



Quick-DNA/RNA™ Microprep Plus Kit

DNA & RNA from any sample

Highlights

- Spin-column purification of DNA and total RNA (including small/microRNAs) from any sample including cells, solid tissue, biological liquids, environmental samples, swabs, and any sample in DNA/RNA Shield™
- Extract DNA and total RNA from low inputs (down to a single cell).
- DNA/RNA Shield™ and Proteinase K are included for unique preservation and lysis technology.
- DNA & RNA are eluted in two separate fractions and are ready for Next-Gen Sequencing, RT/qPCR, etc. DNase I is included.

Catalog Numbers: D7005T, D7005



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







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Product Contents

<i>Quick</i> -DNA/RNA [™] Microprep Plus Kit	D7005T (10 prep)	D7005 (50 prep)
DNA/RNA Lysis Buffer	10 ml	50 ml
DNA/RNA Prep Buffer	5 ml (x2)	50 ml
DNA/RNA Wash Buffer ¹	16 ml (x2) (ready-to-use)	24 ml (x2)
DNase/RNase-Free Water	1 ml (x2)	10 ml
DNase I ² (lyophilized)	50 U	250 U
DNA Digestion Buffer	0.8 ml	4 ml
DNA/RNA Shield [™] (2X concentrate)	5 ml	25 ml
PK Digestion Buffer	1 ml	5 ml
Proteinase K ³ (lyophilized) & Storage Buffer	5 mg (x2)	60 mg
Zymo-Spin™ IC-XM Columns	10	50
Zymo-Spin™ IC Columns	10	50
Collection Tubes	30	50 (x3)
Instruction Manual	1 pc	1 pc

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

¹ Add 96 ml 100% ethanol (104 ml 95% ethanol) to the 24 ml **DNA/RNA Wash Buffer concentrate** (D7005). **DNA/RNA Wash Buffer** (D7005T) is supplied ready-to-use and does not require the addition of ethanol.

² Reconstitute lyophilized **DNase I** with **DNase/RNase-Free Water**, mix by gentle inversion and store frozen aliquots:

[#]E1009-A (250 U), add 275 µl water

[#]E1009-A-S (50 U), add 55 µl water
3 Add Proteinase K Storage Buffer to the lyophilized Proteinase K, see Buffer Preparation, page 5. Store frozen aliquots.

Specifications

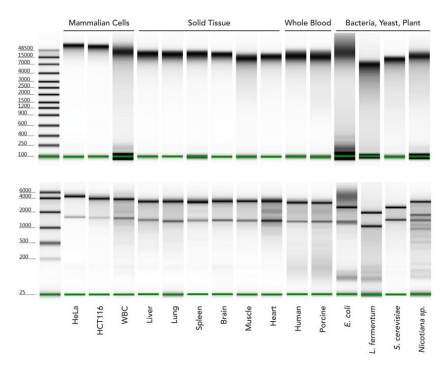
- Sample Sources Any cells (animal, bacterial, blood cells, etc.), all tissues (tough-to-lyse, FFPE, etc.), blood, biological fluids, enzymatic reactions (e.g., DNase I treated) and samples in DNA/RNA Shield™ or other preservation reagents.
- Sample Preservation and Inactivation DNA/RNA Shield™ lyses cells, inactivates nucleases and infectious agents (e.g., virus, pathogens) and is ideal for safe sample storage and transport at ambient temperatures (page 11).
- Size Genomic DNA (≥ 40 kb), mitochondrial and viral DNA (if present) and total RNA including small/microRNAs (≥ 17 nt).
- **Purity –** A_{260}/A_{280} & A_{260}/A_{230} > 1.8. DNA & RNA is ready for Next-Gen Sequencing, RT/qPCR, etc.
- Binding Capacity Zymo-Spin[™] IC-XM and Zymo-Spin[™] IC Column yield up to 5 µg DNA and 10 µg RNA, respectively.
- Compatibility For samples stored in preservation reagents: DNA/RNA Shield™, RNAprotect®, Allprotect®, Universal transport medium/viral transport medium (UTM®/VTM®), PAXgene® and RNAlater™.
- Elution Volume ≥ 6 µl DNase/RNase-Free Water.
- Equipment Needed (user provided) Microcentrifuge, vortex, heat block, water bath or incubator.

Product Description

The Quick-DNA/RNA™ Microprep Plus Kit combines Quick-DNA/RNA™ technology with the addition of DNA/RNA Shield™, a unique preservation and lysis technology, and Proteinase K to enable easy, reliable, and rapid isolation of DNA and total RNA from any low input biological sample including cells (1-10°), all tissues, blood, and other biological fluids.

The procedure uses unique spin-column technology that results in high-quality DNA and total RNA (including small RNAs 17-200 nt) that are ready for Next-Gen Sequencing, RT/qPCR, hybridization, etc.

High-Quality DNA & RNA From Any Sample Type



High quality genomic DNA (top) and total RNA (bottom) are isolated from various sample types including mammalian cells, solid tissue, whole blood, bacteria, yeast, and plant using the *Quick*-DNA/RNA™ Plus kits (Agilent 2200 TapeStation™).

Input Capacity and gDNA/Total RNA Yield

Input	Average gDNA Yield	Average RNA Yield	Kit Capacity
Cells	0.4 μg (per 10 ⁵ cells)	1 μg (per 10 ⁵ cells)	Up to 10 ⁶
HeLa	0.6 µg	1.5 µg	
High Yield Tissue ^{1 (mouse)}	≥ 3 µg (per 1 mg)	≥ 3 µg (per 1 mg)	Up to 2 mg
Spleen	5-7 μg	3-5 µg	
Liver	1.5-3 µg	4-6 μg	
Low Yield Tissue ^{1 (mouse)}	≤ 3 µg (per 1 mg)	≤ 3 µg (per 1 mg)	Up to 5 mg
Brain, Heart	0.5-1.5 μg	0.5-1.5 μg	
Muscle	0.5-1.5 μg	0.5-2 μg	
Lung	1.5-3 µg	1-2 µg	
Intestine	1.5-3 µg	1-3 µg	
Kidney	1.5-3 µg	2-3 µg	
Whole Blood ²	(per 100 μl)	(per 100 µl)	Up to 200 μl
Porcine	0.5-1 μg	1-2 µg	
Human	0.2 - 0.5 μg	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) DNA & RNA Purification.

(I) Buffer Preparation

- Add 96 ml 100% ethanol (104 ml 95% ethanol) to the 24 ml DNA/RNA Wash Buffer concentrate (D7005). DNA/RNA Wash Buffer (D7005T) is supplied ready-to-use and does not require the addition of ethanol.
- ✓ Reconstitute lyophilized DNase I with DNase/RNase-Free Water and mix by gentle inversion. Use immediately or store frozen aliquots:

```
#E1009-A (250 U), add 275 µl water
#E1009-A-S (50 U), add 55 µl water
#E1011-A (1500 U), add 1,500 µl water
```

Reconstitute lyophilized Proteinase K at 20 mg/ml with Proteinase K Storage Buffer and mix by vortexing. Use immediately or store frozen aliquots.

```
#D3001-2-20 (60 mg), add 3.12 ml buffer #D3001-2-5 (5 mg), add 0.26 ml buffer
```

✓ To prepare a 1X solution of DNA/RNA Shield[™], add an equal volume
of nuclease-free water (not provided) to the DNA/RNA Shield[™] (2X
concentrate) (1:1) and mix well.

(II) Sample Preparation

✓ Perform all steps at room temperature and centrifugation at 10,000-16,000 x g for 30 seconds, unless specified.

Samples stabilized and stored in DNA/RNA Shield[™] (cells, tissue, swab, etc.)

If frozen, thaw homogenized sample in **DNA/RNA Shield**^{M} to room temperature (20-30°C). Mix well by vortex. Proceed to the appropriate procedure below based on sample type (omit the step involving the addition of DNA/RNA Shield^{M}).

Cells & Tissue (mammalian)

- For samples (cells or tissue) already stored in DNA/RNA Shield[™], add an equal volume of DNA/RNA Lysis Buffer (1:1), mix well and proceed to purification, page 10.
- Cells: Pellet cell suspension by centrifugation (≤ 500 x g for 1 minute) or process adherent cells directly in culture container. Remove supernatant/media¹ respectively and resuspend cells in DNA/RNA Lysis Buffer (see table below). Proceed to purification, page 10.

Cells	Add DNA/RNA Lysis Buffer	
≤ 10 ⁶	≥ 400 µl	

 Tissue²: Submerge an appropriate amount of fresh or frozen sample (see table below) into DNA/RNA Shield[™] (1X)³ and homogenize^{4,5}.

