**One Call** 

Rapid Isolation of Total RNA from Plants and Animals Using the FastPrep<sup>®</sup> and FastPrep<sup>®</sup> -24 Instruments

One Source

Catalog # 6045-050 50 Preps

Storage: Refrigerated or ambient temperature  $(4^{\circ}C \text{ or } 15 - 30^{\circ}C)$ 

Revision # 6045-050-06APR

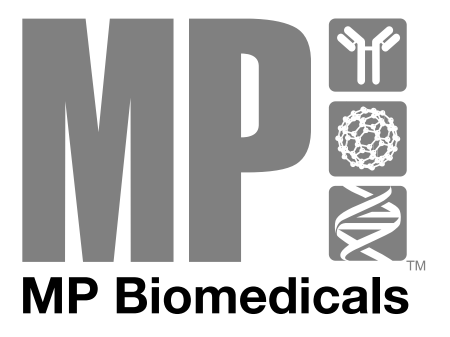
DO NOT expose RNApro<sup>™</sup> Solution to light for extended periods of time. Store in the original bottle in the closed kit box

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### I. Introduction to the FastRNA<sup>®</sup> Pro Green Kit and the FastPrep<sup>®</sup> Instruments

The FastRNA<sup>®</sup> Pro Green Kit is a single reagent extraction method designed to quickly and efficiently isolate total cellular RNA from plant and animal tissues. The RNA*pro*<sup>™</sup> Solution included in the kit is designed to efficiently inactivate cellular RNases during cell lysis to prevent RNA degradation. During use the RNA*pro*<sup>™</sup> Solution is mixed with the sample in a tube containing a specifically selected lysing matrix. The tube is then processed in the FastPrep<sup>®</sup> or FastPrep<sup>®</sup>-24 instrument for 40 seconds at a setting of 6.0 to release the total cellular RNA, DNA and proteins. Following the FastPrep<sup>®</sup> homogenization the RNA is purified and isolated by chloroform extraction and ethanol precipitation. The purified RNA is ready for downstream applications, including RT-PCR and northern analysis. The average RNA yield will vary depending on the source (e.g., seeds versus stems or liver versus brain) and the amount of tissue processed.

The FastPrep<sup>®</sup> and FastPrep<sup>®</sup>-24 Instruments are high-speed, benchtop devices that use a unique, optimized motion to homogenize samples by multidirectional, simultaneous impaction with lysing matrix particles. FastPrep<sup>®</sup> Instruments provide an extremely quick and highly reproducible homogenization that surpasses traditional lysis methods using enzyme digestion, sonication, blending, douncing and vortexing.

FastPrep<sup>®</sup> Instruments in combination with FastPrep<sup>®</sup> kits permit the release and purification of intact DNA, RNA and proteins from virtually any source, including plant samples (seeds, stems, pulp, old leaves, roots), animal samples (tail, liver, pancreas, brain, ear, cells, etc.), bacteria, yeast, spores, and more.

#### 2. Kit Components and User Supplied Materials

#### 2.1 FastRNA® Pro Green Kit Components

RNApro™ Solution	l × 55 milliliter bottle
DEPC-H <sub>2</sub> O	I x I5 milliliter bottle
Lysing Matrix D	$50 \times 2$ milliliter tubes
Short protocol	l each
User manual	l each
MSDS	l each
Certificate of Analysis	l each

#### 2.2 User Supplied Materials

FastPrep® or FastPrep® -24 Instrument (see section 10) Microcentrifuge Pipettmen RNase Erase®,Catalog # 2440-204, recommended Chloroform 100% ethanol 75% ethanol 1.5 or 2.0 ml RNase-free microcentrifuge tubes Agarose Gel loading dye and RNA size marker

### 3. Important Considerations before Use

The presence or introduction of RNase during the procedure may result in sample degradation. It is strongly recommended that the user minimize the potential for RNase contamination by using gloves throughout the procedure, using DEPC-H<sub>2</sub>O and by treating pipettmen, work area, gel box and gel comb with RNase Erase<sup>®</sup>. Additional RNA handling methods and precautions may be found in references I and 2.



