RIBO-prep nucleic acid extraction kit



Instruction Manual

KEY TO SYMBOLS USED

REF Catalogue number Caution Contains sufficient for LOT Batch code <n> tests In vitro diagnostic medical IVD Use-by Date device Consult instructions for VER Version GHS02: Flame Temperature limit Manufacturer GHS05: Corrosion GHS07: Exclamation Date of manufacture mark

1. INTENDED USE

EC REP

RIBO-prep nucleic acid extraction kit is intended for extraction of total RNA/DNA from human biological material for subsequent testing by the nucleic acid amplification techniques (NAT):

- plasma of venous, umbilical cord blood
- serum of venous, umbilical cord blood,
- whole venous, umbilical cord blood
- leukocytes of venous, umbilical cord blood, cerebrospinal fluid (liquor),

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in the European Community

- discharge of the conjunctiva,
- saliva,
- nasopharyngeal and oropharyngeal swabs. sputum,
- endotracheal aspirate bronchial washings
- bronchoalveolar lavage fluid,
- transsudates
- amniotic liquid,
- breast milk.
- pus / liquid or dense necrotic content,
- vesicles and pustules content,
- hair follicles, nail plates,
- tissue (biopsy, surgical, autopsy) native material, paraffin-embedded tissue (biopsy, surgical, autopsy) material,
- urogenital mucous discharge (swab, scrape),
- prostate secretion
- feces / fecal swab,
- anal canal/rectal mucosal discharge
- cultures of microorganisms, isolated by inoculation of human biological material.

 Reagent kit can be used for extraction of total RNA/DNA from ticks, mosquitoes, washes from

environmental objects, water; water sample concentrates (drinking, open sources, sewage) for carrying out prophylactic measures in order to prevent human diseases.

Indications and contra-indications for use of the reagent kit RNA/DNA extraction is used in preanalytical stage of *in vitro* diagnostics by NAT.

2. PRINCIPLE OF NUCLEIC ACID EXTRACTION

Test samples are treated by **Solution for Lysis** to destruct cell membranes, viral envelopes and other biopolymer complexes and release nucleic acids and cellular components. The dissolved RNA/ĎNA precipitates after addition of the **Solution for Precipitation** and centrifugation, while the other components of the lysed clinical material remain in the solution and removed with subsequent washes. The final stage of extraction is dissolution of the pellet in elution buffer, the purified RNA/DNA is transferred into the solution.

The obtained nucleic acid sample is purified and free from inhibitors of amplification, which provides high analytical sensitivity of NAT assay.

3. CONTENT

RIBO-prep nucleic acid extraction kit is produced in 1 form:

variant 100, REF K2-9-Et-100-CE.

variant 100 includes:			
Reagent	Description	Volume, ml	Quantity
Solution for Lysis	clear liquid from colorless to blue grey colour ¹	30	1 vial
Solution for Precipitation	colorless clear liquid	40	1 vial
Washing Solution 3	colorless clear liquid	50	1 vial
Washing Solution 4	Washing Solution 4 colorless clear liquid		1 vial
RNA-buffer	colorless clear liquid	1.2	8 tubes

Variant 100 is intended for RNA/DNA extraction from 100 samples, including controls. While working with nail plates and hair follicles variant 100 is intended for RNA/DNA extraction from 50 samples, including controls

4. ADDITIONAL REQUIREMENTS

- 1.5-ml disposable polypropylene screwed or tightly closed tubes. Screwing caps for tubes.
- Sterile DNase- and RNase-free pipette tips without filter (up to 100 and 200 μ l). Sterile RNase-free pipette tips with aerosol filters (up to 100 μ l, 200 μ l).
- Tube racks for 1.5-ml tubes
- Thermostat with working temperature for 25-100 °C.
- Desktop mircocentrifuge
- Vacuum aspirator with flask for removing supernatant.
- PCR box or Biological cabinet.
- Pipettes (adjustable).
- Refrigerator for 2-8 °C
- Deep-freezer at the temperature from minus 24 to minus 16 °C
- Reservoir for used tips.
 Disposable powder-free gloves and a laboratory coat.

5. GENERAL PRECAUTIONS

The user should always pay attention to the following:

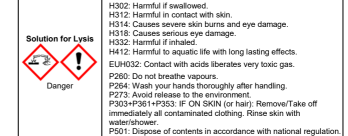
• Use sterile DNase- and RNase-free pipette filter tips and use a new tip for every

- procedure.
- Store all extracted positive material (specimens, controls and amplicons) away from all other reagents and add it to the reaction mix in a distantly separated facility
- Thaw all components thoroughly at room temperature before starting an assay When thawed, mix the components and centrifuge briefly.
- Use disposable protective gloves and laboratory cloths, and protect eyes while samples and reagents handling. Thoroughly wash hands afterwards.

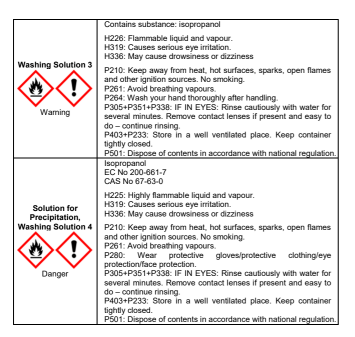
 Do not eat, drink, smoke, apply cosmetics, or handle contact lenses in laboratory work
- Do not use the kit if the internal packaging was damaged or its appearance was changed. Do not use the kit if the transportation and storage conditions according to the Instruction
- Manual were not observed.
- Do not use a kit after its expiration date.
- Dispose of all specimens and unused reagents in accordance with local regulations. Samples should be considered potentially infectious and handled in a biological cabinet
- in compliance 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 inhalation of vapors, samples and reagents contact with the skin, eyes, and mucous membranes. Harmful if swallowed. If these solutions come into contact, rinse the injured area immediately with water and seek medical advice if necessary.

 While observing the conditions of transportation, operation and storage, there are no risks
- of explosion and ignition. Safety Data Sheets (SDS) are available on request.
- The kit is intended for analysis of specified number of samples (see the section "Content"). The kit is ready for use in accordance with the Instruction Manual. Use the kit strictly for
- intended purpose
- Use of this product should be limited to personnel trained in DNA/RNA extraction techniques.
- Workflow in the laboratory must be one-directional, beginning in the Extraction Area and moving to the Amplification and Detection Area. Do not return samples, equipment and reagents to the area where the previous step was performed.

Solution for Lysis has an unpleasant smell. Work with this solution should be performed in a biological cabinet. Contains substance: guanidine thiocyanate



¹ If Solution for Lysis is stored at 2-8 °C, a crystalline precipitate may form.



6. SAMPLING AND HANDLING

See the information about the sampling, conditions of transportation and storage of the test material, the necessity and procedure of its pretreatment before DNA/RNA extraction in the Instruction manual for the PCR kit.

RIBO-prep nucleic acid extraction kit is recommended for RNA and DNA extraction and purification from:

• plasma of

- plasma of venous, umbilical cord blood serum of venous, umbilical cord blood,
- whole venous, umbilical cord blood, leukocytes of venous, umbilical cord blood,

- cerebrospinal fluid (liquor), discharge of the conjunctiva,

- saliva, nasopharyngeal and oropharyngeal swabs,
- sputum, endotracheal aspirate,
- bronchial washings, bronchoalveolar lavage fluid,
- pleural fluid. transsudates
- amniotic liquid,
- breast milk
- pus / liquid or dense necrotic content,
- vesicles and pustules content, hair follicles,
- nail plates
- tissue (biopsy, surgical, autopsy) native material,
- paraffin-embedded tissue (biopsy, surgical, autopsy) material, urogenital mucous discharge (swab, scrape),
- urine.
- prostate secretion,
- feces / fecal swab.
- anal canal/rectal mucosal discharge, cultures of microorganisms, isolated by inoculation of human biological material.
- washes from environmental objects.
- water; water sample concentrates (drinking, open sources, sewage).

Interfering substances and limitations of using test material samples

The information about potential interfering substances and limitations of using test material samples is specified in the Instruction Manual of the PCR kit.

<u>Potential interfering substances</u> Endogenous and exogenous substances that may be present in the biological material used

for the study were selected to assess potential interference (see Table 1).

Model samples of various biological material without adding and with the addition of potentially interfering substances were tested. The concentration of each potentially interfering substance is listed in Table 1.

	Table			Tuble I
Test material	Type of potential interferent	Potential interferent	Tested concentration in a sample	Interference presence
Plasma of venous, umbilical cord blood	Endogenous substances	Hemoglobin	5 g/l (upper limit of normal 1 g/l)	Not detected
		Triglycerides	37 mmol/l (upper limit of normal 3.7 mmol/l)	Not detected
		Bilirubin	210 µmol/l (upper limit of normal 21 µmol/l)	Not detected
		Protein	120 g/l (upper limit of normal 85 g/l)	Not detected
Serum of venous, umbilical cord	Endogenous substances	Bilirubin	210 µmol/l (upper limit of normal – 21 µmol/l)	Not detected
blood, whole venous, umbilical cord blood, leukocytes of venous, umbilical cord blood		Cholesterol	78 µmol/l (upper limit of normal – 7,8 µmol/l)	Not detected
		Triglycerides	37 mmol/l (upper limit of normal – 3.7 mmol/l)	Not detected
	Exogenous substances	Heparin	15 UI/ml	<u>Detected</u>
Cerebrospinal	l Exogenous	Whole blood	Up to 4 %	Not detected
fluid (liquor)	substances	Leukocytes	500 cells/mm ³	Not detected