Tissue	Add DNA/RNA Shield (1X)
High-yield (≤ 2 mg) Low-yield (≤ 5 mg)	≤ 600 µl

- a. For every 300 µl of sample, add 15 µl **Proteinase K** and 30 µl **PK Digestion Buffer**. Mix and incubate at room temperature (20-30°C) for ≥ 30 minutes (homogenized) or 2-5 hours (non-homogenized). Optimization may be required.
- b. To remove particulate debris, centrifuge and transfer the cleared supernatant into a nuclease-free tube (not provided).
- c. Add an equal volume of **DNA/RNA Lysis Buffer** to the supernatant (1:1) and mix well. Proceed to purification, page 10.

¹ If liquid/media cannot be removed, add ≥ 3 volumes DNA/RNA Lysis Buffer to 1 volume liquid sample (3:1) and mix well. Proceed to purification, page 10.

² For examples of sample type input and average yield, see chart on page 4.

³ For a 1X solution of **DNA/RNA Shield**[™], see Buffer Preparation, page 5.

⁴ For efficient homogenization, bead beat samples with ZR BashingBead Lysis Tubes (S6012, S6003), sold separately. See Appendices (page 12) for bead beating parameters. Other types of homogenization can include mortar/pestle, dounce, syringe or tissue grinder, etc.

⁵ Alternatively (if no homogenization), tissue samples can be Proteinase K treated only (proceed to step 3a).

<u>Tough-to-Lyse Samples</u> (bacteria, yeast, insect, swab, soil¹, stool¹, plant¹, seed¹)

1. Add 800 µl of **DNA/RNA Shield**[™] (1X)² to an appropriate amount of sample (see table below) and homogenize³ (e.g., bead beating).

Solid Tissue	Microbes	Add DNA/RNA Shield (1X)
Plant/Seed or Insect (≤ 20 mg)	Bacteria (≤ 2x10 ⁸) Yeast (≤ 2x10 ⁷) Swab, Stool/Soil (≤ 10 mg)	800 µl

- After homogenization, remove particulate debris by centrifugation at max speed. Transfer the cleared supernatant into a nuclease-free tube (not provided).
- 3. Add **DNA/RNA Lysis Buffer** to the supernatant (1:1), mix well and proceed with purification, page 10.

FFPE Tissue

- 1. Remove (trim) excess paraffin wax from ≤ 5 mg FFPE tissue and transfer into a nuclease-free tube (not provided).
- Add 400 μl Deparaffinization Solution⁴ to the sample. Incubate at 55°C for 1 minute. Vortex briefly. Remove the Deparaffinization Solution.
- 3. Add 95 μ l DNase/RNase-Free Water, 95 μ l 2X Digestion Buffer⁴, and 10 μ l Proteinase K. Mix well.
- 4. Incubate at 55°C for 1 hour. Then incubate at 65°C for 15 minutes to de-crosslink the sample.
- 5. Centrifuge to remove insoluble debris and transfer 200 μ l supernatant to a nuclease-free tube (not provided).
- 6. Add **DNA/RNA Lysis Buffer** to the supernatant (1:1) and mix well. Proceed to purification, page 10.

Blood Cells (mammalian, PBMCs, WBCs, etc.)

For blood cells, buffy coat and pelleted PAXgene[®] or RNAlater[™] samples, resuspend in **DNA/RNA Shield** (1X)².

Blood Cells	Add DNA/RNA Shield [™] (1X)	
≤ 0.5 ml blood (≤ 10 ⁶ cells)	300 μΙ	

2. For every 300 μl of sample, add 15 μl **Proteinase K** and 30 μl **PK Digestion Buffer**. Continue to step 3, page 8.

¹ For PCR inhibitor removal, use OneStep PCR Inhibitor Removal Kit (D6030).

² For a 1X solution of **DNA/RNA Shield™**, see Buffer Preparation, page 5.

³ For efficient homogenization, bead beat samples with ZR BashingBead Lysis Tubes (S6012, S6003), sold separately. See Appendices (page 12) for bead beating parameters.

⁴ Deparaffinization Solution (D3067-1-20) and 2X Digestion Buffer (D3050-1-20) are sold separately.

- 3. Mix and incubate at room temperature (20-30°C) for ≥ 30 minutes. Optimization may be required.
- 4. After incubation, vortex sample and centrifuge at max speed for 2 minutes to pellet debris. Transfer 300 μ l of the cleared supernatant to a nuclease-free tube (not provided).
- 5. Add **DNA/RNA Lysis Buffer** to the supernatant (1:1) and mix well. Proceed to purification, page 10.

