



For Professional Use Only

# AmpliSens® hRSV-FEP PCR kit Instruction Manual

# **AmpliSens**®



Ecoli s.r.o., Studenohorska 12 841 03 Bratislava 47 Slovak Republic

Tel.: +421 2 6478 9336 Fax: +421 2 6478 9040



Federal Budget Institute of Science "Central Research Institute for Epidemiology" 3A Novogireevskaya Street Moscow 111123 Russia

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#### 1. INTENDED USE

**AmpliSens®** *hRSV*-FEP PCR kit is an *in vitro* nucleic acid amplification test for qualitative detection of *Human Respiratory Syncytial Virus* RNA in the clinical materials (nasal and oropharyngeal swabs, nasopharyngeal or tracheal sputum or aspirate, and autopsy material) by using end-point hybridization-fluorescence detection of amplified products.



The results of PCR analysis are taken into account in complex diagnostics of disease.

#### 2. PRINCIPLE OF PCR DETECTION

Human Respiratory Syncytial Virus detection by the polymerase chain reaction (PCR) is based on the amplification of the pathogen genome specific region using special Human Respiratory Syncytial Virus primers. In Fluorescent End-Point PCR, the amplified product is detected by using fluorescent dyes. These dyes are linked to oligonucleotide probes that bind specifically to the amplified product during thermocycling. A multichannel rotor-type fluorometer is specially designed to detect fluorescence emission from the fluorophores in a reaction mixture after PCR. It allows detection of the accumulating product without re-opening the reaction tubes after the PCR run. AmpliSens® hRSV-FEP PCR kit is a qualitative test that contains the Internal Control (IC). It must be used in the extraction procedure in order to control the extraction process of each individual sample and to identify possible reaction inhibition. AmpliSens® hRSV-FEP PCR kit uses "hot-start", which greatly reduces the frequency of nonspecifically primed reactions. "Hot-start" is guaranteed by separation of nucleotides and Taq-polymerase by using a wax layer. Wax melts and reaction components mix only at 95 °C.

#### 3. CONTENT

AmpliSens® hRSV-FEP PCR kit is produced in 2 forms:

AmpliSens® hRSV-FEP PCR kit (tubes of 0.2 ml), REF V37-50-R0,2-FEP-CE.

AmpliSens® *hRSV*-FEP PCR kit (tubes of 0.5 ml), **REF** V37-50-R0,5-FEP-CE.

AmpliSens® hRSV-FEP PCR kit includes:

Reagent	Description	Volume (ml)	Amount
PCR-mix-1-FEP/FRT hRSV ready-to-use single-dose test tubes (under wax)	colorless clear liquid	0.008	55 tubes of 0.5 or 0.2 ml
PCR-mix-2-FL	colorless clear liquid	0.77	1 tube
PCR-mix-Background	colorless clear liquid	0.5	1 tube
Mineral oil for PCR	colorless viscous liquid	4.0	1 dropper bottle
Positive Control cDNA hRSV-Flu (C+hRSV-Flu)	colorless clear liquid	0.1	1 tube
Positive Control STI-88 (CS+)	colorless clear liquid	0.1	1 tube
TE-buffer	colorless clear liquid	0.5	1 tube
Negative Control (C-)*	colorless clear liquid	1.2	1 tube
Internal Control STI-rec (IC)**	colorless clear liquid	0.12	5 tubes

<sup>\*</sup> must be used in the extraction procedure as Negative Control of Extraction.

AmpliSens® hRSV-FEP PCR kit is intended for 55 reactions, including controls.

#### 4. ADDITIONAL REQUIREMENTS

- RNA extraction kit.
- Reverse transcription kit.
- Disposable powder-free gloves and laboratory coat.
- Pipettes (adjustable).
- Sterile pipette tips with aerosol barriers (up to 200 μl).
- Tube racks.
- Vortex mixer.
- Desktop centrifuge with a rotor for 2-ml reaction tubes.
- PCR box.
- Personal thermocyclers (for example, Gradient Palm Cycler (Corbett Research, Australia),
   GeneAmp PCR System 2700 (Applied Biosystems, USA), MyCycler (Bio-Rad, USA), Uno-2 (Biometra, Germany), or equivalent).
- Fluorometer (for example, ALA-1/4 (Biosan, Latvia), or equivalent).
- Disposable polypropylene microtubes for PCR (0.5- or 0.2-ml; for example, Axygen, USA).
- Refrigerator for 2–8 °C.
- Deep-freezer for ≤ -16 °C.
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<sup>\*\*</sup> add 10 µl of Internal Control STI-rec (IC) during the RNA extraction procedure directly to the sample/lysis mixture (see RIBO-sorb REF K2-1-Et-50-CE, RIBO-prep, REF K2-9-Et-50-CE protocols).

· Waste bin for used tips.

