

## **EC DECLARATION OF CONFORMITY**

DIESSE DIAGNOSTICA SENESE SPA MANUFACTURER: STRADA DEI LAGHI 39 53035 MONTERIGGIONI (SI) **ITALY EUROPEAN REPRESENTATIVE:** // PRODUCT: **ENZY-WELL TOXOPLASMA IgA** CODE: 91043 **CLASSIFICATION:** IVD IN ANNEX II B ANNEX APPLIED N° IV EXCLUDING (4, 6) CONFORMITY ASSESSMENT ROUTE: WE HEREWITH DECLARE THAT THE ABOVE MENTIONED PRODUCTS MEET THE PREVISIONS OF THE COUNCIL DIRECTIVE 98/79EC FOR IN VITRO DIAGNOSTIC MEDICAL DEVICE. ALL SUPPORTING DOCUMENTATION IS RETAINED UNDER THE PREMISES OF THE MANUFACTURER. TÜV SÜD PRODUCT SERVICE GMBH NOTIFIED BODY: ZERTIFIZIERSTELLE RIDLERST. 65 - 80339 MÜNCHEN GERMANY No. 0123 (EC) CERTIFICATE: VI 056726 0002 Rev. 00 START OF CE-MARKING: **DECEMBER 2003** REVISION: 14 PLACE, DATE OF ISSUE: MONTERIGGIONI, 11 MAY 2022 EXPIRY DATE: 26 MAY 2025 THE PRESENT EC DECLARATION OF CONFORMITY IS ISSUED UNDER THE SOLE AND EXCLUSIVE RESPONSIBILITY OF THE MANUFACTURER.

> CHIARA MUZZI REGULATORY AFFAIRS MANAGER

THIS IS THE CERTIFIED COPY OF THE ORIGINAL DOCUMENT STORED IN ARCHIVE OF DIESSE DIAGNOSTICA SENESE SPA

ISSUED: MONTERIGGIONI, 11/05/2022

MAGDALENA STOCZKO REGULATORY SUPERVISOR

M Stoules

SIGNATURE: