

# NucleoGene QuickEx Total RNA Extraction Kit (For Tissue and Cells) Instructions for Use

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REF

For isolation and purification of Total RNA from Cell and Tissue samples.  
For research use only.  
Not suitable for diagnostic use.  
For professional use only.

NGE024— 50 Test  
100 Test  
250 Test

## Kit Contents

	Supplied Material	50 Test	100 Test	250 Test
1.	Lysis Buffer †	50 ml	100 ml	250 ml
2.	Binding Buffer*	7 ml	15 ml	35 ml
3.	Wash Buffer I*	13 ml	26 ml	65 ml
4.	Wash Buffer II*	5 ml	10 ml	25 ml
5.	DNase I*	1 vial	1 vial	1 vial
6.	DNase I Buffer	2.25 ml	4.5 ml	11.25 ml
7.	Elution Buffer	5 ml	10 ml	25 ml
8.	Spin Column	50	100	250
9.	Collection Tubes	50	100	250
10.	User's Guide	1	1	1

\*Look at "Notes Before Starting" for preparation DNase and Wash Buffer.

†Precipitation may occur during storage. Heat to dissolve the precipitate formed before use.  
Lysis Buffer should be stored at +4° C away from light.

## Storage Conditions & Robustness

Kit components should be stored at room temperature (15-25°C). Do not expose Kit to direct sunlight. When used under these conditions, the kit can be stored for 12 months without any performance loss.

## Intended Use

NucleoGene QuickEx Total RNA Extraction Kit includes spin column and special lysis buffer system for effectively Total RNA(including mirna) extraction in 8 minutes from blood(serum, plasma, whole blood), cell and tissue samples. Specifically designed for Total RNA isolation and purification. Suitable for downstream applications such as purified Total RNA, RT-PCR, northern blotting, primer extension, mRNA selection, cDNA synthesis. Real Time qualitative and quantitative RT-qPCR and others application. NucleoGene QuickEx Total RNA Extraction Kit is designed for research use only. It is not for diagnosis and treatment of disease.

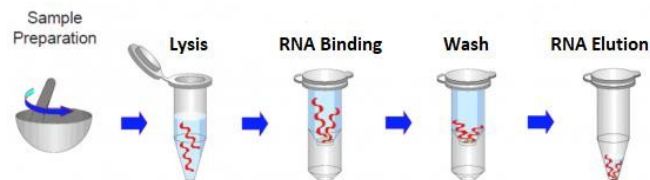
## Product Usage Limits

- NucleoGene QuickEx Total RNA Extraction Kit is for purification of Total RNA of blood, cell and tissue samples. It cannot be used for clinical purposes. Whether the kit is suitable for the user's own experimental design is up to the user's discretion.
- In particular, the ability of this kit to isolate Total RNA of blood, cell and tissue samples has been confirmed.
- The collection, transport and storage of the sample to be used are at least important as purification and are sensitive stages that may affect the results.
- It is designed for professional use by educated personnel only.
- Kits or any kit components with different lot numbers should not be used together.

## Introduction

NucleoGene QuickEx Total RNA Extraction Kit provides a quick and easy spin column system for purifying and enriching RNAs(Total RNAs) and others small cellular RNAs from a wide variety of blood, tissue and cells. Since Total RNAs are vital for regulating gene expression, this kit is opti-

mized for isolation of Total RNA molecules while removing minimizing genomic DNA contamination for improved sensitive downstream applications. NucleoGene QuickEx Total RNA Extraction Kit is improved for cell RNA extraction effectively from blood, cell and tissue samples to be a quick and useful method. Working principle of Kit depends on binding of RNA with its especially improved solutions to spin columns. The superior performance of the kit is due to its ability to lyse the cell walls of blood, cell and tissue samples, as well as its ability to remove PCR inhibitors present in the sample with solutions specially improved for, cell and tissue samples. While wash buffer optimized specifically is improved for purified nucleic acids especially from protein and other inhibitors, the Lysis Buffer solution is improved for lysing the cell wall and remove the inhibitors from the sample. With the complete the isolation process, Total RNA is obtained at high efficiency and purity.



Purified RNA with completed isolation should be used immediately or a freezer at -80 ° C should be preferred for long-term storage.

## Warnings & Precautions

- All clinical specimens and residues and debris arising therefrom must be treated as potential infectious agents and disposed of accordingly.
- All samples should be prepared in Biosafety Level 1 and 2 or Class II Biosafety Cabinets.
- All surfaces should be cleaned daily with a disposable paper towel or napkin by using freshly prepared Sodium Hypochlorite %20 Solution.
- Do not neglect to use laboratory safety consumable materials such as disposable gloves, goggles, visor, disposable cuffs, and disposable mask.
- If any of the kit components come into contact with your skin, wash it with plenty of water immediately. In case any of the kit components come into contact with your mucous membranes such as your eyes or mouth, wash the contact part with plenty of water again; and go to the doctor. If possible, prefer filtered pipette tips.
- Do not mix the kit solutions or waste appeared during isolation with bleach.
- Keep Kit out of contamination sources such as DNA and RNA, especially amplified nucleic acid.
- Do not mix solutions with different lot numbers or do not use or combine products from other companies.
- For more information, please apply the Material Safety Data Sheet(MSDS), which you can request from [www.nucleogene.com](http://www.nucleogene.com).

