Contents

Introduction
Storage and Stability 2
Kit Contents
Before Starting
Mollusc / Arthropod RNA Protocol
DNase digestion Protocol (Optional) 7
Quantization and Storage of RNA
RNA Quality
Troubleshooting Guide
Order Information 11

Introduction

The E.Z.N.A.[®] Mollusc RNA Kit is designed for efficient recovery of total RNA greater than 200 nt from molluscs, arthropods, roundworms, flatworms, and other invertebrate tissue samples rich in mucopolysaccharides. Samples are homogenized and lysed in RNA-Solv Reagent and extracted with chloroform to remove mucopolysaccharides and denature proteins. Following a rapid alcohol precipitation step, binding conditions are adjusted and RNA further purified using HiBind RNA spin columns. In this way salts, proteins and other contaminants are removed to yield high quality total RNA suitable for downstream applications such as reverse transcription, poly (A)+ mRNA selection, and hybridization techniques.

Storage and Stability

All components of the Mollusc RNA Kit are stable for up to 24 months from the date of purchase when stored at 22°C-25°C.For Long time storage, Keep RNA-Solv Reagent at 4°C. During shipment or storage in cool ambient conditions, salts may precipitate in certain buffers. Simply warm to 37°C and mix to dissolve. Contents of the kit should not be refrigerated at any time.

2

Kit Contents

Product	R6875-00	R6875-01	R6875-02
Preps	5	50	200
HiBind [®] RNA Mini column	5	50	200
2 ml Collection tubes	10	100	400
RNA-Solv Reagent	6 ml	60 ml	220 ml
Buffer RB	5 ml	30 ml	100 ml
RNA Wash Buffer I	5 ml	45 ml	175 ml
RNA Wash Buffer II	2 ml	12 ml	45 ml
DEPC water	1 ml	20 ml	50 ml
User Manual	1	1	1

Before Starting

Please take a few minutes to read this booklet thoroughly to become familiar with the protocol. Prepare all materials required before starting to minimize RNA degradation.

	RNA Wash Buffer II Concentrate must be dilute absolute ethanol (96-100%) before use and store a temperature.	
IMPORTANT	R6875-00	Add 8 ml ~96-100% ethanol
	R6875-01	Add 48 ml ~96-100% ethanol
	R6875-02	Add 180 ml ~96-100% ethanol to each bottle

Work with RNA

- Whenever working with RNA, always wear latex gloves to minimize RNase contamination. Change gloves frequently. Use only clean RNase-free disposable plastic pipette tips when using the supplied reagents.
- During the procedure work carefully but quickly.
- Under cool ambient conditions, crystals may form in buffers. This is normal and the bottle may be warmed to redissolve the salt.

E.Z.N.A.™ Mollusc/Arthropod RNA Protocol

Materials to be provided by user:

- Microcentrifuge capable of 12,000 x g
- Nuclease-free microfuge tubes
- Absolute (96%-100%) ethanol
- Isopropyl alcohol (isopropanol)
- Liquid nitrogen for freezing/disrupting samples
- Preheat an aliquot (100 µl per sample) of DEPC-treated water at 65°C.

Note: Use extreme caution when handling liquid nitrogen.

Use only fresh tissue to ensure RNA integrity. Samples preserved in formalin usually yield degraded RNA but may still yield adequate results for RT-PCR of target regions <500 nt.

Arthropods

 Pulverize no more than 50 mg of tissue in liquid nitrogen with mortar and pestle and place the powder in a clean 1.5 ml microcentrifuge tube. If ceramic mortar and pestle are not available, homogenize the sample in the microfuge tube using a disposable microtube pestle (OBI Cat# SSI-1014-39 & SSI-1015-39).Proceed to step 2 below.

Molluscs (and other soft tissued invertebrates)

1. Grind no more than 50 mg tissue in liquid nitrogen with mortar and pestle and place the powder in a clean 1.5 ml microcentrifuge tube. If ceramic mortar and pestle are not available, homogenize the sample in the microfuge tube using a disposable microtube pestle (OBI Cat# SSI-1014-39 & SSI-1015-39). Proceed to step 2 below.

Amount of starting material depends on sample and can be increased if acceptable results are obtained with the suggested 50 mg tissue. For easy to process specimens, the procedure may be scaled up and the volumes of all buffers used increased in proportion. In any event, use no more than 100 mg tissue per HiBind[®] spin-column as RNA binding capacity (100 μ g) may be exceeded. Meanwhile, difficult tissues may require starting with less than 30 mg tissue and doubling all volumes to ensure adequate lysis.

