



### Direct-zol™-96 RNA

TRIzol® In. RNA Out.

#### **Highlights**

- 96-well spin-plate purification of total RNA (including small/ microRNAs) directly from TRIzol®, TRI Reagent® or similar acidguanidinium-phenol based reagents.
- No need for chloroform, phase-separation or precipitation steps.
- RNA is ready for Next-Gen Sequencing, RT-qPCR, etc. DNase I is included.

Catalog Numbers: R2054, R2055, R2056, R2057



Scan with your smart-phone camera to view the online protocol/video.



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Revised on: 5/15/2023

### **Product Contents**

Direct-zol <sup>™</sup> -96 RNA	<b>R2054</b> (2 x 96)	<b>R2055</b> (2 x 96)	<b>R2056</b> (4 x 96)	<b>R2057</b> (4 x 96)
TRI Reagent®	-	200 ml	-	200 ml (x2)
Direct-zol™ RNA PreWash <sup>1</sup> (concentrate)	160 ml	160 ml	160 ml (x2)	160 ml (x2)
RNA Wash Buffer <sup>2</sup> (concentrate)	48 ml	48 ml	48 ml (x2)	48 ml (x2)
DNase I <sup>3</sup> (lyophilized)	1500 U (x4)	1500 U (x4)	1500 U (x8)	1500 U (x8)
DNA Digestion Buffer	4 ml (x2)	4 ml (x2)	16 ml	16 ml
DNase/RNase-Free Water	10 ml	10 ml	30 ml	30 ml
Zymo-Spin <sup>™</sup> I-96 Plate	2	2	4	4
Collection Plate	4	4	8	8
Elution Plate	2	2	4	4
96-Well Plate Cover Foil	2	2	4	4
Instruction Manual	1 pc	1 pc	1 pc	1 pc

**Storage Temperature** - Store all kit components (i.e., buffers, spin-plate) at room temperature. Before use:

<sup>1</sup> Add 40 ml ethanol (95-100%) to the 160 ml **Direct-zol™ RNA PreWash** concentrate.

<sup>2</sup> Add 192 ml 100% ethanol (208 ml 95% ethanol) to the 48 ml RNA Wash Buffer concentrate.

<sup>3</sup> Reconstitute lyophilized DNase I with DNase/RNase-Free Water, mix by gentle inversion and store frozen aliquots:

<sup>#</sup>E1011-A (1500 U), add 275 μl water #E1009-A (250 U), add 55 μl water

### **Specifications**

- Sample Sources Any sample stored and preserved in TRIzol®, TRI Reagent® or similar¹. (animal cells, tissue, bacteria, yeast, biological fluids, samples stored in DNA/RNA Shield™ and in-vitro processed RNA (e.g., transcription products, DNase-treated or labeled RNA)).
- Sample Inactivation TRI Reagent® (provided with R2055 & R2057 only) inhibits RNase activity and inactivates viruses and other infectious agents.
- Size Total RNA including small/microRNAs (≥ 17 nt).
- Purity A<sub>260</sub>/A<sub>280</sub> & A<sub>260</sub>/A<sub>230</sub> > 1.8. RNA is ready for Next-Gen Sequencing, RT-qPCR, etc.
- Binding Capacity 10 µg RNA per well (Zymo-Spin<sup>™</sup> I-96 Plate).
- Compatibility TRIzol®, RNAzol®, QIAzol®, TriPure™, TriSure™ or similar acid-guanidinium-phenol based reagents can be used in place of TRI Reagent®.

Also, compatible with samples in TRIzol®, TRI Reagent® or similar reagent that contain chloroform, 1-bromo-3-chloropropane (BCP), or 4-bromoanisole (BAN), the aqueous phase of phase-separated samples and samples stored in RNAlater™ (page 8). For compatibility with cetyltrimethylammonium bromide (CTAB)-based extraction, see detailed protocol <a href="https://examples.com/here/based-extraction">here</a>.

- Elution Volume ≥ 10 µl DNase/RNase-Free Water.
- Equipment Needed (user provided): Centrifuge with microplate carriers.