#### 5. GENERAL PRECAUTIONS

The user should always pay attention to the following:

- Use sterile pipette tips with aerosol barriers and use new tip for every procedure.
- Store and handle amplicons away from all other reagents.
- Thaw all components thoroughly at room temperature before starting detection.
- When thawed, mix the components and centrifuge briefly.
- Use disposable gloves, laboratory coats, protect eyes while samples and reagents handling. Thoroughly wash hands afterward.
- Do not eat, drink, smoke, apply cosmetics, or handle contact lenses in laboratory work areas.
- Do not use a kit after its expiration date.
- Dispose of all samples and unused reagents in compliance with local authorities' requirements.
- Samples should be considered potentially infectious and handled in a biological cabinet in accordance with appropriate biosafety practices.
- Clean and disinfect all sample or reagent spills using a disinfectant, such as 0.5 % sodium hypochlorite or another suitable disinfectant.
- Avoid contact with the skin, eyes, and mucosa. If skin, eyes, or mucosa contact, immediately flush with water and seek medical attention.
- Material Safety Data Sheets (MSDS) are available on request.
- Use of this product should be limited to personnel trained in DNA amplification techniques.
- The laboratory process must be one-directional, it should begin in the Extraction Area and then move to the Amplification and Detection Area. Do not return samples, equipment, and reagents to the area in which the previous step was performed.



Some components of this kit contain sodium azide as a preservative. Do not use metal tubing for reagent transfer.

#### 6. SAMPLING AND HANDLING



Obtaining samples of biological materials for PCR-analysis, transportation and storage is described in manufacturer's handbook [1]. It is recommended that this handbook is read before starting work.

**AmpliSens®** *hRSV-FEP* PCR kit is intended for the analysis of RNA extracted with RNA extraction kits from nasal and oropharyngeal swabs, nasopharyngeal or tracheal sputum or aspirate, and autopsy material.

<u>Nasal swabs</u> are taken using a probe with a dry cotton swab. Make the patient blow his nose if it is filled with mucus. Insert the probe gently along the external nasal wall by 2–3 cm

till the inferior nasal concha. Then move the probe slightly lower, insert in the inferior nasal meatus under the inferior nasal concha, rotate, and pull it back along the external nasal wall.

When material is obtained, place the working area of the probe with a cotton swab into a sterile disposable tube with 500 µl of transport medium for storage and transportation of respiratory swabs. Break off the terminal part of the probe or cut it off to allow tight closing of the tube cup. Close the tube with the solution and the working area of the probe.

<u>Oropharyngeal swabs</u> are taken using a probe with a dry cotton swab. Obtain swabs by rotating the probe over the surface of tonsils, palatine arches, and the posterior wall of the pharynx after gargling of the oral cavity with water.

When material is obtained, place the working area of the probe with a cotton swab into a sterile disposable tube with 500 µl of transport medium for storage and transportation of respiratory swabs. Break off the terminal part of the probe or cut it off to allow tight closing of the tube cup. Close tube with the solution and the working area of the probe.



It is recommended to combine nasal and throat swabs. To do this, working parts of probes are placed into one tube with 500 µl of medium for storage and transportation of respiratory swabs and studied as one sample.

The <u>sputum</u> is collected into a sterile disposable container after preliminary gargling of the oral cavity with water. <u>Nasopharyngeal or tracheal aspirates</u> are taken by the conventional method and placed in a sterile disposable container.

<u>Sectional material</u> is placed in a sterile disposable container. Material is to be frozen after sampling or analyzed within 1 h.



Only one freeze-thaw cycle is allowed.

#### **Pretreatment of material**

Any operation with studied material is carried out in compliance with local authority's requirements.

All pretreatment manipulations are carried out using adjustable pipettes with disposable tips with aerosol barriers and disposable polypropylene tubes (1.5 ml or 10.0 ml). Disposable plastic dishes (tubes and tips) are to be discarded into a special container with a suitable disinfectant and utilized in compliance with local authority's requirements.

<u>Swabs</u> are used without preliminary preparation.

Sputum must be treated with Mucolysin reagent REF 180-CE according to the Mucolysin instruction manual. 100 µl of the pretreated sputum is used for RNA extraction.

Section material is homogenized with a sterile porcelain mortar and pestle, with subsequent preparation of a 10 % suspension in sterile saline or phosphate buffer. Transfer the suspension to 1.5-ml tube and centrifuge for 5 min at 10,000 rpm. 100 µl of the supernatant is used for RNA extraction. To repeat the test later, freeze the remained

suspension at  $\leq$  – 16 °C.

Additional reagents manufactured by Federal Budget Institute of Science "Central Research Institute for Epidemiology" are required:

- 1. Transport Medium for Storage and Transportation of Respiratory Swabs (unaliquoted)

  REF 959-CE is used for sampling and storage of nasal and oropharyngeal swabs.
- 2. Mucolysin reagent REF 180-CE is used for pretreatment of sputum and aspirates.