To at most order	Type of	Potential	Tested concentration in	Interference	
Test material	potential interferent	interferent	a sample	presence	
Discharge of the conjunctiva	Exogenous substances	Tetracycline	5 mg/ml	Not detected	
tio conjunctiva	Endogenous	Mucin	5 %	Not detected	
Saliva, nasal and	substances	Lugol's solution	0.5 %	Not detected	
oropharyngeal	Exogenous	with glycerin Chlorhexidine	0.5 %	Not detected	
swabs	substances	bigluconate	2.5 %	Not detected	
		aqueous 2.3 %		. 101 40100104	
		Hypertonic sodium chloride	5 % (the drug volume to the volume of the	Not detected	
		solution	test sample)		
		Amoxicyclin+ clavulanic acid	1125 μg/ml + 281 μg/ml	Not detected	
		Cetrifaxone	1500 μg/ml	Not detected	
Sputum, endotracheal		Rifampicin Isoniazid	9-17.5 μg/ml 3-7 μg/ml	Not detected Not detected	
aspirate,	Exogenous substances	Ethambutol	2-5 μg/ml	Not detected	
bronchial washings,		Pyrazinamide	39 μg/ml	Not detected	
bronchoalveolar lavage fluid		Ofloxacin Kanamycin	5.2 μg/ml 22 μg/ml	Not detected Not detected	
3		Para-			
		aminosalicylic acid (PAS)	75 μg/ml	Not detected	
		Cycloserine	25-30 μg/ml	Not detected	
	Endogenous substances	Blood	1:1	Not detected	
Amniotic liquid	Exogenous	Cefazolin	64 µg/ml	Not detected	
•	substances Exogenous	sodium salt	16 % (the drug volume		
Brest milk	substances	"Miramistin"	to the volume of the test sample)	Not detected	
	Endogenous	Blood	1:1	Not detected	
	substances		16 % (the drug volume		
		Metronidasol	to the volume of the test sample)	Not detected	
		Rifampicin	9-17.5 µg/ml	Not detected	
D		Isoniazid	3-7 µg/ml	Not detected	
Pleural liquid, transsudates	Exogenous	Ethambutol Pyrazinamide	2-5 μg/ml 39 μg/ml	Not detected Not detected	
	substances	Ofloxacin	5.2 μg/ml	Not detected	
		Kanamycin	22 μg/ml	Not detected	
		Para- aminosalicylic	75 μg/ml	Not detected	
		acid (PAS)			
	Endogenous	Cycloserine Blood	25-30 μg/ml 1:1	Not detected Not detected	
	substances	Hemoglobin	250 g/l	Not detected	
		Metronidasol	16 % (the drug volume to the volume of the	Not detected	
			test sample)		
Pus /		Rifampicin	9-17.5 µg/ml	Not detected	
liquid or dense		Isoniazid Ethambutol	3-7 μg/ml 2-5 μg/ml	Not detected Not detected	
necrotic contents	Exogenous substances	Pyrazinamide	39 μg/ml	Not detected	
	Substances	Ofloxacin	5.2 μg/ml	Not detected	
		Kanamycin Para-	22 μg/ml	Not detected	
		aminosalicylic	75 μg/ml	Not detected	
		acid (PAS) Cycloserine	25-30 μg/ml	Not detected	
Vesicles and pustules	Exogenous	lodine (Potassium	0.5%	Not detected	
content	substances	iodide)	0.570	Not detected	
	Exogenous	"Hydrocortisone®" ointment for	1 % (the drug weight	Not detected	
Hair follicles	substances	external application	to the volume of the test sample)	Not detected	
	Exogenous	Chlorhexidine	/		
Nail plates	substances	bigluconate aqueous solution	2.5 %	Not detected	
Tissue (biopsy,	Evogenous	•			
surgical, autopsy) native	Exogenous substances	Vancomycin	1500 μg/ml	Not detected	
material Paraffin-		Formaline			
embedded tissue		sour	_	<u>Detected</u>	
(biopsy, surgical, autopsy) material	substances	Buffered formaline	_	Not detected	
Urogenital mucous	Endogenous substances Exogenous substances	Hemoglobin	260 μg/ml		
		Lactoferrin Glycogen	5 μg/ml 120 mg/ml	Not detected	
		Mucin	150 µg/ml		
discharge (swab, scrape)		Clotrimazole	16 % (the drug		
(Swab, scrape)		"Neomycin" + "Nystatin" +	volume to the volume of the test		
		"Polymixin B"	sample)		
	Endogenous substances	Albumin Blood	500 mg/ml 1:1	Not detected Not detected	
Urino		Azithromycin	0.8 mg/ml	Not detected	
Urine	Exogenous	"Rifampicin"	9-17.5 µg/ml	Not detected	
	substances	"Isoniazid" "Ethambutol"	3-7 μg/ml	Not detected	
		Euramputor	2-5 μg/ml	Not detected	

Test material	Type of potential interferent	Potential interferent	Tested concentration in a sample	Interference presence
		"Pyrazinamide"	39 μg/ml	Not detected
		"Ofloxacin"	5.2 μg/ml	Not detected
		"Kanamycin"	22 μg/ml	Not detected
		Para- aminosalicylic acid (PAS)	75 μg/ml	Not detected
		"Cycloserine"	25-30 µg/ml	Not detected
Prostate gland	Endogenous substances	Fructose	10 mg/ml	Not detected
secret	Exogenous substances	Ibuprofen	300 μg/ml	Not detected
		Blood	1:1	Not detected
	Endogenous substances	Mucin (mucus)	Up to 3 % (the drug weight to the volume of the test sample)	Not detected
		Fecal fats	40%	Not detected
		Loperamide	Up to 5 mg/ml	Not detected
	Exogenous substances	Hydrocortisone	Up to 3 % (the drug weight to the volume of the test sample)	Not detected
Feces/ fecal		Rifampicin	9-17.5 μg/ml	Not detected
swab		Isoniazid	3-7 µg/ml	Not detected
		Ethambutol	2-5 μg/ml	Not detected
		Pyrazinamide	39 μg/ml	Not detected
		Ofloxacin	5.2 μg/ml	Not detected
		Kanamycin	22 μg/ml	Not detected
		Para- aminosalicylic acid (PAS)	75 μg/ml	Not detected
		Cycloserine	25-30 μg/ml	Not detected
Anal canal/ rectal mucosal discharge	Exogenous substances	Glycerin	20%	Not detected
Ticks, mosquitoes	Endogenous substances	Bilirubin	210 µmol/l	Not detected
		Cholesterol	78 mmol/l	Not detected
		Triglycerides	37 mmol/l	Not detected
		Hemoglobin	5 g/l	Not detected
		Protein	120 g/l	Not detected
Water, water samples concentrates (drinking, open sources, sewage)	Exogenous substances	Feces	5 %	Not detected

