DIESSE	DECLARATION OF CONFORMITY
MANUFACTURER:	DIESSE DIAGNOSTICA SENESE SPA VIA DELLE ROSE 10 53035 MONTERIGGIONI (SI), ITALY
EUROPEAN REPRESENTATIVE:	//
GENERIC NAME	AUTOMATIC INSTRUMENT FOR ESR ANALYSIS
PRODUCT: CODE:	CUBE 30 touch 10395
TECHNICAL DATA:	110-230 Vac (50-60 Hz); Pwr: 100 VA
CLASSIFICATION:	IVD 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 D	DESIGN, TYPE OF MANUFACTURE OF THE IN VITRO MEDICAL

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
	MEASURE	MENT, CONTROL,	AND LA	BORATORY USE	- PART 1: GEN	IERAL
	REQUIREN	MENTS				

- 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 61010-2-101:2002** 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 LABORATORY USE - EMC REQUIREMENTS - PART 1: GENERAL REQUIREMENTS				
EN 61326-2-6:2006	ELECTRICAL EQUIPMENT FOR MEASUREMENT, CONTROL AND LABORATORY USE - EMC REQUIREMENTS - PART 2-6: PARTICULAR REQUIREMENTS - IN VITRO DIAGNOSTIC (IVD) MEDICAL EQUIPMENT				
AND THEREFORE MEETS THE ESSENTIAL REQUIREMENTS OF THE FOLLOWING COMMUNITY 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 DIRECTIVE – ROHS2 (2011/65/EU)					
NOTIFIED BODY:	NOT NECESSARY				
(EC) CERTIFICATE:	N.A.				
START OF CE-MARKING	G: FEBRUARY 2018				
REVISION:	2				
PLACE, DATE OF ISSUE	: MONTERIGGIONI, 26 NOVEMBER 2020				
EXPIRY DATE:	25 MAY 2022				
THE PRESENT DECLARATION OF CONFORMITY IS ISSUED UNDER THE SOLE AND EXCLUSIVE RESPONSIBILITY OF THE MANUFACTURER.					
	Ch Li				
SIGNATURE:					
	QUALITY & REGULATORY MANAGER				
THIS IS THE CERTIFIED COPY OF THE ORIGINAL DOCUMENT STORED IN ARCHIVE OF DIESSE DIAGNOSTICA SENESE SPA					
ISSUED: MONTERIGGIONI,	26/11/2020 M. Stoales				

MAGDALENA STOCZKO REGULATORY SUPERVISOR