<sup>1</sup> RNAzol®, QIAzol®, TriPure™, TriSure™ or similar acid-guanidinium-phenol reagent.

### **Product Description**

The **Direct-zol**<sup>™</sup>-96 **RNA** provides a streamlined method for the purification of up to 10 μg (per well) of high-quality RNA directly from samples in TRIzol<sup>®</sup>, TRI Reagent<sup>®</sup> or similar reagent<sup>1</sup>. Total RNA, including small RNAs (17-200 nt), is effectively isolated from a variety of sample sources (cells, tissues, serum, plasma, blood, biological liquids, etc.) using this product. The extraction method inactivates viruses and other infectious agents<sup>2</sup>.

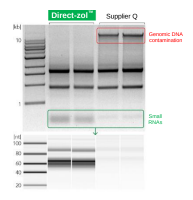
The procedure is easy: simply apply a sample in TRI Reagent® to the **Zymo-Spin™ I-96 Plate**, then bind, wash, and elute the RNA. No phase separation, precipitation, or post-purification steps are necessary. The result is high-quality RNA suitable for subsequent RNA-based methods including Next-Gen sequencing, RT-qPCR, hybridization, etc.

The entire procedure typically takes ~30 minutes (per 2 plates).

Sample in TRIzol®, TRI Reagent® or similar



# Efficient recovery of DNA-free Total RNA



(top) High-quality DNA-free RNA is purified from human epithelial cells using the **Direct- zol**™ procedure compared to a preparation from Supplier Q (1% agarose/TAE gel).

(bottom) Small RNAs are efficiently recovered with the Direct-zol" procedure. However, this is not the case with Supplier Q's prep (Bioanalyzer, Small RNA Chip).

<sup>1</sup> RNAzol®, QIAzol®, TriPure™, TriSure™ or similar acid-guanidinium-phenol reagent.

<sup>2</sup> TRI Reagent® provided with catalog #R2055 and #R2057.

### Input Capacity and Average RNA Yield

Input	Average RNA Yield	Kit Capacity
Cells	1 μg (per 10 <sup>5</sup> cells)	Up to 10 <sup>6</sup>
HeLa	1.5 μg	
High Yield Tissue <sup>1 (mouse)</sup>	≥ 3 µg (per 1 mg)	Up to 2 mg
Spleen	3-5 µg	
Liver	4-6 μg	
Low Yield Tissue <sup>1 (mouse)</sup>	≤ 3 µg (per 1 mg)	Up to 5 mg
Brain, Heart	0.5-1.5 μg	
Muscle	0.5-2 μg	
Lung	1-2 μg	
Intestine	1-3 µg	
Kidney	2-3 μg	
Whole Blood <sup>2</sup>	(per 100 μl)	Up to 200 μI
Porcine	1-2 µg	
Human	0.2-1 μg	

<sup>1</sup> Yield from tissue can vary due to other factors (i.e., organism type, physiological state, and growth conditions. 2 Yield from blood can vary based upon collection, sample preparation, donor, age, and/or health conditions.

#### **Protocol**

The protocol consists of: (I) Buffer Preparation, (II) Sample Preparation and (III) RNA Purification.

The following guidelines are provided for processing various sample types in TRIzol®, TRI Reagent® or similar¹ acid-guanidinium-phenol reagents prior to column purification of the RNA (see page 4 for Input Capacity and Total RNA Yield).

#### (I) Buffer Preparation

- ✓ Add 40 ml ethanol (95-100%) to the 160 ml Direct-zol™ RNA PreWash concentrate.
- ✓ Add 192 ml 100% ethanol (208 ml 95% ethanol) to the 48 ml RNA Wash Buffer concentrate.
- Reconstitute lyophilized DNase I with DNase/RNase-Free Water, mix by gentle inversion and store frozen aliquots:
   #E1011-A (1500 U), add 275 μl water
   #E1009-A (250 U), add 55 μl water

#### (II) Sample Preparation<sup>2</sup>

✓ Perform all steps at room temperature and centrifugation at 10,000-16,000 x g for 1 minute.

