|                              | DECLARATION OF CONFORMITY               |
|------------------------------|---|
| MANUFACTURER:                | DIESSE DIAGNOSTICA SENESE SPA           |
|                              | STRADA DEI LAGHI 39                     |
|                              | 53035 MONTERIGGIONI (SI),               |
|                              | ITALY                                   |
| EUROPEAN REPRESENTATIVE:     | //                                      |
| GENERIC NAME                 | AUTOMATIC INSTRUMENT FOR ESR ANALYSIS   |
| PRODUCT:                     | CUBE 30 touch                           |
| CODE:                        | 10395                                   |
|                              |   |
| TECHNICAL DATA:              | 110-230 Vac (50-60 Hz); Pwr: 100 VA     |
| CLASSIFICATION:              | IVD NOT IN ANNEX II OR SELE-TESTING IVD |
| CLASSIFICATION.              | TVD NOT IN ANNEX II OR SELF-TESTING IVD |
| CONFORMITY ASSESSMENT ROUTE: | ANNEX APPLIED N° III EXCLUDING (6)      |
|                              | ESSENTIAL REQUIREMENTS OF ANNEX I       |

WE HEREWITH DECLARE THAT THE DESIGN, TYPE OF MANUFACTURE OF THE IN VITRO MEDICAL DIAGNOSTIC DEVICE DESCRIBED ABOVE AND THE VERSION DISTRIBUTED ON THE MARKET, CONFORMS TO THE 98/79/EEC DIRECTIVE RELEVANT TO THE IN VITRO MEDICAL-DIAGNOSTICS DEVICES (IVD)

## THIS CERTIFICATE WILL LOSE ITS VALIDITY IN THE EVENT OF:

- MODIFICATIONS MADE TO THE MACHINE IN QUESTION WITHOUT OUR AUTHORIZATION
- INCORRECT USE OF THE INSTRUMENT
- TECHNICAL INTERVENTIONS PERFORMED BY UNAUTHORIZED PERSONNEL
- INSTALLATION OF NON-ORIGINAL SPARE PARTS.

## THE PRODUCT CONFORMS, AS A WHOLE AND IN ITS PARTS, WITH THE FOLLOWING STANDARDS AND THEIR AMENDMENTS:

- EN 61010-1:2010 SAFETY REQUIREMENTS FOR ELECTRICAL EQUIPMENT FOR MEASUREMENT, CONTROL, AND LABORATORY USE - PART 1: GENERAL REQUIREMENTS
- EN 61010-2-081:2015 SAFETY REQUIREMENTS FOR ELECTRICAL EQUIPMENT FOR MEASUREMENT, CONTROL, AND LABORATORY USE - PART 2-081: PARTICULAR REQUIREMENTS FOR AUTOMATIC AND SEMI-AUTOMATIC LABORATORY EQUIPMENT FOR ANALYSIS AND OTHER PURPOSES
- EN 61010-2-101:2017 SAFETY REQUIREMENTS FOR ELECTRICAL EQUIPMENT FOR MEASUREMENT, CONTROL, AND LABORATORY USE - PART 2-101: SAFETY REQUIREMENTS FOR IN VITRO DIAGNOSTIC (IVD) MEDICAL EQUIPMENT

| EN 61326-1:2013   | ELECTRICAL EQUIPMENT FOR MEASUREMENT, CONTROL AND<br>LABORATORY USE - EMC REQUIREMENTS - PART 1: GENERAL<br>REQUIREMENTS  |  |
|---|---|--|
| EN 61326-2-6:2013   | ELECTRICAL EQUIPMENT FOR MEASUREMENT, CONTROL AND<br>LABORATORY USE - EMC REQUIREMENTS - PART 2-6: PARTICULAR<br>REQUIREMENTS - IN VITRO DIAGNOSTIC (IVD) MEDICAL EQUIPMENT |  |
| AND THEREFORE MEETS THE ESSENTIAL REQUIREMENTS OF THE FOLLOWING COMMUNITY<br>DIRECTIVES AND THEIR AMENDMENTS:       |   |  |
| LOW VOLTAGE DIRECTIVE (2014/35/EU)  |   |  |
| ELECTROMAGNETIC COMPATIBILITY DIRECTIVE (2014/30/EU)  |   |  |
| RESTRICTION OF HAZARDOUS SUBSTANCES IN ELECTRICAL AND ELECTRONIC EQUIPMENT<br>DIRECTIVE – ROHS2 (2011/65/EU)        |   |  |
| NOTIFIED BODY:  | NOT NECESSARY   |  |
| (EC) CERTIFICATE:   | N.A.  |  |
| START OF CE-MARKING   | : FEBRUARY 2018   |  |
| REVISION:   | 6   |  |
| PLACE, DATE OF ISSUE  | MONTERIGGIONI, 25 MAY 2022  |  |
| EXPIRY DATE:  | 25 MAY 2027   |  |
| THE PRESENT EC DECLARATION OF CONFORMITY IS ISSUED UNDER THE SOLE AND EXCLUSIVE RESPONSIBILITY OF THE MANUFACTURER. |   |  |
|   | Chhi  |  |
| SIGNATURE:  | CHIARA MUZZI  |  |
|   | REGULATORY AFFAIRS MANAGER  |  |
|   |   |  |
| THIS IS THE CERTIFIED COPY OF THE ORIGINAL DOCUMENT STORED IN ARCHIVE OF DIESSE DIAGNOSTICA SENESE SPA              |   |  |
| ISSUED: MONTERIGGIONI,  | 25/05/2022 M. Storks  |  |

MAGDALENA STOCZKO REGULATORY SUPERVISOR