SHENTEK

Residual Host Cell RNA Sample Preparation Kit User Guide

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Version: A/1

For Research Use Only Product No.: 1201205

Reagents for 100 Extractions

Huzhou Shenke Biotechnology Co., Ltd

(IMPORTANT: Please read this document carefully before experiment.)

1. Product information

■ Product description

SHENTEK® Residual Host Cell RNA Sample Preparation Kit utilizes Magnetic Particle-based separation technique for RNA extraction from biological products. Extracted RNA can be used in various downstream RNA quantitation applications (*E. coli*, 293T etc.). This kit is compatible with automated extraction using our rHCDpurify instrument (recommended) as well as a sophisticated manual sample preparation procedure.

■ Kit contents and storage

WARNING: Please read the Material Safety Data Sheets (MSDSs) and follow the

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	Binding solution	NND016	20 mL × 1 bottle		
I	Wash buffer A	NND014	30 mL × 1 bottle	room temperature	
	Elution solution	NND061	10 mL × 1 bottle	÷	
	Dilution solution	NND062	12 mL × 2 bottles		
П	Magnetic particles	NND030	750 μ L × 2 tube	2 000	
	5M NaCl	NND040	$500 \mu L \times 2 \text{ tubes}$	2-8°C	
III	Proteinase K	NND023	$500 \mu L \times 2 \text{ tubes}$	-20°C	
	RNase inhibitor	NND060	50 μL × 1 tube		

The kit components can be stored at the appropriate conditions for up to 24 months. Please check the expiration date on the labels.

■ Required materials not included in the kit

- Anhydrous Ethanol (Analytical Reagent, AR)
- ➤ 100% Isopropanol (AR)
- ➤ 1 M HCl & 1 M NaOH (optional)
- Low retention filter tips: 1000 μL, 100 μL, 10 μL
- Nonstick, RNase-free microcentrifuge tubes 1.5 mL

■ Related equipment

- Benchtop microcentrifuge
- Magnetic Stand or rHCD purify instrument
- Vortex mixer
- Dry bath incubator

Micropipettes, 1000 μL, 100 μL, 10 μ

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Before first use of the kit:

- Add 40 mL of ethanol to Wash buffer A.
- Prepare a 70% ethanol buffer with Elution solution in a clean tube, label as Wash Buffer B.
- Store Wash buffer A & B at room temperature properly to prevent evaporation by expiration date.

Before each use of the kit:

- ➤ Prepare 100% isopropanol.
- ➤ Set the dry bath temperatures to 55°C, 50°C or 37°C.

 Note: If the reagent is cloudy or contains precipitates, heat at 37°C until it clears.
- ➤ Proteinase K digestion solution preparation, please refer to Table 2.

Protein concentration (mg/mL) in the sample	Proteinase K volume (μL/sample)	Proteinase K Buffer volume (µL/sample)
0-100	10	100
100-200	20	100

Table 2. Proteinase K digestion solution preparation

Note:

- (1) Ignore this step if the test sample does not contain any protein (such as plasmid DNA),
- (2) Prepare the total volume of Proteinase K digestion buffer appropriately more than necessary for experiment to compensate for pipetting loss.
- (3) If Proteinase K buffer is cloudy or contains precipitates, heat at 37°C until it

clears, and mix well.

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Samples preparation

Sample dilution

Test samples may contain high levels of RNA that are above the upper limit of quantification for the residual RNA assay. Please dilute the samples with Dilution Solution before RNA extraction.

Note: You may also dilute samples with a post-extraction working elution solution.

Sample reconstitution

If the sample is a dry powder, it needs to be dissolved with the Dilution Solution or other buffer to reach a high-concentration sample solution, further dilution can be performed with the Dilution Solution before the subsequent steps.

> pH of test samples

The ideal sample pH is neutral, if the sample pH < 5 or > 9, it may affect the purity of extracted RNA. Therefore, test the sample pH and adjust the pH to neutral (pH 6.0-8.0) with 1 M hydrochloric acid or 1M sodium hydroxide before sample preparation.

➤ Negative control sample (NCS)

The NCS is necessary in this assay that are treated along the entire flow with unknown test sample. The NCS serves as a blank in each experiment to evaluate whether there is cross contamination or environmental contamination during sample handling.

