

Quick Start Guide

GenElute™ 96 well Total RNA Purification Kit

RTN9604

Reagents to Prepare

- Add 360 and 150 mL of 96-100% ethanol to each bottle of RNA Wash Buffer 2 and RNA Wash Buffer 3, respectively. After each use, tightly cap to prevent ethanol evaporation.
- Reconstitute RNase-free rDNase by adding 540 μ L of RNase-free Water and incubate for 1 min at room temperature. Gently mix to ensure complete dissolution. If not using the entire 96 well plate, aliquot reconstituted rDNase solution and store at -20 °C.
- For each sample to be processed mix 10 μ L of reconstituted rDNase with 90 μ L of Reaction Buffer for rDNase.

Protocol

All centrifugations carried out at 5,600-6,000 x g.

Harvest and Lyse Pelleted Cells

1. Pellet cells by spinning for 5 min at 500 x g and remove supernatant. Add 300 μ L RNA Lysis Buffer and 3 μ L β -mercaptoethanol (2-ME) to each sample and mix thoroughly. Continue to **Sample Loading**.

Harvest and Lyse Attached Cells

1. Aspirate culture medium and add 130 μ L RNA Lysis Buffer and 1.3 μ L β -mercaptoethanol (2-ME) to each well of cell culture plate and mix thoroughly. Continue to **Sample Loading**.

Harvest and Lyse Tissue

1. Quickly slice and weigh tissue. Transfer to appropriate vessel for homogenization. Add 300 μ L RNA Lysis Solution and 3 μ L β -mercaptoethanol (2-ME) and homogenize until no visible pieces remain. Continue to **Sample Loading**.

Sample Loading

2. Add an equal volume (either 300 μ L or 130 μ L) of RNA Wash Buffer 3 and pipet up and down at least 10-15 times.
3. Place RNA Binding Plate on Square-well Block and transfer lysates to the RNA Binding Plate. Spin for 2 min and empty Square-well Block.

Bind DNA

4. Add 500 μ L RNA Wash Buffer 2 to each well. Place Binding Plate on top of Square-well Block, spin for 2 min, and empty Square-well Block.
5. Carefully add 95 μ L of the prepared rDNase directly to entire membrane and incubate for 15 min at room temperature.

Wash to remove contaminants

6. Add 500 μ L RNA Wash Buffer 1. Place Binding Plate on top of Square-well Block, spin for 2 min, and empty Square-well Block.
7. Add 800 μ L RNA Wash Buffer 2. Place Binding Plate on top of Square-well Block, spin for 2 min, and empty Square-well Block.
8. Add 500 μ L RNA Wash Buffer 3. Place Binding Plate on top of Square-well Block, spin for 2 min, and empty Square-well Block.
9. Place Binding Plate on top of Square-well Block and spin for an additional 10 min to eliminate any trace ethanol. Empty Square-well Block.

Elute purified DNA

10. Place Binding Plate on top of Round-well Block and add 75 μ L RNase-free water to the bottom of each well. Incubate at room temperature for 2 min and spin for 3 min to elute RNA.

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