#### Cells

Lyse animal or gram(-) bacteria cells\* directly in a culture dish\*\* or resuspend pelleted cells in an appropriate volume (see table below) of TRI Reagent® or similar¹ and mix thoroughly. Proceed to RNA Purification (page 7).

Animal	Gram(-) bacteria	Add TRI Reagent®
≤ 10 <sup>5</sup>	-	≥ 100 µl
≤ 10 <sup>6</sup>	≤ 10 <sup>8</sup>	≥ 300 µl

<sup>\*</sup> For cell suspensions, add 3 volumes of TRI Reagent® to 1 volume of cell suspension.

<sup>\*\*</sup> For direct lysis in a dish, add 100 µl for each cm² of culture surface area.

<sup>1</sup> TRIzol®, RNAzol®, QIAzol®, TriPure™, TriSure™ or similar acid-guanidinium-phenol reagent.

<sup>2</sup> RNA yield can vary with sample types, organism, quality and treatment of the starting material (see page 4). To ensure complete lysis and homogenization of a sample, use a sufficient amount of TRIzol®, TRI Reagent® or similar reagent. For detailed processing information, refer to the TRI Reagent® product manual (or manufacturer's instructions for the reagent used).

#### **Tough-to-Lyse Samples**

Tough-to-lyse samples (see table below) can be homogenized in ≥ 800 µl TRIzol®, TRI Reagent® or similar1 with a mortar/pestle, dounce, 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. Proceed to RNA Purification (page 7).

Recommended: Use ZR BashingBead™ Lysis Tubes (materials sold separately; #S6012, #S6003, #S014) for complete lysis and homogenization.

Input	Gram(-) bacteria (optional; easy-to-lyse)	Gram(+) bacteria	<b>Tissue</b> (animal, plant*)	Pathogen (microbes in tissue)
per prep	bacteria (≤ 10 <sup>8</sup> )	bacteria (≤ 10 <sup>8</sup> ) yeast (≤ 10 <sup>7</sup> )	animal: high yield (≤ 2 mg) animal: low yield (≤ 5 mg) plant* (≤ 20 mg)	animal/insect, plant (≤ 5 mg)
lysis beads catalog #	0.5 mm and 0.1 mm; <b>S6012</b>	0.5 mm and 0.1 mm; <b>S6012</b>	2.0 mm; <b>S6003</b>	2.0 mm and 0.1 mm; <b>S6014</b>
high- speed <sup>2,3</sup>	30 sec	5-10 min	30-60 sec	3-5 min
low-speed <sup>3</sup>	5-10 min	20-40 min	3-5 min	5-10 min

<sup>\*</sup>Compatible with CTAB-based RNA extraction methods for polysaccharide-rich and/or phenolics-rich samples (e.g., Pinus, Geranium plants), Please find detailed protocol here.

#### Liquids

Add an appropriate volume of TRI Reagent® or similar1 to a liquid sample and mix thoroughly (see table below). To remove particulate debris (if any), centrifuge and transfer the supernatant into an RNase-free tube. Proceed to RNA Purification (page 7).

Recommended: For biological samples (whole-blood, plasma, serum, buffy coat, PBMCs, WBCs, FACS, etc.) or samples collected in DNA/RNA Shield™4, perform Proteinase K treatment<sup>5</sup> (sold separately) prior to adding TRI Reagent<sup>®</sup>.

Sample		Add TRI Reagent®
Biological liquid (blood, plasma, serum, WBCs, FACs, etc.) or		
<b>Reaction clean-up</b> (DNase I treated RNA, <i>in vitro</i> transcription, labeling, etc.).	100 μΙ	≥ 300 µl
Samples in DNA/RNA Shield™ (biological sample <sup>4,5</sup> or stored purified RNA).	100 μΙ	100 µl

<sup>1</sup> TRIzol®, RNAzol®, QIAzol®, TriPure™, TriSure™ or similar acid-guanidinium-phenol reagent.

<sup>2</sup> Perform high-speed homogenization at 1-minute intervals (including a cooling step for 3-5 minutes), to avoid overheating the machine and/or breaking the tube.