Whole Blood^{1,2} (mammalian)

- Add 200 µl DNA/RNA Shield[™] (2X concentrate) directly to each 200 µl of fresh or frozen blood sample and mix thoroughly³.
- For every 400 μI of reagent/blood mixture, add 8 μI Proteinase K and mix thoroughly. Incubate at room temperature (20-30°C) for 30 minutes.
- After incubation, vortex sample and centrifuge at max speed for 2 minutes to pellet debris. Transfer the cleared supernatant to a new nuclease-free tube (not provided).
- 4. Add an equal volume of isopropanol (1:1) and mix well.
- 5. Proceed to purification, page 10.

Saliva & Buccal Cells

 For saliva and buccal cell samples, add an equal volume of DNA/RNA Shield™ (2X) (1:1).

Saliva & Buccal Cells	Add DNA/RNA Shield [™] (2X)
200 µl (≤ 10 ⁷ cells)	200 μΙ

- For every 400 μl of reagent/sample mixture, add 20 μl Proteinase K and 40 μl PK Digestion Buffer.
- 3. Mix and incubate at room temperature (20-30°C) for ≥ 30 minutes. Optimization may be required.
- After incubation, vortex sample and centrifuge at max speed for 2 minutes to pellet debris. Transfer 400 μl of the cleared supernatant to a nuclease-free tube (not provided).
- Add DNA/RNA Lysis Buffer to the supernatant (1:1) and mix well.
 Proceed to purification, page 10.

¹ Compatible with commonly used anticoagulants (e.g., EDTA, citrate, heparin)

² Up to 1 ml blood per prep can be processed (by reloading the column).

³ To retain protein in the whole blood sample, omit step 2 and continue to step 3.

Urine¹

Generate pellet from up to 40 ml urine by adding 70 µl Urine
 Conditioning Buffer² for every 1 ml of urine and mix by vortex.
 Centrifuge at 3,000 x g for 15 minutes. Discard the supernatant and keep the pellet. Add DNA/RNA Shield™ (1X)³ and mix by pipetting.

Pelleted cells from urine	Add DNA/RNA Shield [™] (1X)
≤ 40 ml urine	300 µl

- For every 300 μl of sample, add 15 μl Proteinase K.
- 3. Mix and incubate at room temperature (20-30°C) for ≥ 30 minutes. Optimization may be required.
- 4. After incubation, vortex sample and centrifuge at max speed for 2 minutes to pellet debris. Transfer 300 μ l of the cleared supernatant to a nuclease-free tube (not provided).
- 5. Add **DNA/RNA Lysis Buffer** to the supernatant (1:1) and mix well. Proceed to purification, page 10.

¹ Warm up urine sample at 37°C for 5-10 minutes if there is visual precipitation or cloudiness. Samples that contain bacterial contamination will not be clear.

² Urine Conditioning Buffer (D3061-1-8, D30601-1-140) is sold separately.

³ For a 1X solution of **DNA/RNA Shield**[™], see Buffer Preparation, page 5.

(III) DNA and RNA Purification (in two separate fractions)

- ✓ Perform all steps at room temperature and centrifuge at 10,000-16,000 x g for 30 seconds, unless specified.
- Transfer the sample into a Zymo-Spin[™] IC-XM Column¹ in a Collection Tube and centrifuge. Save the flow-through for RNA purification and the filter for DNA purification!

For **whole-blood samples only**: Discard the flow-through from step 1 (above). Then transfer the **Zymo-Spin IC-XM™ Column** into a nuclease-free tube (not provided). Add 200 µl **DNA/RNA Lysis Buffer** directly to the matrix of the filter, let stand for 5 minutes and then centrifuge. <u>Save the flow-through and the filter column!</u> Then proceed to purification below.

DNA Purification

(DNA is in the filter.)

2a. Transfer the Zymo-Spin™ IC-XM Column¹ into a new Collection Tube.

RNA Purification

(RNA is in the flow-through.)

2b. Add 1 volume ethanol (95-100%) to the flow-through and mix well.

Example: Add 400 μl ethanol to 400 μl flow-through.

Then transfer the mixture into a **Zymo-Spin**[™] **IC Column**¹ in a **Collection Tube** and centrifuge². Discard the flow-through.

At this point, **DNase I** treatment (incolumn) can be performed (see page 12).

- 3. Add 400 µl **DNA/RNA Prep Buffer** to the column and centrifuge. Discard the flow-through.
- 4. Add 700 μl **DNA/RNA Wash Buffer** to the column and centrifuge. Discard the flow-through.
- 5. Add 400 µl **DNA/RNA Wash Buffer** and centrifuge the column for 2 minutes to ensure complete removal of the wash buffer. Then carefully, transfer the column into a nuclease-free tube (not provided).
- Add 15 μl DNase/RNase-Free Water directly to the column matrix and centrifuge.