2. Add 1 ml of RNA-Solv Reagent and vortex vigorously to make sure that all

4

6

5

4. **Centrifuge at 12,000 x g for 15 minutes at 4°C.** The mixture separates into a lower phenol-chloroform phase, an interphase, and an upper aqueous phase. RNA remains entirely in the aqueous phase.

clumps are dispersed. RNA cannot be effectively extracted from clumped tissue.

Add 0.2 mL of chloroform per 1 mL of RNA-Solv Reagent. Cap sample tubes

securely and shake vigorously for 15 seconds. Incubate at room temperature for 3

Incubate at room temperature for 2-3 minutes.

3.

minutes.

- 5. Transfer no more than 80% of the aqueous phase to a fresh tube. Add one volume of isopropanol and mix to precipitate RNA. Incubate at room tempature for 10 min and centrifuge 12,000 x g for 3 min at 4°C. Carefully discard as much supernatant as possible without disturbing RNA pellet. Invert the microfuge tube on a paper towel for 1 min to allow residual liquid to drain; it is not necessary to dry the pellet.
- Add 100 μl of sterile DEPC-treated water and vortex to resuspend the pellet. A brief incubation at 65°C may be necessary to effectively dissolve the RNA.

Note: All of the centrifugation step below should be carried out at room temperature.

- Adjust binding condition: Add 350µl Buffer RB/2-mercaptoethanol followed by 250 µl absolute ethanol and vortex to mix. If RB buffer is used in step to dissolve the RNA pellet, add 350µl of 70% ethanol. Vortex to mix. Note: RB buffer recommended at this step to avoid RNA degradation. Add 20 µl 2mercaptoethanol per 1 ml of Buffer RB before use. This mixture can be made and stored at room temperature for 1 week.
- Apply entire mixture, including any precipitation that may have formed, to an HiBind[®] RNA column assembled in a 2 ml collecting tube (supplied). Centrifuge ≥ 10,000 x g for 30 sec at room temperature. Discard flow-through liquid and reuse collecting tube in next step.
- 9. Add 500 μ I RNA Wash Buffer I and centrifuge at 10,000 x g for 30 sec. Discard both

flow-through liquid and collecting tube.

Optional on-membrane DNase I digestion: This is the starting point to perform DNase I digestion. See Page 7 for detailed protocol.

 Place column in a clean 2ml collection tube (supplied), and add 400 µl Wash Buffer II diluted with ethanol. Centrifuge at 10,000 x g for 30 sec at room temperature and discard flow-through. Reuse the collection tube in step.

Note: Wash Buffer II Concentrate must be diluted with absolute ethanol before use. Refer to label on bottle for directions.

- Wash column with a second 400 µl of Wash Buffer II as in step 10. Centrifuge and discard flow-through. Then with the collection tube empty, centrifuge the spin cartridge for 2 min at full speed to completely dry the HiBind[™] matrix.
- 12. Elution of RNA. Transfer the column to a clean 1.5 ml microfuge tube (not supplied with kit) and elute the RNA with 30-50 μl of DEPC-treated water (supplied with kit). Make sure to add water directly onto column matrix. Centrifuge 1 min at maximum speed. A second elution into the same tube may be necessary if the expected yield of RNA >50 μg. To maintain a high RNA concentration, use the first eluate for the second elution.

Note: RNA may be eluted with a greater volume of water. While additional elutions increase total RNA yield, the concentration will be lowered since more than 80% of RNA is recovered with the first elution.

DNase digestion Protocol (Optional)

Since HiBind[®] RNA resin and spin-column technology actually removes most of DNA without the DNase treatment, it is not necessary to do DNase digestion for most downstream applications. However, certain sensitive RNA applications might require further DNA removal. Following steps provide on-membrane DNase I digestion:(see DNase I cat.# E1091for detail information)

- 1. Follow the standard protocol until the samples completely pass through the HiBind RNA column (step1-9). Prepare the following:
 - a. For each HiBind[®] RNA column, prepare the DNase I digestion reaction mix as follows:

OBI DNase I Digestion Buffer	73.5 µl
RNase-free DNase I (20 Kunitz unites/µI)	1.5 µl
Total volume	75 µl