<sup>3</sup> High-speed homogenizers (e.g., MP Bio FastPrep-24™, Bertin Precellys, etc.). Low-speed homogenizers (e.g., Vortex Genie, etc.).

<sup>4</sup> DNA/RNA Shield™ reagent (R1100, R1200) or DNA/RNA Shield™ Blood Collection Tube (R1150).

<sup>5</sup> For Proteinase K treatment, see page 9.

#### (III) RNA Purification

- ✓ Perform all steps at room temperature and centrifugation at ≥ 2,500 x g for 5 minutes (spin-plate).
- ✓ Do not use the **96-Well Plate Cover Foil** on the spin-plate during RNA Purification. If necessary, use an Air Permeable Sealing Cover (#C2011-8); sold separately.
- 1. Add an equal volume ethanol (95-100%) to a sample lysed in TRI Reagent® or similar reagent¹ and mix well².

Example: Add 400 µl ethanol to 400 µl mixture (sample lysed in TRI Reagent®).

- Transfer the mixture to a well of the Zymo-Spin<sup>™</sup> I-96 Plate<sup>3</sup>
  mounted on a Collection Plate and centrifuge<sup>4</sup>. Mount the spinplate onto a new collection plate and discard the flow-through.
- 3. **DNase I**<sup>5</sup> treatment (recommended)
  - (D1) Add 400 µl RNA Wash Buffer to each well and centrifuge.
  - (D2) For each sample/well to be treated, add 5 µl **DNase I** (6 U/µl)\*, 35 µl **DNA Digestion Buffer** and mix by gentle inversion in an RNase-free tube (not included). Add 40 µl directly to the column matrix of each well.
  - (D3) Incubate at room temperature (20-30°C) for 15 minutes. Proceed to step 4.
- 4. Add 400 μl/well **Direct-zol<sup>™</sup> RNA PreWash**<sup>6</sup> to the plate and centrifuge. Discard the flow-through and repeat this step.
- Add 800 µl/well RNA Wash Buffer<sup>6</sup> to the plate and centrifuge.
   Discard the flow-through. To ensure complete removal of the wash buffer, centrifuge the plate again. Then mount the plate onto an Elution Plate.
- Add 25 μl/well of DNase/RNase-Free Water directly to the matrix and centrifuge.

Alternatively, for highly concentrated RNA use ≥ 10 µl elution.

The eluted RNA<sup>7</sup> can be used immediately or stored frozen.
Use the **96-Well Cover Foil** to prevent the eluate from evaporation.

<sup>1</sup> TRIzol®, RNAzol®, QIAzol®, TriPure™, TriSure™ or similar acid-guanidinium-phenol reagent.

<sup>2</sup> Mix well by pipetting up and down several times and/or by shaking (vortexing) at ~1,300 rpm. Optimization may be required.

<sup>3</sup> The well capacity is 800 μl. Reload the plate to process > 800 μl.

<sup>4</sup> At this point, proteins can be purified from the flow-through (see page 9).

<sup>5</sup> Prior to use, reconstitute the lyophilized **DNase I** (Buffer Preparation, page 5). \* Unit definition – one unit increases the absorbance of a high molecular weight DNA solution at a rate of 0.001 A260 units/ml of reaction mixture at 25°C.

<sup>6</sup> Before use, add ethanol to the buffer concentrate (Buffer Preparation, page 5).

<sup>7</sup> For complete removal of PCR (RT) inhibitors from plant, soil and fecal samples, use the OneStep™ PCR Inhibitor Removal Kit (D6030).

### **Appendices**

# RNA purification from aqueous phase after TRI Reagent® extraction

For samples that have already been phase separated in TRI Reagent<sup>®1</sup> or similar<sup>2</sup>, simply transfer the aqueous phase<sup>3</sup> containing RNA into an RNase-free tube. Add an equal volume ethanol (95-100%) to the aqueous phase (1:1) and mix thoroughly. Transfer up to 800 µl of the mixture into a plate/tube and proceed to RNA Purification (page 7, step 2).