The final sample volume after the addition of RNA*pro*<sup>TM</sup> Solution to the tube must maintain at least a  $\frac{1}{4}$  inch (5 mm) air space in the matrix tube during FastPrep<sup>®</sup> Instrument processing. Sample loss or tube failure may result from overfilling the matrix tube. The matrix tube caps must be secure, but not over-tightened, to prevent sample leakage. If the sample is too large for processing in a single tube, divide the sample and process using multiple tubes.

The average RNA yield will vary depending on the source (e.g., seeds versus stems or liver versus brain) and the amount of tissue processed. RNA yield from plant samples will also vary depending on the sample age (e.g., young versus mature leaves or root tip versus mature root). Standardizing and optimizing the amount of plant or animal tissue will result in reduced variability among operators and between experiments.

Confirm prior to centrifugation of the sample in the matrix sample tube that the matrix sample tubes spin freely and will not scrape the microcentrifuge wall during centrifugation. The use of other manufactured tubes in the FastPrep<sup>®</sup> Instruments is not recommended and may result in sample loss or instrument failure.

Add the RNApro<sup>TM</sup> Solution to the sample as soon as possible to initiate RNase inhibition. Samples, both FastPrep<sup>®</sup> Instrument homogenized and non-homogenized, are stable in RNApro<sup>TM</sup> Solution overnight at room temperature or  $4^{\circ}$ C.

Plant or animal cell variability in protein content and mucopolysaccharide composition may result in carryover into the aqueous solution following chloroform extraction. While this may not compromise downstream applications the user may adapt the protocol to include an additional chloroform (isoamyl alcohol may be included with the chloroform [CHCl3:IAA, 24:1, v:v]) extraction after Step 9 (Quick Protocol) or in step 11 (Detailed Protocol) to reduce the potential carryover.

A single 40 second run at a speed setting of 6.0 in the FastPrep<sup>®</sup> or FastPrep<sup>®</sup> 24 Instrument is sufficient to lyse most plant or animal samples. If the user experimentally determines that additional processing time is required, the sample should be incubated on ice in the Lysing Matrix tube for at least 2 minutes between successive FastPrep<sup>®</sup> Instrument homogenizations to prevent sample heating and possible RNA degradation.

The FastRNA® Pro Green Kit is designed to selectively purify total cellular RNA from DNA and protein. Experiments have indicated the RNA is sufficiently pure for use in RT-PCR and Northern analysis, however, it is recommended the user incorporate DNase I treatment of the RNA prior to use in applications where absolute control of DNA contamination is essential. Use DNase I at the concentration recommended by the manufacturer and incubate at 37°C for 30 minutes. The DNase I is inactivated by incubation at 75°C for 5 minutes or by addition of EDTA to 25 mM followed by phenol/chloroform extraction and precipitation.

### 4. Safety Precautions

The RNApro<sup>™</sup> Solution contains components that, when in contact with human tissue or during inhalation, may cause irritation or burning. Wear personal protective equipment to prevent skin contact (e.g., gloves, lab coat, and eye protection) and prevent inhalation of reagent vapors and consumption of liquid during use. Consult the enclosed Material Safety Data Sheet for additional details.

### 5. Quick Protocol for Experienced Users

- For each 100-300 mg sample to be processed, add 1 ml RNApro<sup>™</sup> Solution to a green-cap tube containing Lysing Matrix D.
- Add 100 300 mg plant or up to 100 mg animal tissue sample to the tube containing RNApro<sup>™</sup> Solution and Lysing Matrix D.