## Other Required Materials

- Ethanol (%96-100)
- Chloroform
- Microcentrifuge
- Vortex mixer
- Microcentrifuge tubes (1.5 ml)
- Microcentrifuge tubes (2.0 ml)
- Micropipette set and sterile filtered micropipette tips
- Spatula

## Notes Before Starting

If the kit is kept in a non air conditioned environment, Lysis buffer solution can precipitate at room temperature in cold weather. In such a case to dissolve the precipitate, the Lysis buffer solution is held at 50° C in a water bath until dissolving completely. Be sure of all solutions are at the room temperature before and after protocol implementation, if they are not, allow them to reach room temperature.



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DNase I is shipped at room temperature and should be stored at -20°C for extended periods after receiving the kit. Add absolute ethanol to Wash Buffer then mix by shaking for a few seconds. Check the box on the bottle. Be sure and close the bottle tightly after each use to avoid ethanol evaporation. Prepare Phosphate Buffered Saline (PBS, pH7.2) and 0.10-0.25% Trypsin for cultured cells. Yield and quality of RNA will be higher when fresh samples or samples which have been flash frozen and stored at -80° C are used.

#### Preparation of Solutions;

**Binding Buffer:** Add 28 ml of 96% or absolute ethanol for 50 tests  
Add 55 ml of 96% or absolute ethanol for 100 tests  
Add 140 ml of 96% or absolute ethanol for 250 tests

**Wash Buffer I:** Add 13 ml of 96 % or absolute ethanol for 50 tests  
Add 26 ml of 96 % or absolute ethanol for 100 tests  
Add 65 ml of 96 % or absolute ethanol per bottle for 250 tests

**Wash Buffer II:** Add 20 ml of 96 % or absolute ethanol for 50 tests  
Add 40 ml of 96 % or absolute ethanol for 100 tests  
Add 100 ml of 96 % or absolute ethanol per bottle for 250 tests

**DNase I:** Add 250 µl of Nuclease Free Water for 50 tests  
Add 500 µl of Nuclease Free Water for 100 tests  
Add 1250 µl of Nuclease Free Water for 250 tests

#### Protocol

##### Control the Reagents

Before starting to Total RNA isolation, control reagents that are free of leakage. To mix the solutions, shake the reagents slightly. If reagents precipitate (specially Lysis solution), they dissolve at 50 ° C by heating.

##### Sample Preparation

##### Tissue(Tough-to-lyse samples)

Tough-to-lyse samples (see table below) can be homogenized in ≥ 800 µl Lysis Buffer with a mortar/pestle, syringe, tissue grinder, or bead beating with a high-speed homogenizer. To remove particulate debris from homogenized tissue, centrifuge and transfer the supernatant into a new nuclease-free tube.

Tissue	Gram(+) bacteria	Pathogen (microbes in tissue)
animal: high yield (≤ 100 mg) animal: low yield (≤ 100 mg) plant (≤ 100 mg)	bacteria (≤ 1x10 <sup>7</sup> ) yeast (≤ 1x10 <sup>7</sup> )	animal/insect, plant (≤ 25 mg)

#### Cell

**Lyse animal or gram(-) bacteria cells\* directly in a culture dish\*\* or resuspend pelleted cells in an appropriate volume (see table below) of Lysis Buffer and mix thoroughly.**

**\* For cell suspensions, add 3 volumes of Lysis Buffer to 1 volume of cell suspension.**

**\*\* For direct lysis in a dish, add 100 µl for each cm<sup>2</sup> of culture surface area.**

##### \*Adherent Cultured Animal Cells

- Remove the culture medium and wash cells in PBS.
- Aspirate PBS then add 0.10-0.25% Trypsin in PBS.
- Once cells have detached, add the medium and transfer to a 15 ml centrifuge tube.
- Proceed with Suspension Cultured Animal Cells.

##### \*Suspension Cultured Animal Cells

- Transfer cells (up to 1 x 10<sup>6</sup>) to a 2 ml microcentrifuge tube.
- Harvest by centrifugation for 5 minutes at 300 x g.
- Carefully remove the supernatant completely by aspiration then proceed to Lysis.

##### \*Bacterial Cells

- Transfer 2 ml bacterial culture (up to 1 x 10<sup>6</sup>) to a 2 ml microcentrifuge tube.
- Harvest by centrifugation for 5 minutes at 14000 x g.
- Carefully remove the supernatant completely by aspiration then proceed to Lysis.