#### RNA extraction from samples stored in RNAlater™

#### Cells

Pellet cells<sup>4</sup> at up to 5,000 x g and remove the RNAlater<sup>™</sup> (supernatant) prior to RNA extraction. Then lyse the cell pellet in TRI Reagent<sup>®</sup> (Sample Preparation, page 5).

Note: To extract RNA from cells without reagent removal, use 10 volumes of TRI Reagent® per sample volume. Proceed to phase separation and process the aqueous phase. Simply transfer the aqueous phase containing RNA into an RNase-free tube. Then add an equal volume ethanol (95-100%) to the aqueous phase (1:1) and mix thoroughly. Transfer up to 800 µl of the mixture into a plate/tube and proceed to RNA Purification (page 7, step 2).

#### Tissue

Remove tissue from RNAlater™ using forceps. Eliminate any excess reagent or crystals that may have formed and proceed immediately with extraction in TRI Reagent® (Sample Preparation, page 6).

<sup>1</sup> For detailed processing information, refer to the TRI-Reagent® product manual (or manufacturer's instructions for the reagent used).

<sup>2</sup> TRIzol®, RNAzol®, QIAzol®, TriPure™, TriSure™ or similar acid-guanidinium-phenol reagents.

<sup>3</sup> Alternatively, the aqueous phase can be processed with the RNA Clean & Concentrator™ (R1015).

<sup>4</sup> Different cells may react differently to centrifugation forces and it is recommended to test the pelleting procedure with non-valuable samples first. Diluting RNAlater™ by 50% with cold PBS reduces solution density allowing for lower forces during cell pelleting (e.g., 500 x g).

#### **Protein Purification**

The protein content in the flow-through after the RNA binding to the column can be purified (see RNA Purification, page 7, step 2):

- 1. Add 4 volumes of cold acetone (-20°C) to the flow-through (4:1) and mix.
- 2. Incubate the samples for 30 minutes on ice.
- 3. Centrifuge at max speed for 10 minutes. Discard the supernatant. Keep the pellet.
- Add 400 µl ethanol (95-100%) to the protein pellet. Centrifuge at max speed for 1 minute. Discard the supernatant.
- 5. Air-dry the protein pellet for 10 minutes at room temperature.
- Resuspend and vortex the pellet in a buffer appropriate for 6. downstream application (e.g., SDS-PAGE sample loading buffer).

#### **Proteinase K Treatment**

Proteinase K treatment can be performed on protein-rich samples stored in DNA/RNA Shield™ (2X concentrate; #R1200) (e.g., tissue, blood cells, plasma, serum, saliva, sputum, etc.) using Proteinase K Set (#D3001-2-5, D3001-2-20; sold separately).

Add 10 µl Proteinase K (reconstituted) to 1 ml DNA/RNA Shield sample (scale proportionally) and mix by inversion. Then incubate at room temperature (20-30°C) for 30 minutes (homogenized) or 2-5 hours (nonhomogenized). Optimization may be required.

#### **Compatibility with CTAB-Based Methods**

The Direct-zol RNA kits are compatible with CTAB-based RNA extraction methods for polysaccharide-rich and/or phenolics-rich samples (e.g., Pinus, Geranium plants). Please find detailed protocol here.

<sup>1</sup> Yield from tissue can vary due to other factors (i.e., organism type, physiological state and growth conditions.

<sup>2</sup> Yield from blood can vary based upon collection, sample preparation, donor, age, and/or health conditions.

# **Ordering Information**

Product Description	Catalog No.	Size
<b>Direct-zol™-96 RNA</b>	R2054	2 x 96 preps.
(TRI Reagent <sup>®</sup> <u>not</u> included)	R2056	4 x 96 preps.
Direct-zol™ -96 RNA	R2055	2 x 96 preps.
(supplied with TRI Reagent®)	R2057	4 x 96 preps.

