







EC Certificate

Full Quality Assurance System Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6) (List A and B and devices for self-testing)

No. V1 056726 0002 Rev. 00

Manufacturer:

Diesse Diagnostica Senese SpA

Strada dei Laghi 39 53035 Monteriggioni (SI) ITALY

Product Category(ies): Reagents for determination of infection markers and detection of tumoral marker PSA

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device families in accordance with IVDD Annex IV. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of List A devices an additional Annex IV (4) certificate is mandatory. All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:V1056726 0002 Rev. 00

Report no.:	
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Valid from: Valid until: 2022-04-29 2025-05-26

Date,

2022-04-29

Christoph Dicks Head of Certification/Notified Body



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Model(s):

ELISA reagents, kits, and controls for Rubella, Toxoplasma, CMV, and Chlamydia of: CHORUS product line, ENZYWELL product line Complement Fixation Test reagents, kits, and controls for Rubella, Toxoplasma, CMV, and Chlamydia of: Manual procedure product line, SERAMAT product line ELISA reagents, kits, and controls for detection of tumoral marker PSA of CHORUS product line

Facility(ies):

Diesse Diagnostica Senese SpA Via del Pozzo 5, 53035 Monteriggioni (SI), ITALY

Diesse Diagnostica Senese SpA Strada dei Laghi 39, 53035 Monteriggioni (SI), ITALY

Diesse Diagnostica Senese SpA Via delle Rose 10, 53035 Monteriggioni (SI), ITALY