Alternatively, for highly concentrated DNA/RNA use ≥ 6 µl elution.

The eluted DNA/RNA³ can be used immediately or stored frozen.

¹ To process samples > 700 μ l, columns may be reloaded.

² Optional: At this point, proteins can be purified from the flow-through (page 12).

³ For complete removal of PCR (RT) inhibitors from plant, soil, and fecal samples, use the OneStep™ PCR Inhibitor Removal Kit (D6030).

Appendices

Sample stabilization and storage in DNA/RNA Shield™

<u>Liquid samples (e.g., whole-blood)</u>: Add 3 volumes **DNA/RNA Shield**[™] (1X)¹ to 1 volume sample (3:1). Mix well.

<u>Solid samples (e.g., tissue)</u>: Submerge sample (≤ 10% (v/v or w/v)) in **DNA/RNA Shield**[™] (1X)¹ and homogenize (Appendices, page 12).

Store samples in **DNA/RNA Shield**^{$^{\text{TM}}$} at ambient temperature for ≥ 1 month or long term at frozen temperature. DNA/RNA Shield $^{\text{TM}}$ is directly compatible with most guanidinium-based extraction methods (e.g., no need to remove reagent from the stored sample prior to extraction).

<u>Samples in RNAprotect, Allprotect, RNAlater, PAXgene, UTM/VTM, saline or PBS</u>

✓ RNAProtect®, Allprotect®: Add 3 volumes of DNA/RNA Lysis Buffer to 1 volume of sample (3:1). Mix well and/or homogenize base on sample type (see Sample Preparation, page 6), then proceed to purification, page 10.

✓ RNAlater™:

- a. Cells Pellet² by centrifugation at up to 5,000 x g and remove RNAlater (supernatant). Proceed to Sample Preparation, page 6.
- b. Tissue Transfer into a new tube with forceps and remove any excess RNAlater[™]. Proceed to Sample Preparation, page 6.

Alternatively, for liquid samples from which RNAlater cannot be removed, add 1 volume of nuclease-free water (or PBS) to 1 volume liquid sample (1:1) and mix. Then add 4 volumes **DNA/RNA Lysis Buffer** to 1 volume sample/water (or PBS) mixture (4:1). Mix again and proceed to Total RNA Purification, page 10.

- ✓ <u>PAXgene®</u>: Refer to manufacturer's instructions to remove the reagent then proceed to Sample Preparation, Blood Cells, page 7.
- ✓ <u>Swab samples in UTM®/VTM®</u>, <u>saline or PBS</u>: Remove swab and add 3 volumes of **DNA/RNA Lysis Buffer** to 1 volume sample (3:1). Mix well and proceed to purification, page 10.

Optional: To inactivate pathogens, store at room temperature prior to purification, add 1 volume of DNA/RNA Shield™ (2X concentrate) to 1 volume liquid sample (1:1) and mix well. Then proceed to Sample Preparation, Samples in DNA/RNA Shield™, page 6.

<u>Liquids/Reaction Clean-up</u> (DNase I treated RNA, in vitro transcriptions, etc.)

Add 150 μ l **DNA/RNA Lysis Buffer** to a \geq 50 μ l liquid sample (3:1) and mix well. Proceed to purification, page 10.

¹ For a 1X solution of **DNA/RNA Shield**[™], see Buffer Preparation, page 5.

² 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).

(Appendices continued)

DNase I Treatment (in-column)

- 1. Following RNA binding step (page 10, step 2b), add 400 µl **DNA/RNA Wash Buffer** to the column, centrifuge and discard the flow-through.
- 2. To treat each sample, prepare **DNase I Reaction Mix** (see table below) in a nuclease-free tube (not provided) and invert gently to mix.
- 3. Add 40 µl directly into the column matrix and incubate at room temperature (20-30°C) for 15 minutes. Proceed with the purification protocol (page 10, step 3).

DNase I Reaction Mix

DNase I (reconstituted; 1 U/µI) ¹	5 µl
DNA Digestion Buffer	35 µl

Protein Purification: Acetone Precipitation of Proteins

- ✓ After the RNA binding to the column (page 10, step 2b), the protein content (denatured) in the <u>flow-through</u> can be purified:
- 1. Add 4 volumes of cold acetone (-20°C) to 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).