7. WORKING CONDITIONS

RIBO-prep nucleic acid extraction kit should be used at the temperature from 20 to 28 $^{\circ}\text{C}$ and relative humidity from 15 to 75 %.

8. PROTOCOL

8.1. RNA and DNA Extraction

It is allowed to use another types of biological material, volumes of investigated sample according to the *Instruction manual* for the used PCR kit.

To work with RNA, it is necessary to use only disposable sterile plastic consumables that have a special marking RNase-free, DNase-free.

Procedure (for all types of biological material except nail plates and hair follicles) 1. Warm Solution for Lysis (if stored at 2-8 °C) at the temperature 65 °C until crystals

NOTE: Solution for lysis has unpleasant odor. Work in the PCR box

- 2. Take required number of 1.5-ml disposable tubes with tightly closable caps (including one tube for Negative Control of Extraction (C-) and one tube for Positive Control of Extraction (PCE) if they are provided for analysis). Mark the tubes.
- Add 10 μ I of Internal Control (if it is provided for analysis) to each tube and then add 300 μ I of Solution for Lysis.

NOTE: It is allowed to change the volume of Internal Control according to the Instruction manual for the used PCR kit.

4. Add 100 µl of prepared samples to the tubes with Solution for Lysis and Internal Control (if used) using pipette tips with aerosol filters. Add 100 µl of Negative Control (if it is provided for analysis) to the tube labeled C-. Add 90 µl of Negative Control and 10 µl of Positive Control (if it is provided for analysis) to the tube labeled PCE.

It is allowed to change the volume and dilution of Negative Control (C-) and

Positive Control of Extraction (PCE) according to the Instruction manual for NOTE: the used PCR kit.

- 5. Thoroughly mix the contents of the tubes by vortexing, then centrifuge tubes for 5 s to be sure there are no drops on the cap, and incubate them at 65 °C for 5 min.
 6. Add 400 μl of Solution for Precipitation, tightly close the tubes and mix them by vortexing.
- Centrifuge all tubes for 5 min at 13,400 rpm
- Carefully remove the supernatant without disturbing the pellet using a vacuum aspirator and 200-µl tips. Use a new tip for each tube.
- Add 500 µl of Washing Solution 3 to each tube, tightly close the tubes and turn them
 carefully upside down 3–5 times to wash the pellet. This procedure can be performed simultaneously for all the tubes: cover the tubes placed in a rack with a lid or another rack, press them, and turn the rack.
- 10. Centrifuge all the tubes at 13,400 rpm for 1–2 min.

 11. Carefully remove the supernatant without disturbing the pellet using a vacuum aspirator and 10-ul tips. Use a new tip for each tube.
- 12.Add **200 µl of Washing Solution 4** to each tube, tightly close the tubes and turn them carefully upside down and back 3–5 times to wash the pellet.

 13.Centrifuge all tubes at 13,400 rpm for 1–2 min.

 14.Carefully remove the supernatant without disturbing the pellet using a vacuum aspirator
- and 10-µl tips. Use a new tip for each tube.

 15.Incubate all tubes with open caps at 65 °C for 5 min (to dry the pellet).
- 16.Add 50 µl of RNA buffer into each tube. Mix the tubes by vortex. Then incubate them at 65 °C for 5 min occasionally stirring by vortex.

Increasing of the elution volume is allowed (for example up to 90 µl) NOTE: according to the Instruction manual for the used PCR kit

17. Centrifuge the tubes at 13,400 rpm for 1 min. The supernatant contains purified RNA and DNA. Samples are ready for reverse transcription and PCR.

