

## **EC DECLARATION OF CONFORMITY**

MANUFACTURER: DIESSE DIAGNOSTICA SENESE SPA

STRADA DEI LAGHI 39 53035 MONTERIGGIONI (SI)

**ITALY** 

EUROPEAN REPRESENTATIVE:

GENERIC NAME RFID TRANSPONDER DEVICES

PRODUCT AND CODE TEST DEVICE 10K 10290

TEST DEVICE 5K 10291
TEST DEVICE 1K 10292
TEST DEVICE NEXT 10K 10297
TEST DEVICE NEXT 5K 10296
TEST DEVICE NEXT 1K 10294
TEST DEVICE NEXT 500 10293
TEST DEVICE NEXT 50 10295

TEST DEVICE ORIGINAL 4K 10270

WE HEREWITH DECLARE THAT TEST DEVICE PRODUCTS COMPLY WITH DIRECTIVE 2014/53/EU AND THE RELEVANT APPLICABLE SECTOR STANDARDS LISTED BELOW, WHEN USED WITH THE ESR ANALYZERS PRODUCED BY DIESSE DIAGNOSTICA SENESE S.P.A.

ETSI EN 300 330 V2.1.1 Short Range Devices (SRD); Radio equipment in the frequency range 9

kHz to 25 MHz and inductive loop systems in the frequency range 9 kHz to 30 MHz; Harmonised Standard covering the essential requirements of

article 3.2 of Directive 2014/53/EU

ETSI EN 301 489-1 V1.9.2 Electromagnetic compatibility and Radio spectrum Matters (ERM);

ElectroMagnetic Compatibility (EMC) standard for radio equipment and

services; Part 1: Common technical requirements

ETSI EN 301 489-3 V2.1.1 ElectroMagnetic Compatibility (EMC) standard for radio equipment and

services; Part 3: Specific conditions for Short-Range Devices (SRD) operating on frequencies between 9 kHz and 246 GHz; Harmonised Standard covering the essential requirements of article 3.1(b) of Directive

2014/53/EU

EN 55035:2017+A11:2020 Electromagnetic compatibility of multimedia equipment. Immunity

requirements

EN 61000-4-2:2009 Electromagnetic compatibility (EMC) - Part 4-2: Testing and

measurement techniques - Electrostatic discharge immunity test

EN 61000-4-3:2006 Electromagnetic compatibility (EMC) - Part 4-3: Testing and

measurement techniques - Radiated, radio-frequency, electromagnetic

field immunity test

+A1:2008+A2:2010

(2016-11)

(2011-09)

(2019-03)

START OF CE-MARKING: NOVEMBER 2021

EXCEPT:

2022: TEST DEVICE ORIGINAL 4K

REVISION: 2

PLACE, DATE OF ISSUE: MONTERIGGIONI, 03 OCTOBER 2022

EXPIRY DATE: //

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

SIGNATURE:

CHIARA MUZZI REGULATORY AFFAIRS MANAGER

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ISSUED: MONTERIGGIONI, 03/10/2022

MAGDALENA STOCZKO REGULATORY SUPERVISOR

Stoules



MANUFACTURER:	DIESSE DIAGNOSTICA SENESE SPA STRADA DEI LAGHI 39 53035 MONTERIGGIONI (SI), ITALY
EUROPEAN REPRESENTATIVE:	//
PRODUCT: CODE:	ESR CONTROL CUBE 10436
CLASSIFICATION:	IVD NOT IN ANNEX II OR SELF-TESTING IVD
CONFORMITY ASSESSMENT ROUTE:	ANNEX APPLIED N° III
WE HEREWITH DECLARE THAT THE ABOVE MENTIONED PRODUCTS MEET THE PREVISIONS OF THE COUNCIL DIRECTIVE 98/79EC FOR IN VITRO DIAGNOSTIC MEDICAL DEVICE. ALL SUPPORTING DOCUMENTATION IS RETAINED UNDER THE PREMISES OF THE MANUFACTURER.	
NOTIFIED BODY:	NOT NECESSARY
(EC) CERTIFICATE:	N.A.
START OF CE-MARKING:	NOVEMBER 2016
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.	
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SIGNATURE:	CHIARA MUZZI REGULATORY AFFAIRS MANAGER
	REGULATORT AFFAIRS MANAGER

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ISSUED: MONTERIGGIONI, 25/05/2022

> MAGDALENA STOCZKO REGULATORY SUPERVISOR

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