Homogenization with ZR BashingBead Lysis Tubes

- ✓ Recommended for complete and efficient homogenization of tough-to-lyse samples (e.g., tissue, plant, seed, microbes, etc.). Lysis tubes sold separately.
- For high-speed homogenizers (e.g., MP Bio FastPrep-24, Bertin Precellys) and low-speed homogenizers (e.g., Vortex Genie), bead-beating time optimization may be required.

	Tissue		Microbes	
Input	Mammalian	Plant/Seed or Insect	Bacteria, Swab, Yeast, Stool/Soil	
Cat. no.	S6003	S6003	S6012	
(lysis bead size)	(2.0 mm)	(2.0 mm)	(0.5 mm and 0.1 mm)	
High-speed	30-60 sec	3-5 min	30-60 sec	
Low-speed	3-5 min	15-20 min	5-10 min	

¹ 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.

Ordering Information

Product Description	Catalog No.	Size
Quick-DNA/RNA [™] Microprep Plus Kit	D7005T D7005	10 preps 50 preps
Individual Kit Components	Catalog No.	Amount
DNA/RNA Lysis Buffer	D7001-1-50 D7001-1-200	50 ml 200 ml
DNA/RNA Prep Buffer	D7010-2-25 D7010-2-50	25 ml 50 ml
DNA/RNA Wash Buffer (concentrate)	D7010-3-12 D7010-3-24	12 ml 24 ml
DNase/RNase-Free Water	W1001-6 W1001-30	6 ml 30 ml
DNase I Set (lyophilized) (250 U supplied with DNA Digestion Buffer, 4 ml)	E1010	1 set
DNA/RNA Shield™ (2X concentrate)	R1200-25 R1200-125	25 ml 125 ml
PK Digestion Buffer	R1200-1-5 R1200-1-20	5 ml 20 ml
Proteinase K (lyophilized) & Storage Buffer	D3001-2-5 D3001-2-60	5 mg 60 mg
Zymo-Spin [™] IC-XM Columns	C1103-50	50
Zymo-Spin [™] IC Columns	C1004-50 C1004-250	50 250
Collection Tubes	C1001-50	50

Complete Your Workflow

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

ZR BashingBead Lysis Tubes	
2.0 mm beads #S6003	Plant/animal tissue
0.1 + 0.5 mm beads #S6012	Microbes
0.1 + 2.0 mm beads #S6014	Microbes in tissue/insects

✓ For isolation of DNA/RNA from any sample:

Quick-DNA/RNA Plus kits	
Microprep Plus #D7005	From 1 cell and up
MagBeads #R2130	Automatable (Tecan, Hamilton, Kingfisher, etc.)

✓ For clean-up (purification) and concentration of any RNA sample. (e.g., from the aqueous phase of TRIzol® extractions) or from any enzymatic reaction (e.g., DNase I treated RNA):

RNA Clean & Concentrator kits	
Microprep #R1013-R1014	DNase I Set included
MagBeads #R1082	Automatable (Tecan, Hamilton, Kingfisher, etc.)

✓ 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 DNA/RNA Shield™ and/or DNA/RNA Lysis Buffer to ensure complete lysis and homogenization until lysate is transparent (see image).			
Low purity (A ₂₅₀ /A ₂₃₀ nm, A ₂₅₀ /A ₂₉₀ nm)	Sample handling:			
	 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:			
	 Increase the volume DNA/RNA Shield[™] and/or DNA/RNA Lysis Buffer (proportionally) to ensure complete lysis and homogenization. Be sure to centrifuge and pellet any cellular debris then process the cleared lysate. 			
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 nucleic acid recovery. Use less input material and/or increase the volume DNA/RNA Shield™ and/or DNA/RNA Lysis Buffer.			
	High-protein content (blood, plasma/serum, etc.)			
	- Perform Proteinase K treatment to the sample prior to purification. See appropriate sample preparation protocol.			
DNA contamination	To remove DNA:			
	- Perform in-column DNase I treatment (page 12) or perform DNase I treatment post-purification, then re-purify the treated sample.			
	-For future preps, increase the volume of DNA/RNA Shield™ and/or DNA/RNA Lysis Buffer to ensure complete lysis and homogenization of the sample.			
RNA degradation	To prevent RNA degradation:			
	- Immediately collect and lyse fresh sample into DNA/RNA Shield™ and/or DNA/RNA Lysis Buffer ensure stability. Homogenized samples can be stored frozen for later processing.			

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

Notes



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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.

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