

SIGNATURE:

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: | VACU-TEC FLAG |
| CODE: | 10235 |
| 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. |
| DAGIC LIDI DI | 007700173 FCDTUDECO0 14 |
| BASIC UDI-DI | 803389132 ESRTUBES00 JL |
| UDI-DI | 08033891320751 |
| 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 | |
| • • | ENT AND OF THE COUNCIL ON IN VITRO DIAGNOSTIC MEDICAL ATION 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. | |
| | Varia Centra Alan |

MARIA CLAUDIA ALCARO
PERSON RESPONSIBLE FOR THE REGULATORY
COMPLIANCE

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

ISSUED: MONTERIGGIONI, 2022-05-23

M. Stoales

MAGDALENA STOCZKO REGULATORY SUPERVISOR

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