- 3. Process the tube in the FastPrep<sup>®</sup> or FastPrep<sup>®</sup> -24 Instrument for 40 seconds at a setting of 6.0.
- 4. Remove and centrifuge the tube at a minimum of 12,000  $\times$  g for 5 minutes at 4°C.
- 5. Transfer the upper phase (~ 750 µl) to a new microcentrifuge tube. Avoid transferring the debris pellet and lysing matrix.
- 6. Incubate the transferred sample 5 minutes at room temperature.
- Add 300 µl of chloroform (NO isoamyl alcohol). Vortex 10 seconds and incubate 5 minutes at room temperature.
- Centrifuge the tubes at a minimum of 12,000 x g for 5 minutes at 4°C.
- 9. Transfer the upper phase (without disturbing the interphase) to a new microcentrifuge tube.
- 10. Add 500  $\mu l$  of cold absolute ethanol; invert 5X to mix and store at -20°C for at least 30 minutes.
- 11. Centrifuge at a minimum of 12,000  $\times$  g for 15 minutes at 4°C and remove the supernatant.
- Wash the pellet with 500 µl of cold 75% ethanol (made with DEPC-H<sub>2</sub>O).
- 13. Remove the ethanol, air dry 5 minutes at room temperature (DO NOT completely dry the RNA) and resuspend the RNA in 100  $\mu$ l of DEPC-H<sub>2</sub>O.
- 14. Incubate 5 minutes at room temperature.
- 15. Determine the RNA concentration:

- a. Dilute 5  $\mu l$  of RNA into 495  $\mu l$  of DEPC-H  $_2O$
- b. Read the  $OD_{260}$  using DEPC-H<sub>2</sub>O as a blank
- c. Calculate the sample  $\mu$ g RNA per ml using the formula: (OD<sub>260</sub>)(40  $\mu$ g/ml/per OD)(100 [dilution factor]) =  $\mu$ g RNA per ml
- 16. Aliquot and store the RNA solution at -70°C.
- RNA integrity can be analyzed visually using denaturing or non-denaturing 1.2 % agarose gel electrophoresis (See Figure | & 2).

#### 6. Detailed Protocol

- For each 100-300 mg sample to be processed, add 1 ml RNApro<sup>™</sup> Solution to a green-cap tube containing Lysing Matrix D.
- Add 100 300 mg plant or up to 100 mg animal tissue sample to the tube containing RNApro<sup>™</sup> Solution and Lysing Matrix D.
- 3. Securely close the cap to prevent leakage in the next step. NOTE: The volume of the sample with 1 ml of RNApro<sup>™</sup> Solution must provide approximately ¼ inch (5 mm) airspace in the matrix tube to allow for effective homogenization and to prevent sample leakage and/or tube failure. DO NOT overfill the matrix tube. If the sample is too large for processing in a single tube, divide the sample and process using multiple tubes.
- 4. Process the sample tube in the FastPrep® or FastPrep® -24 Instrument for 40 seconds at a setting of 6.0. If the user experimentally determines that additional processing time is required, the sample should be incubated on ice in the Lysing Matrix tube for at least 2 minutes between successive FastPrep® Instrument homogenizations to prevent sample heating and possible RNA degradation.



- 5. Remove the sample tube and centrifuge at a minimum of 12,000 x g for 5 minutes at 4°C or room temperature.
- 6. Transfer the upper phase to a new microcentrifuge tube. Avoid transferring the debris pellet and lysing matrix.
- 7. Incubate the transferred sample 5 minutes at room temperature to increase RNA yield.
- Add 300 µl of chloroform (NO isoamyl alcohol). Vortex 10 seconds.
- 9. Incubate 5 minutes at room temperature to permit nucleoprotein dissociation and increase RNA purity.
- 10. Centrifuge the tubes at a minimum of 12,000  $\times$  g for 5 minutes at 4°C.
- 11. Transfer the upper phase to a new microcentrifuge tube without disturbing the interphase. If a portion of the interphase is transferred, repeat the centrifugation with the upper phase, and transfer the new upper phase to a clean microcentrifuge tube.

NOTE: Samples containing large amounts of cellular mucopolysaccharides can be re-extracted with chloroform (isoamyl alcohol may be included with the chloroform [CHCI3:IAA, 24:1, v:v]) to increase RNA purity. Alternatively, a lithium chloride precipitation may be used (see the Troubleshooting section).