Note:

- 1. DNase I is very sensitive for physical denaturaion, so do not vortex this DNase I mixture. Mix gently by inverting the tube. Prepare the fresh DNase I digestion mixture before RNA isolation.
- 2. OBI DNase I digestion buffer is supplied with OBI RNase-free Dnase set.
- 3. Standard Dnase buffers are not compatible with on-membrane Dnase digestion.

b. Pipet 75 µl of the DNase I digestion reaction mix directly onto the surface of HiBind[®] RNA resin in each column. Make sure to pipet the Dnase I digestion mixture directly onto the membrane. Dnase I digestion will not be complete if some of the mix stick to the wall or the O-ring of the HiBind[®] RNA column.

c. Incubate at room temperature(25-30°C) for 15 minutes

Place column in a clean 2ml collection tube, and add 500 µl RNA Wash Buffer

 (If on-membrane DNase digestion was performed in the previous step, wait
 at least 5 minutes before proceeding). Centrifuge and discard flow-through.
 Reuse the collection tube in step 7.

 Place column in the same 2ml collection tube, and add 400 µl RNA Wash Buffer II diluted with ethanol. Centrifuge and discard flow-through. Reuse the collection tube in step 4.

Note: Wash Buffer II Concentrate must be diluted with absolute ethanol before use. Refer to label on bottle for directions.

- 4. Wash column with a second 400 µl of Wash Buffer II as in step 5. Centrifuge and discard flow-through. Then with the collection tube empty, centrifuge the spin cartridge for 1 min at full speed to completely dry the HiBind[®] matrix.
- 5. Elution of RNA. Transfer the column to a clean 1.5 ml microfuge tube (not supplied with kit) and elute the RNA with 30-50 μl of DEPC-treated water (supplied with kit). Make sure to add water directly onto column matrix. Centrifuge 1 min at maximum speed. A second elution may be necessary if the expected yield of RNA >50 μg. Alternatively, RNA may be eluted with a greater volume of water. While additional elutions increase total RNA yield, the concentration will be lowered since more than 80% of RNA is recovered with the first elution. Pre-heating the water to 70°C before adding to column and incubating column 5 min at room temperature before centrifugation may increase yields.

8

Quantization and Storage of RNA

To determine the concentration and purity of RNA, measure absorbance at 260 nm and 280 nm in a spectrophotometer. 1 O.D. unit measured at 260 nm corresponds to 40 μ g of RNA per ml. The ratio of A₂₆₀/A₂₈₀ of pure nucleic acids is 2.0, while for pure protein it is approximately 0.6. A ratio of 1.8-2.0 corresponds to 90%-100% pure nucleic acid. Store RNA samples at -70°C in water. Under such conditions RNA prepared with the E.Z.N.A. system is stable for more than a year.

RNA Quality

It is highly recommended that RNA quality be determined prior to all analyses. The quality of RNA can be assessed by denaturing agarose gel electrophoresis and ethidium bromide staining. Several sharp bands should appear on the gel. These are the 28S and 18S ribosomal RNA bands as well as certain populations of mRNA and possibly viral RNA bands. If these bands smear towards lower molecular weight RNAs, then the RNA has undergone major degradation during preparation, handling, or storage. RNA molecules less than 200 bases in length do not efficiently bind the HiBind matrix, thus the method enriches high quality mRNA.

Troubleshooting Guide

Problem	Cause	Suggestion	
Little or no RNA eluted	RNA remains on the column	 Repeat elution. Pre-heat DEPC-water to 70° C prior to elution. Incubate column for 2 min with water prior to centrifugation. 	
	Column is overloaded	Reduce quantity of starting material.	
Clogged column	Incomplete disruption or lysis of tissue.	 Completely disrupt sample in liquid nitrogen. Increase centrifugation time. Reduce amount of starting material 	
Precipitated RNA will not dissolve.	High nucleic acid and polysaccharide content.	 Reduce amount of starting material. Generally it is best to start with 50-100 mg at first. To avoid RNA degradation, do not increase incubation time for resuspension. 	
Degraded RNA	Source	 Freeze starting material quickly in liquid nitrogen and store at -70°C without thawing. Follow protocol closely, and work quickly. Make sure that 2-mercaptoethanol is added to Buffer MRL. 	
	RNase contamination	 Ensure not to introduce RNase during the procedure. Check buffers for RNase contamination. 	
Problem in downstream applications	Salt carry-over during elution	 Ensure Wash Buffer II has been diluted with 100% ethanol as indicated on bottle. Diluted Wash Buffer II must be stored at room temperature. Repeat wash with Wash Buffer II. 	
DNA contamination	Co-purification of DNA	 Digest with RNase-free DNase and inactivate at 75°C for 5 min. 	
Low Abs ratios	RNA diluted in acidic buffer or water	 DEPC-treated water is acidic and can dramatically lower Abs260 values. Use TE buffer (pH 8) to dilute RNA prior to spec analysis. 	