Individual Kit Components	Catalog No.	Amount
TRI Reagent®	R2050-1-50 R2050-1-200	50 ml 200 ml
Direct-zol RNA PreWash (concentrate)	R2050-2-40 R2050-2-160	40 ml 160 ml
RNA Wash Buffer (concentrate)	R1003-3-24 R1003-3-48	24 ml 48 ml
Zymo-Spin I-96 Plates	C2004	2 plates
<b>DNase I</b> (lyophilized) (250 U supplied with DNA Digestion Buffer, 4 ml)	E1010	1 set
DNase/RNase-Free Water	W1001-10 W1001-30	10 ml 30 ml
Collection Plate (capacity 1.2 ml/well)	C2002	2 plates
96-Well Block (capacity 2 ml/well)	P1001-2	2 plates
Elution Plate (capacity 0.35 ml/well)	C2003	2 plates
96-Well Plate Cover Foil	C2007-2	2
DNA/RNA Shield™ (2X concentrate)	R1200-25 R1200-125	25 ml 125 ml
Proteinase K Set (w/ Storage Buffer)	D3001-2-5 D3001-2-20	5 mg 20 mg

## **Complete Your Workflow**

✓ For tough-to-lyse samples in TRIzol, use ZR BashingBead Lysis Tubes:

ZR BashingBead Lysis Tubes	
2.0 mm beads #S6003	For plant/animal tissue
0.1 + 0.5 mm beads #S6012	For microbes
0.1 + 2.0 mm beads #S6014	For microbes in tissue/insects

✓ The only direct, high-throughput and automatable RNA purification from sample lysates in TRIzol (DNase I Set included with all formats):



Direct-zol RNA kits	
Microprep #R2060-R2063	From 1 cell and up
Miniprep #R2050-R2053	Up to 50 ug RNA
Miniprep Plus #R2070-R2073	Up to 100 ug RNA
96-well #R2054-R2057	Spin-plate
MagBeads #R2100-R2105	Automatable (Tecan, Hamilton, Kingfisher, etc.)

✓ For RNA clean-up (purification) from the aqueous phase (e.g., TRIzol, TRI Reagent or similar) or from any enzymatic reaction (e.g., DNase I treated RNA):



RNA Clean & Concentrator kit	
#R1013-R1014	DNase I Set included

✓ For NGS:

Zymo-Seq RiboFree Total RNA Library Prep kit		
#R3000	12 preps	
#R3003	96 preps	

# **Troubleshooting Guide**

Problem	Possible Causes and Suggested Solutions			
Precipitation, viscous	Incomplete lysis and/or high-mass input:			
lysate	- If precipitation occurs (upon adding ethanol to the lysate) or if the lysate is extremely viscous, increase the volume of TRIzol®, TRI Reagent® or similar reagent to ensure complete lysis and homogenization until lysate is transparent (see image).			
Low purity	Sample handling:			
(A <sub>260</sub> /A <sub>230</sub> nm, A <sub>260</sub> /A <sub>280</sub> nm)	- 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.			
	Incomplete lysis and/or cellular debris:			
	<ul> <li>Increase the volume of TRIzol®, TRI Reagent® or similar to ensure complete lysis and homogenization. Be sure to centrifuge any cellular debris and then process the cleared lysate.</li> </ul>			
Low yield	Sample input:			
	- 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 TRIzol®, TRI Reagent® or similar reagent.			
	High-protein content (blood, plasma/serum, etc.)			
	- Perform Proteinase K treatment to the sample prior to adding TRIzol®, TRI Reagent® or similar reagent (Sample preparation, page 6).			
DNA contamination	To remove DNA:			
	- Perform in-column DNase I treatment (page 7) or perform DNase I treatment post-purification (R1013, page 4), then re-purify the treated sample.			
	-For future preps, increase the volume of TRIzol®, TRI Reagent® or similar reagent to ensure complete lysis and homogenization of the sample.			
RNA degradation	To prevent RNA degradation:			
	-Immediately collect and lyse fresh sample into TRIzol®, TRI Reagent® or similar reagent to ensure RNA stability. Homogenized samples in TRIzol®, TRI Reagent® or similar can be stored frozen for later processing.			

For technical assistance, please contact 1-888-882-9682 or email tech@zymoresearch.com

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