#### 7. WORKING CONDITIONS

AmpliSens® hRSV-FEP PCR kit should be used at 18-25 °C.

#### 8. PROTOCOL

#### 8.1. RNA extraction

It is recommended to use the following nucleic acid extraction kits:

- RIBO-sorb, **REF** K2-1-Et-50-CE.
- RIBO-prep, **REF** K2-9-Et-50-CE.



Extract RNA according to the manufacturer's instructions.

Add 10 µl of Internal Control STI-rec to each tube.

#### 8.2. Reverse transcription

It is recommended to use the following kit for complementary DNA (cDNA) synthesis from RNA:

• REVERTA-L, **REF** K3-4-50-CE.



Carry out the reverse transcription according to the manufacturer's instructions.

#### 8.3. Preparing the PCR

The total reaction volume is 25  $\mu$ I, the volume of cDNA sample is 10  $\mu$ I.

#### 8.3.1. Preparing tubes for PCR

- Prepare the required number of tubes with PCR-mix-1-FEP/FRT hRSV and wax for amplification of cDNA from clinical and control samples.
- 2. Add **7 μl** of **PCR-mix-2-FL** to the surface of the wax layer of each tube ensuring that it does not fall under the wax and mix with **PCR-mix-1-FEP/FRT** *hRSV*.
- 3. Add above 1 drop of mineral oil for PCR (about 25 μl).
- 4. Prepare 2 tubes with PCR-mix-1-FEP/FRT hRSV and mark them as Background. Add 17 µI of PCR-mix-Background to the surface of the wax layer of each tube ensuring that it does not fall under the wax and mix with PCR-mix-1-FEP/FRT hRSV. Add above 1 drop of mineral oil for PCR.

- 5. Using tips with aerosol barrier, add **10 μl** of **cDNA samples** obtained in RNA reverse transcription reaction.
- 6. Carry out the control amplification reactions:

NCA - Add 10  $\mu l$  of TE-buffer to the tube labeled NCA (Negative Control of

Amplification).

C+<sub>hRSV-Flu</sub> - Add 10 μI of Positive Control cDNA hRSV-Flu to the tube labeled C+<sub>hRSV-</sub>

Flu (Positive Control of Amplification).

CS+ - Add 10 μI of Positive Control STI-88 to the tube labeled CS+.

#### 8.3.2. Amplification

Run the following program in the thermocycler (see Table 1). When the temperature reaches 95 °C (pause mode), insert tubes into cells of the thermocycler and press the button to continue.

It is recommended to sediment drops from walls of tubes by short centrifugation (1–3 s) before placing them in the thermocycler.

Table 1
Programming thermocyclers at cDNA amplification of *Human Respiratory Syncytial virus* 

	Thermocyclers with active temperature adjustment:				nent:				
Terzik (DNA-Technology)		GeneAmp PCR System 2700 (Applied Biosystems), Gradient Palm Cycler (Corbett Research), MyCycler (Bio-Rad)			Thermocyclers with block temperature adjustment:  Biometra				
Step	Tempe- rature	Time	Cycles	Tempe- rature	Time	Cycles	Tempe- rature	Time	Cycles
0	95 °C	pause		95 °C	pause		95 °C	pause	
1	95 °C	5 min	1	95 °C	5 min	1	95 °C	5 min	1
	95 °C	10 s		95 °C	10 s		95 °C	25 s	
2	54 °C	20 s	42	54 °C	25 s	42	54 °C	40 s	42
	72 °C	10 s		72 °C	25 s		72 °C	25 s	
3	72 °C	1 min	1	72 °C	1 min	1	72 °C	1 min	1
4	10 °C	stora	age	10 °C	sto	rage	10 °C	stora	age

#### 9. DATA ANALYSIS

Detection is conducted with an ALA-1/4 fluorescence detector.



Please read Ala-1 Operating Manual before use of this kit.

Program the detector according to the manufacturer's manual and Guidelines.

#### 9.1. Results interpretation

1. When the analysis is complete the results are automatically shown in the table as follows:

pos – positive result;

neg – negative result;

eq – equivocal result (signal is in grey zone);

**nd** – invalid result (specific signal and IC signal are absent in the sample).

2. The result of analysis is considered reliable only if the results obtained for Positive and

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#### Results for controls

		Result of autom		
Control	Stage for control	FAM channel	HEX channel	Interpretation
		(IC)	(samples)	
C-	RNA extraction	+	RSv - neg	OK
NCA	Amplification	-	RSv - nd	OK
C+ <sub>hRSV-Flu</sub>	Amplification	-	RSv - pos	OK
CS+	Amplification	+	RSv - neg	OK

#### 10. TROUBLESHOOTING

Results of analysis are not taken into account in the following cases:

- PCR and detection of the samples with the nd result (except for NCA) are to be repeated.
   If the same result is obtained, analysis should be repeated starting from the RNA extraction stage. The nd result for NCA is normal.
- 2. PCR and detection of the samples with the **eq** result are to be repeated. If the same result is obtained, the samples are considered to be **positive**.
- 3. If no positive signal is detected for Positive Controls of Amplification, this may indicate that the temperature profile programming of the used instrument was incorrect, the configuration of the PCR reaction was incorrect, the storage conditions for kit components did not comply with the manufacturer's instruction, or that the reagents kit has expired. Check programming of the used Instrument, storage conditions, and the expiration date of the reagents and then repeat PCR analysis.
- 4. If a positive signal is detected for C- in the HEX channel or for NCA in both channels, this indicates contamination of reagents or samples. In such cases, the results of analysis must be considered as **invalid**. Analysis should be repeated and measures to detect and eliminate the source of contamination are to be taken.

If you have any further questions or if you encounter problems, please contact our Authorized Representative in the European Community.

#### 11. TRANSPORTATION

AmpliSens® hRSV-FEP PCR kit should be transported at 2–8 °C for no longer than 5 days.

#### 12. STABILITY AND STORAGE

All components of the **AmpliSens**<sup>®</sup> *hRSV*-FEP PCR kit are to be stored at 2-8 °C when not in use. All components of the **AmpliSens**<sup>®</sup> *hRSV*-FEP PCR kit are stable until labeled expiration date. The shelf life of reagents before and after the first use is the same, unless otherwise stated.



PCR-mix-1-FEP/FRT *hRSV* is to be stored away from light.

#### 13. SPECIFICATIONS

#### 13.1. Sensitivity

Analytical Sensitivity of **AmpliSens**® *hRSV*-FEP PCR kit is not less than 1x10<sup>3</sup> genome equivalents per 1 ml of sample (GE/ml).



The claimed analytical features of **AmpliSens®** *hRSV*-FEP PCR kit are guaranteed only when additional reagents kits RIBO-sorb, RIBO-prep and REVERTA-L (manufactured by Federal Budget Institute of Science "Central Research Institute for Epidemiology") are used.

#### 13.2. Specificity

The analytical specificity of **AmpliSens**® *hRSV*-FEP PCR kit is assured by selection of specific primers and probes as well as stringent reaction conditions. The primers and probes were checked for possible homologies to all sequences published in gene banks by sequence comparison analysis. The clinical specificity of **AmpliSens**® *hRSV*-FEP PCR kit was confirmed in laboratory clinical trials.

#### 14. REFERENCES

 Handbook "Sampling, Transportation, and Storage of Clinical Material for PCR Diagnostics", developed by Federal State Institute of Science "Central Research Institute for Epidemiology" of Federal Service for Surveillance on Consumers' Rights Protection and Human Well-Being, Moscow, 2008.

#### 15. QUALITY CONTROL

In compliance with Federal Budget Institute of Science "Central Research Institute for Epidemiology" ISO 13485-Certified Quality Management System, each lot of **AmpliSens®** *hRSV-FEP* PCR kit has been tested against predetermined specifications to ensure consistent product quality.

# 16. KEY TO SYMBOLS USED

REF	Catalogue number	Σ	Sufficient for
LOT	Batch code		Expiration Date
IVD	In vitro diagnostic medical device	[]i	Consult instructions for use
VER	Version		Keep away from sunlight
	Temperature limitation	NCA	Negative control of amplification
	Manufacturer	C-	Negative control of extraction
	Date of manufacture	C+ <sub>hRSV-Flu</sub> , CS+	Positive control of amplification
EC REP	Authorised representative in the European Community	IC	Internal control
$\triangle$	Caution		

# **List of Changes Made in the Instruction Manual**

VER	Location of changes	Essence of changes
	Cover page	The phrase "For Professional Use Only" was added
	Intended use	The phrase "The results of PCR analysis are taken into account in complex diagnostics of disease" was added
	Content	New sections "Working Conditions" and "Transportation" were added
13.12.10		The "Explanation of Symbols" section was renamed to "Key to Symbols Used"
13.12.10	Stability and	The information about the shelf life of reagents before and after the first use was added
	Storage	Information that PCR-mix-1-FEP/FRT <i>hRSV</i> is to be kept away from light was added
	Key to Symbols Used	The explanation of symbols was corrected
Text	Positive Control STI was changed to Positive Control STI-88	
	TEXT	Positive Control cDNA <i>hRSV</i> -Flu (C+) was changed to Positive Control cDNA <i>hRSV</i> -Flu (C+ <sub>hRSV</sub> -Flu)
22.06.11 RT	Cover page, text	The name of Institute was changed to Federal Budget Institute of Science "Central Research Institute for Epidemiology"