RNA Pure Kit

For research use only

 Catalogue Numbers
 Quantity

 IB47640
 4 rxns

 IB47641
 50 rxns

 IB47642
 100 rxns



Introduction

The RNA Pure Kit uses a simple and efficient spin column procedure to purify Total RNA stored in RNase-free water, elution buffer or TE Buffer following extraction using acid-guanidinium-phenol-chlorofom based methods such as TRIzol® Reagent and IBI Isolate. Contaminants such as RNases, DNA and residual phenol are effectively removed using a simple 4 step procedure. The high-quality, total RNA is eluted in RNase-free Water or TE (RNase-free) and is ready for use in a variety of sensitive downstream applications.

Quality Control

The RNA Pure Kit is tested on a lot-to-lot basis. Following RNA purification using the RNA Pure Kit, 10 μ l from a 50 μ l eluate of purified RNA is analyzed by electrophoresis on a 0.8% agarose gel.

Advantages

- Purify up to 50 µg of total RNA within 10 minutes
- Recovery: up to 80% of high quality RNA (A260/A280 = 1.9-2.0)
- Elution volume: 20-50 µl
- Compatibility: purify RNA stored in RNase-free water, elution buffer or TE Buffer following extraction using IBI Isolate, TRI-Reagent®, TRIzol®, RNAzol® and QIAzol® etc.

Applications

RT-PCR, Northern Blotting, Primer Extension, mRNA Selection, cDNA Synthesis, RNase Protection Assay

Caution

During operation, always wear a lab coat, disposable gloves, protective goggles and (anti-fog) procedure mask. Disposable/non-disposable glassware, plasticware and automatic pipettes should be sterile (RNase-free) and used only for RNA procedures.

Additional Requirements

absolute ethanol, 1.5 ml microcentrifuge tubes (RNase-free)

Components and Storage

Item	Volume	Product	Shipping	Storage
RNA Pure Buffer	3 ml	PR004	room temperature	dry at room temperature
	30 ml	PR050		
	60 ml	PR100		
Wash Buffer ¹ (Add Ethanol)	1 ml (4 ml)	PR004	room temperature	dry at room temperature
	12.5 ml (50 ml)	PR050		
	25 ml (100 ml)	PR100		
RNase-free Water	1 ml	PR004	room temperature	dry at room temperature
	6 ml	PR050		
	6 ml	PR100		
PR Columns	4 pcs	PR004	room temperature	dry at room temperature
	50 pcs	PR050		
	100 pcs	PR100		
2 ml Collection Tubes	4 pcs	PR004	room temperature dry at room	dry at room temperature
	50 pcs	PR050		
	100 pcs	PR100		

¹Add absolute ethanol (see the bottle label for volume) to Wash Buffer prior to initial use

RNA Purification Protocol Procedure

Please read the entire instruction manual prior to starting the Protocol Procedure.

1. Sample Preparation

- 1. Transfer up to 100 µl of RNA product (stored in RNase-free water, elution buffer, TE buffer) to a 1.5 microcentrifuge tube (RNase-free).
- 2. Add 5 volumes of RNA Pure Buffer to 1 volume of the sample then shake vigorously.

2. RNA Binding

- 1. Add an equal volume of 70% ethanol (if the sample mixture is 600 µl, add 600 µl of 70% ethanol) to the sample mixture from step 1.
- 2. Shake the mixture vigorously and break up any precipitate with a pipette.
- 3. Place a PR Column in a 2 ml Collection Tube then transfer 500 µl of the ethanol-added mixture to the PR Column.
- 4. Centrifuge at 14-16,000 x g for 1 minute then discard the flow-through and transfer the remaining mixture to the same PR Column.
- 5. Centrifuge at 14-16,000 x g for 1 minute then discard the flow-through and place the **PR Column** back in the 2 ml Collection Tube.

3. RNA Wash

- 1. Add 600 µl of Wash Buffer (make sure ethanol was added) to the CENTER of the PR Column.
- 2. Centrifuge at 14-16,000 x g for 30 seconds then discard the flow-through and place the PR Column back in the 2 ml Collection Tube.
- 3. Centrifuge at 14-16,000 x g for 3 minutes to dry the column matrix.

4. RNA Elution

- 1. Place the dried PR Column in a clean 1.5 ml microcentrifuge tube (RNase-free).
- 2. Add 20-50 µl of RNase-free Water or TE (RNase-free) to the CENTER of the column matrix.
- 3. Let stand for 2 minutes or until the RNase-free Water or TE (RNase-free) is absorbed completely by the matrix.
- 4. Centrifuge at 14-16,000 x g for 2 minutes to elute the purified RNA.

Troubleshooting

Problem	Cause	Solution
	A. Incorrect RNA elution	A. Make sure RNase-free Water is added to the
Low Yield		center of the RB Column and is absorbed
		completely.
Degraded RNA	A. Incorrect sample storage temperature	A. Extracted RNA should be stored at -70°C.
Low RNA A260/A280	A. Incomplete wash step	A. Wash the RB Column with ethanol added Wash
		Buffer 2 times.
Eluted RNA does not	A. Residual ethanol contamination	A. Following the wash step, dry the RB Column with
perform well in downstream		additional centrifugation at 14-16,000 x g for 5
applications		minutes or incubate at 60°C for 5 minutes.

Related RNA Extraction Products

RNA Extraction and Purification		
Product	Package Size	Catalogue Number
Total RNA Mini Kit (Blood/Cultured Cell)	50/100/300 preps	IB47321/322/323
Total RNA Mini Kit (Tissue)	50/100/300 preps	IB47301/302/303
Total RNA Mini Kit (Plant)	50/100/300 preps	IB47341/342/343
rBAC Mini RNA Bacteria Kit	100/300 preps	IB47421/422
rYeast Total RNA Mini Kit	100/300 preps	IB47411/412
miRNA Isolation Kit	100 preps	IB47371
IBI Isolate	100/200 rxns	IB47601/602
IBI Tri-Isolate	100/200 rxns	IB47631/632
RNA Pure Kit	50/100 rxns	IB47641/642
Virus DNA/RNA Purification		
Product	Package Size	Catalogue Number
Viral Nucleic Acid Extraction Kit II	50/100/300 preps	IB47401/402/403

For additional product information, please visit www.ibisci.com. Thank you!