Order Information

Product No.	Product Name	Description
Blood DNA Kits		
D3481-01/02 D3491-01/02	Blood DNA Kit I (Proteinase included)	Isolation of total cellular DNA from fresh and dry blood
D3482-01/02 D3492-01/02	Blood DNA Kit II (OB Protease included)	Isolation of total cellular DNA from fresh and dry blood
D3493-01/02	Blood DNA Midiprep Kit (Proteinase not included)	Isolation of total cellular DNA from up to 10ml blood.
D3494-01/02	Blood DNA Midiprep Kit (OB Proteinase included)	Isolation of total cellular DNA from up to 10ml blood.
D2492-01/02	Blood DNA Maxiprep Kit (OB Proteinase included)	Isolation of total cellular DNA from up to 30ml blood.
D1192-01/02	EZ 96™ Blood DNA Kit	Isolation of total cellular DNA from fresh and dry blood with 96 well plate
Tissue DNA kits		
D3395-01/02 D3495-01/02	Tissue DNA Kit I (Proteinase not included)	Isolation of total cellular DNA from tissue samples.
D3396-01/02 D3496-01/02	Tissue DNA Kit II (OB Protease included)	Isolation of total cellular DNA from tissue samples.
D1196-01/02	EZ 96™ Tissue DNA Kit	Isolation of total cellular DNA from tissue samples with 96 well plate
D3592-01/02	Forensic DNA Kit	Isolation of genomic DNA from forensic samples. OB collection paper included.

Product Number	Product Name	Description		
E.Z.N.A.™ Total RNA	E.Z.N.A.™ Total RNA Miniprep Kits			
R6634-01/02 R6834-01/02	E.Z.N.A.™ Total RNA Kit	Total RNA isolation from animal cells or tissues.		
R6627-01/02 R6827-01/02	E.Z.N.A.™ Plant RNA Kit	Total RNA Isolation from plant samples		
R6640-01/02 R6840-01/02	E.Z.N.A.™ Fungal RNA Kit	Total RNA Isolation from fungal samples		
R6670-01/02 R6870-01/02	E.Z.N.A.™ Yeast RNA Kit	Total RNA Isolation from yeast samples		
R6850-01/02 R6950-01/02	E.Z.N.A.™ Bacterial RNA Kit	Total RNA Isolation from yeast samples		
R6675-01/02 R6875-01/02	E.Z.N.A.™ Mollusc RNA Kit	Total RNA Isolation from mollusc, invertebrates samples.		
E.Z.N.A.™ Total RNA	Midi/maxi Kits			
R6664-01/02	E.Z.N.A.™ Total RNA Midi Kit	Total RNA isolation from animal cells or tissues		
R6693-01/02	E.Z.N.A.™ Total RNA Maxi Kit	Total RNA isolation from animal cells or tissues		
R6615-01/02	E.Z.N.A.™ Blood RNA Midi Kit	Total RNA isolation from blood samples		
R6616-01/02	E.Z.N.A.™ Blood RNA Maxi Kit	Total RNA isolation from blood samples		
R6628-01/02	E.Z.N.A.™ Plant RNA Midi Kit	Total RNA isolation from plant samples		
Other RNA isolation kit, Reagent and supplies				
R6511-01/02	mRNA Enrichment kit	mRNA isolation		
R6830-01/02	RNA-Solv™ reagent	Single reagent for total RNA isolation		
R6248-01/02 R6249-01/02	E.Z.N.A.™ RNA Probe purification kit	RNA Probe purification		
R6376-01/02	E.Z.N.A.™ Poly-Gel RNA Isolation Kit	Isolate RNA from poly-acrylamide gel		
R6500-01/02	E.Z.N.A.™ Oligo (dT) Cellulose	High capacity oligo(dT) cellulose		