Note: If the sample is diluted, please use the same buffer for negative controls.

Sample extraction recovery control (ERC)
 Sample extraction recovery control (ERC) is used to evaluate the efficiency,

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Sample digestion

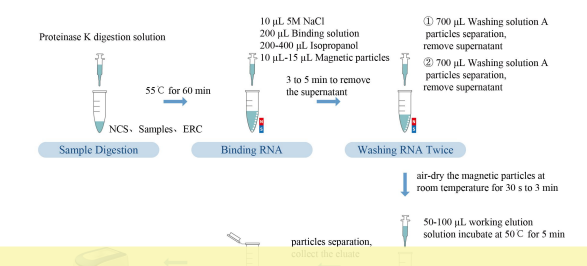
Note: If it is known that the test sample does not contain protein (such as plasmid DNA), please ignore this step.

For each sample tube:

- 1. Add 100 μ L sample and 110 μ L or other volumes Proteinase K digestion solution, then vortex and centrifuge briefly in a microcentrifuge.
- 2. Incubate at 55°C for 60 min in dry bath incubator.
- 3. After this step, machine extraction or manual extraction can be selected following individual step blow. For machine extraction procedure, a maximum digestion volume of 300 μ L can be managed in rHCDpurify system.

■ RNA extraction (Manual)

1. The sample contains protein



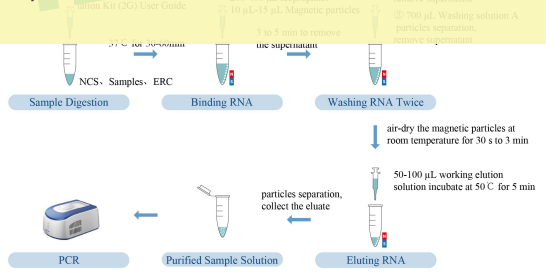
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Note: After sample preparation, the following RNA extraction experiment should be processed as soon as possible.

Binding RNA

For each sample tube:

1. Add 10 μ L 5 M NaCl and 200 μ L Binding solution, vortex and brief spin for 10 seconds (s) in a microcentrifuge.

2. Add 200 μL to 400 μL isopropanol, vortex and spin for 10 s. *Note:*

The volume of isopropanol is approximately 1/3 to 1/2 of the total liquid volume.

3. Add 10 μ L-15 μ L Magnetic particles, vortex the tubes vertically at medium speed for 5 minutes to bind the nucleic acids. Spin for 10 s and place the tubes on the magnetic stand.

(1) Refore use, vortex the Magnetic particles for 5 seconds until

the particles are completely re-suspended Magnetic particles

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Note

- (1) Normally, the particles separation takes 3-5 minutes.
- (2) Do not disturb the Magnetic particles when aspirating the supernatant. During extraction, always place the magnetic bead pellet toward the magnet stand.

Washing RNA

For each tube of Magnetic particles (bind with nucleic acids):

- 1. Add 700 μ L of Wash buffer A, vortex for 10 s to mix well, then spin for 10 s and place the tubes on the magnetic stand. Wait until the solution is clear and the particles are completely separated. Discard the supernatant without disturbing the Magnetic particles.
- 2. Add $700 \,\mu\text{L}$ of wash buffer B, vortex for $40 \,\text{s}$ to mix well, then spin for $10 \,\text{s}$ and place the tubes in the magnetic stand. Wait until

the solution is clear and the particles are completely separated. Discard the supernatant without disturbing the Magnetic particles.

3. To remove the supernatant completely, spin for 10 s in a microcentrifuge and place the tubes in the magnetic stand. Wait until the particles are completely separated, carefully use a 10 μ L volume pipette to remove the remaining liquid.

Note: When removing the supernatant, avoid removing the Magnetic particles together with the supernatant.

4. With the cap open, air-dry the magnetic particles at room temperature for 30 seconds to 3 min to remove any residual ethanol.

Note: The drying time depends on the environmental conditions. It could be shorter in higher temperature or low humidity condition,

while slightly longer in lower temperature or high humidity

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during incubation to ensure complete resuspension of the

Magnetic particles.