Procedure (for nail plates and hair follicles)

- 1. Warm Solution for Lysis (if stored at 2-8 °C) at the temperature 65 °C until crystals disappear.
- 2. Take required number of 1.5 ml disposable tubes with tightly closable caps (including one tube for Negative Control of Extraction (C-) and one tube for Positive Control of Extraction (PCE) if they are provided for analysis). Mark the tubes.
- Add Internal Control (if it is provided for analysis) to each tube in a volume 1.5 times more than recommended volume for other types of biological material.

It is allowed to change the volume of Internal Control according to the Instruction manual for the used reagents kit for carrying out the amplification.

- 4. Add to each tube per one examination, 2 samples of nail plates (about 2x10 mm in size)
- or 2-3 hair follicles with hair shaft no more than 1-2 cm long. 5. Add 100 μ l of Negative Control (if it is provided for analysis) to the tube labeled C–. Add 85 μl of Negative Control and 15 μl of Positive Control (if it is provided for analysis) to the tube labeled PCE.

It is allowed to change the volume and dilution of **Negative Control (C–)** and **Positive Control of Extraction (PCE)** according to the *Instruction manual* for NOTE: the used PCR kit.

- 6. Add 600 μl of Solution for Lysis to the tubes. Make sure that test material is completely immersed into the solution. Tightly close the tubes
- Mix C- and PCE tubes thoroughly by vortexing, then centrifuge tubes for 5 s to be sure there are no drops on the cap.
- 8. Put the tubes with test and control samples to the thermostat and incubate them at 65 °C
- 9. After incubation mix thoroughly by vortexing, then centrifuge tubes for 2 min at 13,000 rpm.
- 10.Take necessary number of 1.5-ml disposable polypropylene screwed tubes (including controls of the extraction if they are provided for analysis). Mark the tubes.
- 11.Carefully remove the supernatant and add **400 µl** of it to the prepared tubes. 12.Continue the extraction according to the protocol for other biological materials (see above), starting with entry 5.

The purified RNA/DNA can be stored:

– at 2-8 °C for 24 h:

- at the temperature from minus 24 to minus 16 °C for 1 year.

8.2. Amplification

It's recommended to use AmpliSens® PCR kits and REVERTA-L reverse transcription reagents kit.

Carry out the amplification according to the manufacturer instruction.

9. TROUBLESHOOTING

These troubleshooting rules may be helpful in explaining any questions that may arise. False negatives with extraction product:

- Degradation of the nucleic acid contained in the sample. It is necessary to use a new
- sample. Store samples under appropriate conditions.

 Loss of nucleic acid pellet. Carefully discard the washing solution trying not to disturb the nucleic acid pellet.
- Degradation of the extracted nucleic acid. It is necessary to use DNase- and RNase-free

False positives with extraction product:

- Contamination during sample extraction. It is necessary to open one test tube at a time.
 Avoid spilling the contents of the test tube. Always change tips.
- Contamination of the reagents prepared for the step. It is necessary to repeat the test.
- Contamination of the extraction zone by amplicons. It is necessary to clean surfaces and instruments using aqueous detergents, wash lab coats, and replace test tubes and tips in use. Use different laboratory coats in different areas.

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

10. TRANSPORTATION

RIBO-prep nucleic acid extraction kit should be transported at 2-25 °C for no longer than 5 days

11. STABILITY AND STORAGE

All components of RIBO-prep nucleic acid extraction kit are to be stored at 2-8°C, when not in use. All components of RIBO-prep nucleic acid extraction kit are to be stable until the expiration date on the label. The shelf life of reagents before and after the first use is the same, unless otherwise stated.

