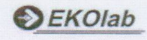


Certificat Certificate

N° 2007/28642.5

AFNOR Certification certifies that the management system implemented by: AFNOR Certification удостоверяет, что система менеджмента организации:







for the following activities: для следующих областей деятельности:

DEVELOPMENT, PRODUCTION, STORAGE AND SALE OF MEDICAL DEVICES FOR IN-VITRO DIAGNOSTICS.

РАЗРАБОТКА, ПРОИЗВОДСТВО, ХРАНЕНИЕ И РЕАЛИЗАЦИЯ МЕДИЦИНСКИХ ИЗДЕЛИЙ ДЛЯ IN-VITRO ДИАГНОСТИКИ.

> has been assessed and found to meet the requirements of: проверена и признана соответствующей требованиям стандарта:

#### ISO 13485:2016

and is developed on the following locations: и действует на следующих площадках.

142530, RUSSIA, MOSCOW REGION, ELEKTROGORSK CITY, Budennogo str., 1-1A 142530, РОССИЯ, МОСКОВСКАЯ ОБЛАСТЬ, г. ЭЛЕКТРОГОРСК, ул. Буденного, 1-1А

This certificate is valid from (year/month/day) Данный сертификат действителен с (год/ месяц/ день)

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STED130-2017 vs. 01



Declaration of Conformity

### DECLARATION OF CONFORMITY

#### 1) Manufacturer (Name, department): CJSC EKOlab

Address: 1 Budennogo Str., Elektrogorsk, Moscow region, 142530, Russia

1 6

# 2) European authorized representative: CEpartner4U BV,

Address: ESDOORNLAAN 13, 3951DB MAARN, THE NETHERLANDS; (on product labels printed as: CEpartner4U, ESDOORNLAAN 13, 3951DB MAARN, THE NETHERLANDS. www.cepartner4u.com)

3) Product(s) (name, type or model/batch number, etc.):

Rabbit plasma

### 4) The product(s) described above is in conformity with:

Title	Document No.	
In vitro Diagnostic Medical Devices		98/79/EC
Directive		

5) Additional information (conformity procedure, Notified Body, CE certificate, etc.):

Conformity assessment procedure for CE marking: In vitro Diagnostic Medical Device Directive, Annex III

Registration nr. : pending

Elektrogorsk, Russia; 2017-11-03

(Place & date of issue (yyyy-mm-dd))



(name; function and signature of manufacturer)



Declaration form: Standard ISO/IEC 17050-1:2010



Declaration of Conformity

STED130-2017 vs. 01

Page: 2 of 2

## Appendix

7 4

Date: 2017-11-08

List of devices.

8

Device name	Type/ model/ref number	Risk class / rule <sup>1</sup>	Code: EMDS/GMDN	First date of CE- compliance
Rabbit plasma		Low risk	15011290/0	2017-11-08



<sup>1</sup> See EDMS codes: <u>http://www.edma-ivd.be/</u> (products classification)/Preference GMDN code

Cepartner4U

Certificate number: 2017-IVD/193

## **Certificate of CE-Notification**

This is to certify that, in accordance with the *In Vitro* Diagnostic Medical Device Directive 98/79/EC, **CEpartner4U BV** agrees to perform all duties and responsibilities as the Authorized Representative for

### CJSC EKOlab

1 Budennogo Str., Elektrogorsk, Moscow region, 142530, Russia

as stipulated and demanded by the aforementioned Directive. The Dutch Competent Authorities have accepted the manufacturer's medical device registrations by CEpartner4U as listed on the product list attached to the manufacturer's Declaration of Conformity:

### Device group: Rabbit plasma

IVD devices were registered under number:

Registration number Rabbit plasma: NL-CA002-2017-43242

#### with Dutch Competent Authorities as a consequently this IVD devices were entered in EUDAMED by Dutch Competent Authorities

The manufacturer has provided CEpartner4U with all necessary documentation, together with an appropriate Declaration of Conformity that the IVD medical devices fulfil the essential requirements of Directive 98/79/EC.

2017-12-18

Olga Teirlinck Consultant CEpartner4U BV

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Esdoornlaan13 3951 DB Maarn NL tel: +31 (0)343 442 524 www.cepartner4u.nl

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