

FEDERAL SERVICE FOR SUPERVISION OF CONSUMER RIGHTS PROTECTION AND HUMAN WELFARE

FEDERAL BUDGET INSTITUTE OF SCIENCE  
«CENTRAL RESEARCH INSTITUTE FOR EPIDEMIOLOGY»

111123, Moscow, 3A Novogireevskaya street, Tel.: +7 495 974 96 42, Fax: +7 495 305 54 23,  
e-mail: obtk@pcr.ru



EC DECLARATION OF CONFORMITY

Directive 98/79/EC of the European Parliament and of the Council of 27<sup>th</sup> of October 1998 on  
*In Vitro* Diagnostic Medical Devices

Federal Budget Institute of Science “Central Research Institute for Epidemiology” hereby under own responsibility declares that the products covered by the declaration conform with Essential Requirements listed in Annex I of EC Directive 98/79/EC (IVD Directive). Supporting documentation is retained under the premises of the manufacturer.

The quality management system meets the requirements of the standard EN ISO 13485 “Medical devices – Quality management systems – Requirements for regulatory purposes” and is certified by Institute for testing and certification, Inc. (certificate No. 21 0023 SJ, valid until 26.04.2024).

<b>Manufacturer:</b>	Federal Budget Institute of Science “Central Research Institute for Epidemiology”
<b>Authorized Representative:</b>	Ecoli Dx, s.r.o. Purkyňova 74/2 Praha 1, 110 00 Czech Republic Tel: +420 325 209 912 Cell: +420 739 802 523 E-mail: ecoli@ecoli.sk
<b>Product Name:</b>	Annex for this Declaration
<b>Description:</b>	Reagent kits for qualitative detection and quantification of DNA (RNA) of different infectious agents or HLA B*5701 DNA in human specimens
<b>Classification:</b>	Article 9, paragraph 3 of EC Council Directive 98/79/EC on <i>in Vitro</i> Diagnostic Devices Annex II List B IVDs (According to EC Declaration of Conformity List)
<b>Conformity Assessment Route:</b>	Annex IV (IVDD) Full QA System
<b>Notified Body:</b>	Institute for testing and certification, Inc. třída Tomáše Bati 299 Louky, 763 02 Zlin, Czech Republic E-mail: itc@itczlin.cz Notified Body No. 1023
<b>EC Certificate:</b>	No. 11 0040 QS/NB revision j, valid until 2024-05-26
<b>Place, Date of Issue:</b>	Zlin, Czech Republic, 2021-04-27

Signed

Full name: Vasily G. Akimkin  
Title: Director



Valid from 2021-04-27

Valid until 2024-05-26

№№№	Description	Model(s)
1.	AmpliSens® <i>Rubella virus</i> -FRT PCR kit	variant FRT-50 F
2.	AmpliSens® <i>Toxoplasma gondii</i> -FRT PCR kit	variant FRT-50 F
3.	AmpliSens® <i>CMV</i> -FEP PCR kit	variant FEP (0.2-ml tubes)
4.	AmpliSens® <i>CMV</i> -FRT PCR kit	variant FRT-100 F
5.	AmpliSens® <i>HSV / CMV</i> -MULTIPRIME-FRT PCR kit	variant FRT-100 F
6.	AmpliSens® <i>CMV</i> -screen/monitor-FRT PCR kit	variant FRT-100 F
7.	AmpliSens® <i>EBV / CMV / HHV6</i> -screen-FRT PCR kit	variant FRT-100 F
8.	AmpliSens® <i>Chlamydia trachomatis</i> -FEP PCR kit	variant FEP (0.2-ml tubes)
9.	AmpliSens® <i>Chlamydia trachomatis</i> -FRT PCR kit	variant FRT variant FRT-100 F
10.	AmpliSens® <i>C.trachomatis / Ureaplasma / M.genitalium</i> -MULTIPRIME-FEP PCR kit	variant FEP (0.2-ml tubes)
11.	AmpliSens® <i>C.trachomatis / Ureaplasma / M.genitalium</i> -MULTIPRIME-FRT PCR kit	variant FRT-100 F
12.	AmpliSens® <i>C.trachomatis / Ureaplasma / M.hominis</i> -MULTIPRIME-FEP PCR kit	variant FEP (0.2-ml tubes)
13.	AmpliSens® <i>C.trachomatis / Ureaplasma / M.hominis</i> -MULTIPRIME-FRT PCR kit	variant FRT-100 F
14.	AmpliSens® <i>C.trachomatis / Ureaplasma / M.genitalium / M.hominis</i> -MULTIPRIME-FRT PCR kit	variant FRT-100 F
15.	AmpliSens® <i>N.gonorrhoeae / C.trachomatis / M.genitalium / T.vaginalis</i> -MULTIPRIME-FRT PCR kit	variant FRT-100 F
16.	AmpliSens® <i>N.gonorrhoeae / C.trachomatis / M.genitalium</i> -MULTIPRIME-FRT PCR kit	variant FRT-100 F
17.	AmpliSens® Genoscreen HLA B*5701-FRT PCR kit	variant FRT
18.	AmpliSens® <i>Mycoplasma pneumoniae / Chlamydophila pneumoniae</i> -FEP PCR kit	variant FEP (0.2-ml tubes)
19.	AmpliSens® <i>Mycoplasma pneumoniae / Chlamydophila pneumoniae</i> -FRT PCR kit	variant FRT-100 F
20.	AmpliSens® <i>T.vaginalis / N.gonorrhoeae / C.trachomatis</i> -MULTIPRIME-FRT PCR kit	variant FRT-100 F