- 12. Add 500  $\mu l$  of cold absolute ethanol to the sample; invert 5X to mix and store at -20°C for at least 30 minutes.
- 13. Centrifuge at a minimum of 12,000  $\times$  g for 15 minutes at 4°C and remove the supernatant. The RNA will appear as a white pellet in the tube. If the pellet is floating the sample may be recentrifuged to place the pellet at the tube bottom.

- Wash the pellet with 500 µl of cold 75% ethanol (made with DEPC-H<sub>2</sub>O).
- 15. Remove the ethanol, air dry 5 minutes at room temperature (DO NOT completely dry the RNA) and resuspend the RNA in 100  $\mu$ l of DEPC-H<sub>2</sub>O for short-term storage.

RNA resuspended in DEPC- $H_2O$  is generally stable for up to a year at -80°C. For longer term storage RNA samples may be stored at -20°C as ethanol precipitates. Ethanol precipitates must be pelleted and the RNA resuspended in aqueous solution prior to use.

NOTE: RNA does not evenly distribute in ethanol and can lead to inconsistent RNA amounts between samples when equal volumes are pipetted. In situations where precise amounts of RNA are required it is best to precipitate the total amount of RNA required, resuspend the RNA in DEPC-H<sub>2</sub>O and measure the concentration by OD<sub>260</sub> before proceeding.

- Incubate 5 minutes at room temperature to facilitate RNA resuspension.
- 17. Determine the RNA concentration:
  - a. Dilute 5  $\mu l$  of the purified RNA into 495  $\mu l$  of DEPC-H  $_2O$
  - b. Read the  $OD_{260}$  using DEPC-H<sub>2</sub>O as a blank
  - c. Calculate the sample  $\mu$ g RNA per ml using the formula: (OD<sub>260</sub>)(40  $\mu$ g/ml/per OD)(100 [dilution factor]) =  $\mu$ g RNA per ml

Spectrophotometer accuracy is greatest between ~ 0.2 and ~ 0.8. If the OD reading is below the range, add more RNA sample (e.g., 20  $\mu$ I RNA + 480  $\mu$ I DEPC-H<sub>2</sub>O) or concentrate the RNA by precipitation into a smaller volume. If the OD reading is above the range, use less RNA or dilute the sample prior to use.



18. Aliquot and store the RNA solution at  $-70\mu$ C.

19. The RNA integrity and an estimation of yield can be determined by analyzing a portion of the RNA sample using gel electrophoresis. Add I µg RNA in 9 µl DEPC-H<sub>2</sub>O, heat to 65°C for 5 minutes, add gel loading buffer and load the sample on a 1.2% agarose gel containing 2.2M formaldehyde in MOPS buffer. The sample is run at  $\sim$  80 volts for 30 minutes. Ethidium bromide may be added to the denatured RNA sample at 10 µg per milliliter prior to gel loading or the gel may be ethidium bromide stained and destained following electrophoresis and visualized under UV light. The quality of the RNA is determined by the appearance of ribosomal RNAs as sharp, distinct bands. Heterogeneous-sized messenger RNA may appear as diffuse ethidium staining between and below the ribosomal bands. Small RNA species such as tRNA and 5S RNA may be present in varying amounts at the dye front.

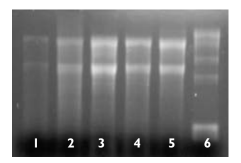
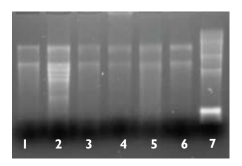


Figure 1:

Rat total RNA extracted with the FastRNA® Pro Green Kit. Approximately 2% of the total RNA isolated from 100 mg frozen tissue was loaded on to a 1.2% denaturing agarose gel (1XMOPS). Lane 1: tail; Lane 2: kidney; Lane 3: liver; Lane 4: ear; Lane 5: brain; Lane 6: 0.24-9.5kb RNA Ladder.



