

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 acid-guanidinium-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.



Table of Contents

Product Contents	01
Specifications	02
Product Description	03
Input Capacity & Average Yield Chart	04
Protocol	05
(I) Buffer Preparation	05
(II) Sample Preparation	05
Cells: Animal, Bacteria	05
Tough-to-Lyse Samples: Tissue, Microbes, etc.	06
Blood, Rxn Clean-up, DNA/RNA Shield Samples	06
(III) RNA Purification	07
Appendices	08
RNA Purification from TRIzol Aqueous Phase	08
RNAlater Samples	08
Protein Purification	09
Proteinase K Treatment	09
Ordering Information	10
Complete Your Workflow	11
Troubleshooting Guide	12
Notes	13
Guarantee	17

Product Contents

Direct-zol™ -96 RNA	R2054 (2 x 96)	R2055 (2 x 96)	R2056 (4 x 96)	R2057 (4 x 96)
TRI Reagent®	-	200 ml	-	200 ml (x2)
Direct-zol™ RNA PreWash ¹ (concentrate)	160 ml	160 ml	160 ml (x2)	160 ml (x2)
RNA Wash Buffer ² (concentrate)	48 ml	48 ml	48 ml (x2)	48 ml (x2)
DNase I ³ (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™ 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:

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

2 Add 192 ml 100% ethanol (208 ml 95% ethanol) to the 48 ml **RNA Wash Buffer** concentrate.

3 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**

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_{260}/A_{280} & $A_{260}/A_{230} > 1.8$. RNA is ready for Next-Gen Sequencing, RT-qPCR, etc.
- **Binding Capacity** – 10 µg RNA per well (**Zymo-Spin™ 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).

- **Elution Volume** – ≥ 10 µl **DNase/RNase-Free Water**.
- **Equipment² Needed** (user provided): Centrifuge with microplate carriers.

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

² For samples > 700 µl, the vacuum manifold can be used at step 2 only (page 7). Mount the plate onto the manifold and load sample. Then centrifuge the plate to remove any residual buffer/sample. Proceed with the protocol by centrifuge.

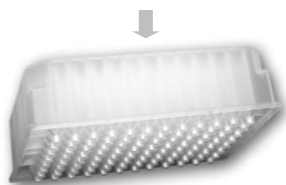
Product Description

The **Direct-zol™-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®, TRI Reagent® or similar reagent¹. 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 agents².

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).

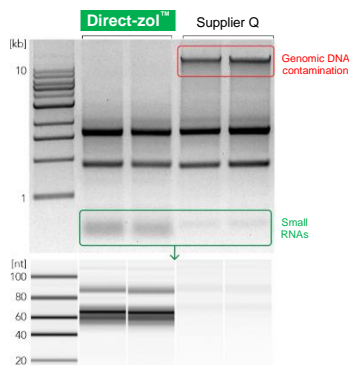
Sample in TRIzol®, TRI Reagent® or similar



Spin
Wash
Elute

Ready to use RNA

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).

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

2 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 ⁵ cells)	Up to 10 ⁶
HeLa	1.5 µg	
High Yield Tissue ¹ (mouse)	≥ 3 µg (per 1 mg)	Up to 2 mg
Spleen	3-5 µg	
Liver	4-6 µg	
Low Yield Tissue ¹ (mouse)	≤ 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 ²	(per 100 µl)	Up to 200 µl
Porcine	1-2 µg	
Human	0.2-1 µg	

1 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 Preparation²

- ✓ 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 ⁵	-	≥ 100 µl
≤ 10 ⁶	≤ 10 ⁸	≥ 300 µl

* For cell suspensions, add 3 volumes of TRI Reagent® to 1 volume of cell suspension.

** For direct lysis in a dish, add 100 µl for each cm² of culture surface area.

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

2 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	Tissue	Pathogen (microbes in tissue)
per prep	bacteria (≤ 10 ⁸)	bacteria (≤ 10 ⁸) yeast (≤ 10 ⁷)	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; S6012	0.5 mm and 0.1 mm; S6012	2.0 mm; S6003	2.0 mm and 0.1 mm; S6014
high- speed ^{2,3}	30 sec	5-10 min	30-60 sec	3-5 min
low-speed ³	5-10 min	20-40 min	3-5 min	5-10 min

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 treatment⁵ (sold separately) prior to adding TRI Reagent®.

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

1 TRIzol®, RNAzol®, QIAzol®, TriPure™, TriSure™ or similar acid-guanidinium-phenol reagent.
2 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.
3 High-speed homogenizers (e.g., MP Bio FastPrep-24™, Bertin Precellys, etc.). Low-speed homogenizers (e.g., Disruptor Genie, etc.).
4 DNA/RNA Shield™ reagent (R1100, R1200) or DNA/RNA Shield™ Blood Collection Tube (R1150).
5 For Proteinase K treatment, see page 9.

