

EC Certificate

mdc medical device certification GmbH

Notified Body 0483
herewith certifies that

**ravo Diagnostika GmbH
Oltmannsstraße 5
79100 Freiburg
Germany**

for the scope

**in vitro diagnostic devices for determination of
antibodies specific for toxoplasmosis
ToxoTool M-I ISAGA
ToxoTool A-I ISAGA**

has introduced and applies a

Quality System

for the design, manufacture and final inspection.

The mdc audit has proven that this quality system
meets all requirements according to

**Annex IV – excluding Section 4 and 6
of the Council Directive 98/79/EC**

of the European Parliament and of the Council of
27 October 1998 on in vitro diagnostic medical devices.

The surveillance will be held as specified in Annex IV, Section 5.

Valid from	2022-05-05
Valid until	2025-05-26
Registration no.	D1052800015
Report no.	P22-00227-228273
Stuttgart	2022-05-05


Head of Certification Body



Benannt durch/Designated by
Zentralstelle der Länder
für Gesundheitsschutz
bei Arzneimitteln und
Medizinprodukten
www.zlg.de
ZLG-BS-247.10.05