#### Figure 2:

Plant total RNA extracted with the FastRNA<sup>®</sup> Pro Green Kit. Approximately 2% of the total RNA isolated from 100 mg tissue was loaded on to a 1.2% denaturing agarose gel (1XMOPS). Lane 1: wheat seed; Lane 2: tomato leaf; Lane 3: red soybean seed; Lane 4: oat seed; Lane 5: iceplant leaf; Lane 6: barley seed; Lane 7: 0.24-9.5kb RNA Ladder.

### 7. Troubleshooting

#### 7.1 Degraded RNA or Lower than Expected RNA Yields

RNA purified using the FastRNA® Pro Green Kit and analyzed by denaturing or non-denaturing agarose gel electrophoresis will appear as 2 distinct ribosomal RNA (rRNA) bands of approximately equal fluorescent intensity using ethidium bromide staining. Messenger RNA (mRNA), which typically represents approximately less than 1% of the total cellular RNA and is heterogeneous length, will not be visible as distinct bands. rRNA is used as a marker to assess sample RNA degradation. Degraded RNA may appear as unequal fluorescent intensity between bands, a single band may be completely lacking or a heterogeneous fluorescent smear may appear below the rRNA bands or throughout the gel lane.

Recommended precautions include cleaning all instruments and work area with RNase Erase<sup>®</sup> (Catalog # 2440-204) prior to use. Use disposable sterile plastic containers when possible. Glassware should be thoroughly cleaned, rinsed with DEPC-H<sub>2</sub>O and baked at 250°C for 4 hours prior to use to remove RNase. Sterile, plugged micropipettes are recommended (see I, 2 for additional suggestions).

Certain plant or animal cells may contain elevated RNase levels. Reduce the exposure time to RNase by adding  $pro^{TM}$  Solution to each sample as soon as possible following sample harvest. Process fewer samples to shorten the time before complete cellular lysis and exposure to the RNase inactivating activity of  $pro^{TM}$  Solution.

Plant or animal samples stored for extended duration at room temperature, frozen or refrigerated for extended periods will contribute to reduced RNA yield and integrity. RNApro<sup>TM</sup> Solution can permeate samples and will protect RNA from degradation for at least 24 hours before it is processed in the FastPrep<sup>®</sup> Instrument. However, higher yields of RNA will always result when samples are homogenized immediately after addition of RNA**pro**<sup>TM</sup> Solution.

Artifactual RNA degradation may occasionally occur during gel electrophoresis due to a gel that was not RNase free, running the gel at too high voltage or from using depleted running buffer. Rerun the samples with a known intact RNA sample using freshly prepared reagents.

RNA degradation may occur due to RNase contamination introduced into the DEPC-H<sub>2</sub>O following use. Prepare fresh DEPC-H<sub>2</sub>O in an RNase free container (1,2). RNApro<sup>TM</sup> Solution contains RNase inactivating components and will not become contaminated during use.

#### 7.2 No Pellet after Ethanol Precipitation

The purified RNA may not appear as a pellet but may instead adhere to the side of the tube. The RNA may not be visible and it may appear that RNA has not been purified. Complete the RNA purification and confirm the RNA concentration by  $OD_{260}$  and integrity by gel electrophoresis. Adhering to the tube wall will not affect the RNA purity, size or use in subsequent applications.

The pellet may not be firmly attached to the side of the tube and may be observed floating in the solution or at the solution surface. Recentrifuge the sample in the same tube and exercise caution to not lose the pellet when removing the supernatant.

Confirm the correct amount of plant or animal tissue was used to isolate RNA. Total RNA yield will vary with the sample age, source (e.g., root versus pulp) and storage conditions. In a controlled experiment, titrate the RNA yield from increasing amounts of sample. If using mature plant tissue, perform the control experiment of processing an immature, rapidly growing



portion of the plant, which in general has greater amounts of RNA available for isolation. If limited amounts of sample are available or the RNA yield is low, the DEPC-H<sub>2</sub>O resuspension volume may be reduced to 50 or 25  $\mu$ l to concentrate the RNA (Quick Protocol step 13 and Detailed Protocol step 15).