(III) RNA Purification

- ✓ Perform all steps at room temperature and centrifugation at $\geq 2,500 \times g$ 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.
 2. Transfer the mixture to a well of the **Zymo-Spin™ I-96 Plate**³ mounted on a **Collection Plate** and centrifuge⁴. Mount the spin-plate onto a new collection plate and discard the flow-through.
 3. **DNase I**⁵ treatment (recommended)
 - (D1) Add 400 μ l **RNA Wash Buffer** to each well and centrifuge.
 - (D2) For each sample/well to be treated, add 5 μ l **DNase I** (6 U/ μ l)*, 35 μ l **DNA Digestion Buffer** and mix by gentle inversion in an RNase-free tube (not included). Add 40 μ l 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 μ l/well **Direct-zol™ RNA PreWash**⁶ to the plate and centrifuge. Discard the flow-through and repeat this step.
 5. Add 800 μ l/well **RNA Wash Buffer**⁶ 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**.
 6. Add 25 μ l/well of **DNase/RNase-Free Water** directly to the matrix and centrifuge.

Alternatively, for highly concentrated RNA use $\geq 10 \mu$ l elution.

The eluted RNA⁷ can be used immediately or stored frozen.

Use the **96-Well Cover Foil** to prevent the eluate from evaporation.

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

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

3 The well capacity is 800 μ l. Reload the plate to process > 800 μ l.

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

5 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.

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

7 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®¹ or similar², simply transfer the aqueous phase³ 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 RNeasy™

Cells

Pellet cells⁴ at up to 5,000 x g and remove the RNeasy™ (supernatant) prior to RNA extraction. Then lyse the cell pellet in TRI Reagent® (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 RNeasy™ using forceps. Eliminate any excess reagent or crystals that may have formed and proceed immediately with extraction in TRI Reagent® (Sample Preparation, page 6).

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

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

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

4 Different cells may react differently to centrifugation forces and it is recommended to test the pelleting procedure with non-valuable samples first. Diluting RNeasy™ 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.
4. 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.
6. Resuspend and vortex the pellet in a buffer appropriate for 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 (non-homogenized). Optimization may be required.

1 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.

Ordering Information

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

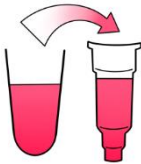
Complete Your Workflow

- ✓ 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

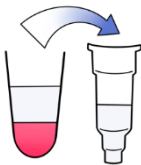
- ✓ The only **direct**, high-throughput and automatable RNA purification from sample lysates in TRIzol (DNase I Set included with all formats):



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

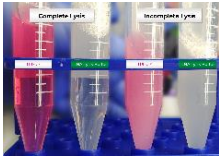
#R1013-R1014	DNase I Set included
--------------	----------------------

- ✓ 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 lysate	<p>Incomplete lysis and/or high-mass input:</p> <ul style="list-style-type: none"> - 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 (A_{260}/A_{230} nm, A_{260}/A_{280} nm)	<p>Sample handling:</p> <ul style="list-style-type: none"> - 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. <p>Incomplete lysis and/or cellular debris:</p> <ul style="list-style-type: none"> - 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.
Low yield	<p>Sample input:</p> <ul style="list-style-type: none"> - 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. <p>High-protein content (blood, plasma/serum, etc.)</p> <ul style="list-style-type: none"> - Perform Proteinase K treatment to the sample prior to adding TRIzol®, TRI Reagent® or similar reagent (Sample preparation, page 6).
DNA contamination	<p>To remove DNA:</p> <ul style="list-style-type: none"> - 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	<p>To prevent RNA degradation:</p> <ul style="list-style-type: none"> - 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

Notes

[illegible]

Notes

[illegible]

Notes

This image shows a single sheet of white paper with horizontal ruling lines. The lines are evenly spaced and run across the width of the page. There are no margins, text, or other markings on the paper.

Notes

This image shows a single sheet of white paper with horizontal ruling lines. The lines are evenly spaced and run across the width of the page. There are no margins, text, or other markings on the paper.



100% satisfaction guarantee on all Zymo Research products, or your money back.

Zymo Research is committed to simplifying your research with quality products and services. If you are dissatisfied with this product for any reason, please call 1(888) 882-9682.

Integrity of kit components is guaranteed for up to one year from date of purchase. Reagents are routinely tested on a lot-to-lot basis to ensure they provide the highest performance and reliability.

This product is for research use only and should only be used by trained professionals. It is not for use in diagnostic procedures. Some reagents included with this kit are irritants. Wear protective gloves and eye protection. Follow the safety guidelines and rules enacted by your research institution or facility.

TM Trademarks of Zymo Research Corporation

Direct-zolTM product technologies are subject to U.S. and foreign patents or are patent pending. U.S. Patent Nos. 9,051,563 B2; and 9,206,469 B2 and foreign equivalents. Direct-zol[®] is a registered trademark of Zymo Research Corporation. Other trademarks: TRI Reagent[®], TRIzol[®] and RNAzol[®] (Molecular Research Center, Inc.), QIAzol[®] (Qiagen GmbH), TriPure[™] (Roche, Inc.), TriSure[™] (Bioline Ltd.), RNAlater[®] (Ambion, Inc.), Bioanalyzer (Agilent Technologies, Inc.).

*The **BEAUTY** of **SCIENCE** is to Make Things **SIMPLE**®*

