050101, W01050501, W01050604) Removal of class C device: Anti-Toxoplasma gondii IIFT (IgM) EUROI_PDQ2_HX_2025-10-23_extsigned-signed.pdf	2026-01-21



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