##### \*Yeast Cells

- Transfer 2 ml yeast culture (up to 1 x 10<sup>6</sup>) to a 2 ml microcentrifuge tube.
- Harvest by centrifugation for 5 minutes at 14000 x g.
- Carefully remove the supernatant completely by aspiration then proceed to Lysis.



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## RNA EXTRACTION

1. Ensure that the sample and Lysis Buffer (1 ml for 100 mg Tissue or  $1 \times 10^6$  cells) are fully mixed and 5 minutes incubation at room temperature, later centrifuge for 10 minutes at 12,000 xg +4°C. Carefully withdraw the supernatant and transfer it to a new (1.5-2 ml) microcentrifuge tube.

2. Add 200 µl of Chloroform per 1 ml Lysis Buffer and mix well, then incubate for 2 minutes at room temperature and centrifuge at 12,000 xg for 15 minutes at in a refrigerated centrifuge set at +4 C. 3 phases will form in the tube, transfer the entire clear phase at the top to a new 1.5-2 ml microcentrifuge tube that does not contain nuclease.

3. Add an equal volume binding buffer (eg transferred supernatant 250 ul+250 ul binding buffer) and mix thoroughly by vortex.

4. Transfer up the 700 µl of the mixture into a Spin Column in a Collection Tube and centrifuge at 11,000 xg for 30 sec. Discard the pass-through and place the spin column back in the same collection tube. (Repeat this step using the remainder of the sample, if it is necessary.)

### DNase I treatment (optional)

\* In an RNase-free tube (not included), add 5 µl DNase I (6 U/l), 45 µl DNase I Buffer and mix by gentle inversion. Add the mix directly to the column matrix. Incubate at room temperature (20-30°C) for 15 minutes. Proceed to step 5.

5. Add 500 µl of Wash I Buffer to the Spin Column and centrifuge at 11,000 xg for 30 seconds. Discard the pass-through and reinsert the column back into same collection tube.

6. Add 500 µl of Wash II Buffer to the Spin Column and centrifuge at 14,000 xg for 1 min. Discard the pass-through and reinsert the column back into same collection tube.

7. Centrifuge at  $\geq 10,000$  xg for an additional 1 min at room temperature to remove residual wash buffer. Transfer the column to a new 1.5 ml microcentrifuge tube.

8. To elute the RNA, add 100-50 µl of Elution Buffer directly to the center of the spin column and centrifuge at 11,000 xg for 2 min.

**\*Alternatively, for highly concentrated RNA use  $\geq 30$  µl elution. The eluted RNA can be used immediately or stored frozen (-80 °C)**

## Troubleshooting

<b>Precipitation, viscous lysate</b>	<b>Incomplete lysis and/or high-mass input:</b> <ul style="list-style-type: none"> <li>- If precipitation occurs (upon adding ethanol to the lysate) or if the lysate is extremely viscous, increase the volume of Lysis Buffer to ensure complete lysis and homogenization until lysate is transparent.</li> </ul>
<b>Low purity (A260/A230 nm, A260/A280 nm)</b>	<b>Sample handling:</b> <ul style="list-style-type: none"> <li>- Ethanol and/or salt contamination. After centrifugation steps, carefully remove the column from the collection tube to prevent buffer carryover. Alternatively, blot emptied collection tube with a tissue or towel. Make sure lysate and wash buffers have passed completely through the matrix of the column. This may require centrifuging at a higher speed and/or longer time.</li> </ul> <b>Incomplete lysis and/or cellular debris:</b> <p>Increase the volume of Lysis Buffer to ensure complete lysis and homogenization. Be sure to centrifuge any cellular debris and then process the cleared lysate.</p>
<b>Low yield</b>	<b>Sample input:</b> <ul style="list-style-type: none"> <li>- Too much input or incomplete lysis/homogenization can cause cellular debris to clog or overload the column and result in compromised RNA recovery. Use less input material and/or increase Lysis Buffer.</li> </ul> <p>High-protein content (blood, plasma/serum, etc.)</p> <ul style="list-style-type: none"> <li>- Perform Proteinase K treatment to the sample prior to adding Lysis Buffer</li> </ul>
<b>DNA contamination</b>	<b>To remove DNA:</b> <ul style="list-style-type: none"> <li>- Perform in-column DNase I treatment or perform DNase I treatment post-purification then repurify the treated sample.</li> <li>- For future preps, increase the volume of Lysis Buffer to ensure complete lysis and homogenization of the sample.</li> </ul>
<b>RNA degradation</b>	<b>To prevent RNA degradation:</b> <ul style="list-style-type: none"> <li>- Immediately collect and lyse fresh sample into Lysis Buffer to ensure RNA stability. Homogenized samples in Lysis Buffer can be stored frozen for later processing.</li> </ul>

