

DIESSE EU DECLARATION OF CONFORMITY

MANUFACTURER:	DIESSE DIAGNOSTICA SENESE SPA STRADA DEI LAGHI 39 53035 MONTERIGGIONI (SI), ITALY	
SINGLE REGISTRATION NUMBER	IT-MF-000013311	
AUTHORIZED REPRESENTATIVE:	//	
PRODUCT: CODE:	VACU-TEC "S" 10200/S	
INTENDED PURPOSE:	Single-use vacuum tube containing anticoagulant (sodium citrate) to be used for manual sampling and whole blood collection for the determination of Erythrocyte Sedimentation Rate (ESR) in DIESSE VES instruments. All the tubes must be used by professional laboratory users only.	
BASIC UDI-DI	803389132 ESRTUBES00 JL	
UDI-DI	08033891320539	
RISK CLASS:	CLASS A	
CLASSIFICATION RULE:	RULE 5C	
CONFORMITY ASSESSMENT ROUTE:	ARTICLE 17, ANNEX II and ANNEX III	
WE HEREWITH DECLARE THAT THE ABOVE MENTIONED PRODUCT IS IN CONFORMITY WITH REGULATION		

(EU) 2017/746 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL ON IN VITRO DIAGNOSTIC MEDICAL DEVICE. ALL SUPPORTING DOCUMENTATION IS RETAINED UNDER THE PREMISES OF THE MANUFACTURER.

REFERENCE TO ANY CS APPLIED:	NOT APPLICABLE
NOTIFIED BODY:	NOT NECESSARY
(EU) CERTIFICATE:	//
REVISION:	0
PLACE, DATE OF ISSUE:	MONTERIGGIONI, 23 MAY 2022
EXPIRY DATE:	//

THE PRESENT EU DECLARATION OF CONFORMITY IS ISSUED UNDER THE SOLE AND EXCLUSIVE RESPONSIBILITY OF THE MANUFACTURER.

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MARIA CLAUDIA ALCARO PERSON RESPONSIBLE FOR THE REGULATORY COMPLIANCE

SIGNATURE:

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

ISSUED: MONTERIGGIONI, 2022-05-23

M. Stonks

MAGDALENA STOCZKO REGULATORY SUPERVISOR