12. REFERENCES

- 1. H. Zetzsche, H.-P. Klenk, M.J. Raupach, T. Knebelsberger, B. Gemeinholzer. Comparison of methods and protocols for routine DNA extraction in the DNA Bank Network // DNA Bank Network. URL: https://www.dnabank-network.org. [date of the application: 2020/02/051
- Gubala V. et al. Point of care diagnostics: status and future //Analytical chemistry. 2012. T. 84. №. 2. C. 487-515.
- Boom R. et al. Rapid and simple method for purification of nucleic acids //Journal of clinical microbiology. 1990. T. 28. №. 3. C. 495-503.
- เกาแบบเบเบฐง. เฮฮบ. 1. ∠อ. № 3. 0. 495-3043.
 4. Boom R. et al. Improved silica-guanidiniumthiocyanate DNA isolation procedure based on selective binding of bovine alpha-casein to silica particles //Journal of Clinical Microbiology. 1999. T. 37. № 3. C. 615-619.

13. QUALITY CONTROL

In compliance with Federal Budget Institute of Science "Central Research Institute for Epidemiology" ISO 13485-Certified Total Quality Management System, each lot of RIBO-prep nucleic acid extraction kit has been tested against predetermined specifications to ensure consistent product quality.

Please contact our Authorized representative in the European Community if side effects, undesirable reactions, facts and circumstances that pose a threat to the life and health of citizens and medical workers are identified during the use of the reagent kit.

	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		
		New sections "Working Conditions" and		
	Content	"Transportation" were added		
	Contont	The "Explanation of Symbols" section was renamed to "Key to Symbols Used		
	Ctability and Ctarage	The information about the shelf life of open		
10.12.10	Stability and Storage	reagents was added		
10.12.10	Key to Symbols Used	The explanation of symbols was corrected		
		The color of Solution for Lysis was changed into blue		
		The volume of Washing Solution 3 was changed		
	Content	into 25 ml (for variant 50) The reference «If Solution for Lysis is stored at 2-		
		8 °C, a crystalline precipitate may form» was		
		added		
04.07.11	Cover page, text	The name of Institute was changed to Federal Budget Institute of Science "Central Research		
VV	Cover page, text	Institute for Epidemiology"		
06.09.11	8. Protocol	Procedure of extraction was corrected (heating at		
RT	8.1. RNA/ DNA extraction	65 °C for 5 min was added in Section 8.1, article 4).		
12.07.12		Information about an unpleasant smell of		
BM	General precautions	Solution for Lysis and the necessity to work in a		
31.03.15	General precautions,	biological cabinet was added		
ME	14. Key to symbols used	Information about hazards was corrected		
10.05.17	Through the text	Correction according to the template Information about hazards was rewritten		
PM	 General precautions, Key to symbols used 	according to the Regulation 1272/2008/EC.		
27.03.18	3. Content	The color of the reagent was specified		
PM	Through the text	The text formatting was changed		
09.04.20 KK		The phrase "Not for use in the Russian		
	Footer	Federation" was added		
21.10.20 KK	Footer, 3. Content	The information about variant 50, REF K2-9-Et- 50-CE was deleted		
11.03.21	o. comen	The name, address and contact information for		
VA	_	Authorized representative in the European		
	Interfering substances	Community was changed		
	and limitations of using	The sections were added		
12.08.21 EM	test material samples Principle of nucleic acid			
LIVI	extraction			
	Through the text	Corrections according to the template		
	1. Intended use	"Indications and contra-indications for use of the reagent kit" subsection was added		
31.05.22	General precautions	The phrase "for single use" was deleted		
KK	2. Contra productions	The Authorized representative in the European		
	13. Quality control	Community was specified for the contact in case		
—	A loke ode d	of undesirable effects when using the reagent kit The intended use was specified. The list of		
	1. Intended use	biological material was expanded.		
18.07.24 rec HM 5.4	3. Content	Information about the number of extraction reactions while working with nail plates and hair		
	J. COMMON	follicles was added		
	4. Additional	The list of additional requirements was specified.		
	requirements	Information about hazards and precautionary		
	5. General precautions	statements list for Solution for lysis was rewritten		
	6. Sampling and	according to the Regulation (EU) 2020/878 List of potentially interfering substances was		
	handling	expanded		
	8. Protocol	Working procedure was rewritten		
-	12. References Through the text	References were renewed Corrections according to the template		
17.02.25		Information about hazards for Washing Solution		
HM	5. General precautions	4 was corrected		

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