Technical Bulletin

Total RNA and Protein Extraction Kit (Cat. VRP-01)

Description:

- This kit is designed for rapidly isolating the highest yields and quality of total RNA and protein from cells/tissue samples, offering a simple, fast, environmental-friendly protocol for total RNA/protein extractions without using vacuum filtration and toxic organic solvents such as xylene, phenol or chloroform. This kit is specially designed for preparing the high quality of RNA/protein from small amounts of sample material for RT-PCR, qPCR, microarrays, western blots and other RNA/protein assays. The kit is compatible with cells/tissues from LCM samples.
- Key features:
 - 1. Obtain the highest yield and integrity of total RNA and total protein within 1 hour.
 - 2. No filter column or vacuum filtration is required, able to avoid the loss of RNA during extracting.
 - 3. Suitable for extracting the pure total RNA/protein from the small/tiny tissues or LCM samples.

Kit components and protocol: (keep all samples and buffers on ice during proceeding)

Quantity (50 extractions)	Storage	
10.0 mL	-20°C	
1.25 mL	-20°C	
10.0 mL	-20°C	
1.25 mL	-20°C	
	Quantity (50 extractions) 10.0 mL 1.25 mL 10.0 mL 1.25 mL	Quantity (50 extractions) Storage 10.0 mL -20°C 1.25 mL -20°C 10.0 mL -20°C 1.25 mL -20°C 1.25 mL -20°C 1.25 mL -20°C

- Add 200ul ice-cold <u>one-step fast lysis buffer-A</u> and 25ul <u>one-step I-Lysis buffer-B</u> into a clean 1.5ml mcrocentrifuge tube containing 50mg-200mg frozen/ fresh tissues or 1x10e7 cells. Incubate on ice for 15 minutes. Vortex vigorously at highest speed for 20 seconds every 5 minutes.
- **2. (Optional)** Using a clean plastic pestle to homogenize the tissues for 10-20 strokes to completely disrupt the tissues and may obtain higher yields of total RNA and proteins.
- Centrifuge at highest speed (\$\sigma 13,000 xg\$) for 5 minutes at 4°C and transfer all the supernatant into a clean new 1.5mL tube. Discard the pellet.

Note, Discard the lipoproteins/lipids that may form an upper layer after centrifugation.

- **4.** Add 200ul **<u>RNA Precipitation Solution (2x)</u>** into the supernatant, mix thoroughly by pipette up and down several times.
- Centrifuge at 5,000 xg for 2 minutes at 4°C and transfer the supernatant into a clean new 1.5mL tube and incubate at -20°C for 20 minutes or overnight. Save the pellet tube on ice and label as total protein.
- 6. Centrifuge the supernatant at highest speed (∽13,000 x g) for 10 minutes at 4°C. <u>The pellet is</u> the total RNA, and the supernatant contains protein fragments. Keep the pellet (RNA) on ice or continue the step 8 for RNA cleanup.
- 7. Transfer the supernatant into the <u>total protein</u> tube from step 5, and add 25ul <u>protein extraction buffer-C</u>, incubate on ice for 10 minutes, vortex vigorously at highest speed for 20 seconds every 5 minutes. This suspension contained the total protein is ready for most protein assays: 1D and 2D electrophoresis and Western blotting.

Note, centrifuge at highest speed (>13,000 xg) for 2 minute at 4°C prior to measure the protein concentration using a spectrometer or BCA assays. Store the protein solution at -80°C.

Technical Bulletin

- 8. Simply rinse the RNA pellet with 500ul ice cold 80% ethanol for 2 times without resuspending the RNA pellet. Centrifuge at highest speed (∽13,000 xg) for 10 minute at 4°C if the pellets are resuspended. Air-dry the RNA pellet for 5-10 minutes.
- **9.** Dissolve the RNA pellets in 20ul nuclease-free H₂O or TE. Centrifuge at highest speed (\cdots 13,000 xg) for 2 minute at 4°C prior to measure the RNA concentration with 260/280nm.
- 10. Store all the RNA/protein extracts at -80°C.





Precautions and Disclaimer:

This product and procedure described are intended for R&D use only. Purchase of this product does not convey a license to perform any patented process.