Note:

- (1) Vortex the mixture of Magnetic particles and eluent, then swing to the bottom of the tube.
- (2) If the mixture splashed to the cap, centrifuge the tube and mix well by vortex.
- 2. After incubation, Quickly spin the eluate tube for 1 min, then place the tubes on the magnetic stand. Wait until the particles are completely separated, carefully transfer the eluate to a clean microcentrifuge tube.
- 3. Quickly spin the eluate tube for 10 s and then place it on the magnetic stand. Wait until the particles are completely separated, carefully transfer the eluate to a nonstick 1.5 mL microcentrifuge

tube and label the corresponding sample.

Note: Note: Transfer the eluate completely and avoid leaving any residuals behind.

Precautions

- 1. During washing and eluting RNA, centrifugation should be performed immediately after vortex to ensure no magnetic particles or liquid left on the tube caps or walls.
- 2. When open the centrifuge tube, hold the tube in one hand, gently open the cap with your thumb without splashing.
- 3. Place the tubes in the magnetic stand with the pellet against the magnet, and rotate the tubes slowly during the process to accelerate the Magnetic particles aggregation.
- 4. Do not over dry the Magnetic particles when removing the

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5. Please perform the subsequent asset on the subsequent

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Table 3. 96 deep well plate layout

Group 1					Group 2						
1	2	3	4	5	6	7	8	9	10	11	12
S1											
S2						S1 -ERC					
S3						S2 -ERC					
S4						S3 -ERC					
S5						S4 -ERC					
S6						S5 -ERC					
						S6 -ERC					
NCS											

> Column 1 or 7: Binding solution 200 μL/well, 10 μL 5 M NaCl, isopropanol

 $200~\mu L$ to $400~\mu L/well$ and all samples after digestion

- Column 2 or 8: Washing buffer A 700 μL/well
- Column 3 or 9: Washing buffer B 700 μL/well
- > Column 4 or 10: Magnetic particles 15 μL/well
- Column 5 or 11: working elution solution 100 μL/well *Note*:

The volume of isopropanol is approximately 1/3 to 1/2 of the total liquid volume.

Program setting

- a. Power button on → click "login" to enter account and password → enter the main page
- b. Wipe the interior of the instrument with a 75% ethanol→click on "UV light"→

Note: This step can be set before the extraction preparation operation.

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deep well plate and transfer all the purified sample solution to the corresponding new 1.5 mL tube.

Important points to note:

- a. Before starting the program, it is compulsary to add plastic sleeve.
- b. UV sterilization is required for at least 15 minutes before and after instrument operation. The interval of two extractions will need more than 30 minutes.
- c. After the program is completed, the sample eluente needs to be transferred immediately to a clean 1.5 mL tube.
- d. Please try to conduct subsequent testing on the same day of sample RNA extraction to ensure an accurate test result.

Troubleshooting

Problem	Possible cause	Suggested Solution
Problem	Possible cause	Suggested Solution
	Ethanol not added in	Add ethanol to Wash buffer A
	Wash buffer A	according to the user manual
	Magnetic pellet over	The drying time depends on the lab
	drying	environment. Shorten or extend the
		incubation time according to above
		description.
	Magnetic particles are	Vortex the microcentrifuge tube with
	attached too tightly to	the eluate, until the Magnetic
	the tube walls during	particles fall off the tube walls and
	the elution.	suspended in the eluate; if the
		Magnetic particles are still attached
		to the tube walls, incubate the
Low recovery		microcentrifuge tube at 50°C for 2
of nucleic		min, then vortex until the Magnetic

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		stand, resuspend the Magnetic
		particles by pipetting gently at the
		tube bottom until the particles are
		attached to the walls.
	Store Magnetic	Store the Magnetic particles at
	particles -20°C caused	2-8°C.
	the performance drop	
	Inaccurate spiking or	Use low retention filter tips and
Unstable	aspiration	calibrate the pipette regularly to
recovery rate		guarantee precise measurement.
	Residual Magnetic	Centrifuge again and place it in the
	particles left in the	magnetic stand, then carefully
	sample after elution	transfer the eluate to a clean 1.5 mL
		tube.

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Huzhou Shenke Biotechnology Co., Ltd. www.shentekbio.com

Address: 8th Floor, 6B Building, No.1366 Hongfeng Road, Huzhou313000, Zhejiang Province, China

E-mail: info@shentekbio.com

Phone: +1 (908) 822-3199 / (+86) 400-878-2189