#### 7.3 Genomic DNA Contamination

Genomic DNA contamination will appear as a high molecular weight smear on a denaturing gel or as ethidium bromide stained material in the gel loading well. In the event genomic DNA contamination occurs, re-extract the RNA sample with chloroform or chloroform:isoamyl alcohol (24:1, v:v). The lower phase of the chloroform extraction contains genomic DNA and should be carefully avoided when removing the top RNA containing phase. Leaving a small volume of the top phase in the tube will prevent accidental DNA contamination.

#### 7.4 Mucopolysaccharide/ Carbohydrate Contamination

Samples containing large amounts of cellular mucopolysaccharides can be re-extracted after the initial chloroform extraction with a second chloroform extraction. Isoamyl alcohol may be included with the chloroform [CHCl3:IAA, 24:I, v:v]) to increase RNA purity. Refer also to Lithium Chloride Precipitation in the Troubleshooting section.

#### 7.5 Lithium Chloride Precipitation

Lithium chloride (LiCl) may be used to precipitate RNA while excluding carbohydrate, DNA, proteins and transcription inhibitors. Lithium chloride has historically been used to precipitate RNA greater than  $\sim$  300 nucleotides from tRNA and 5S RNA. Lithium chloride precipitation may be incorporated into the FastRNA®

Pro Green Kit procedure. Following ethanol precipitation of the RNA and resuspension in 100  $\mu$ I DEPC-H<sub>2</sub>O, add lithium chloride to a final concentration of 2 – 3 M (e.g., 0.2 volumes [20  $\mu$ I] RNase free 8 M lithium chloride). Add 2.5 volumes RNase free absolute ethanol (250  $\mu$ I). Mix the solution and store on ice at least 2 hours. Centrifuge for 15 minutes at minimum of 12,000 rpm at 4°C. Remove the supernatant and wash the pellet with 75% cold RNase free ethanol. The ethanol wash step is critical to prevent LiCl inhibition of cell-free translation and in vitro transcription. Air dry and resuspend the RNA in 100  $\mu$ I DEPC-H<sub>2</sub>O.

### 8. Recommended Reference Format for Publications

Total RNA was isolated from \_\_\_\_\_ mg of \_\_\_\_\_cells using the FastRNA® Pro Green Kit (MP Biomedicals, Irvine CA) and the FastPrep® -24 Instrument (MP Biomedicals, Irvine CA). Samples have been homogenized for \_\_\_\_\_ seconds at a speed setting of \_\_\_\_\_.

### 9. References

1. Molecular Cloning, Sambrook and Russell. Cold Spring Harbor Laboratory Press, 3rd Edition, 2001.

2. Current Protocols in Molecular Biology, John Wiley & Sons, Inc., 2002, www.currentprotocols.com.

### 10. Related Products

Description	Size	Catalog #
FastPrep <sup>®</sup> -24 Instrument	100-230V	6002-500
FastPrep® FP100A Instrument	100V	6001-100
FastPrep <sup>®</sup> FP120A Instrument	120V	6001-120
FastPrep <sup>®</sup> FP220A Instrument	220V	6001-220
FastRNA® Pro Red Kit (Yeast)	50 preps	6035-050
FastRNA® Pro Blue Kit (Bacteria)	50 preps	6025-050
FastDNA® Kit	100preps	6540-400
FastDNA® SPIN Kit	100 preps	6540-600
FastDNA® SPIN Kit for Soil	50 preps	6560-200
FastPROTEIN™ Blue Matrix	50 preps	6550-400
FastPROTEIN™ Red Matrix	50 preps	6550-600
RNase Erase®	500 ml	2440-204
Lysing Matrix D	50 x 2ml tubes	6913-050
Lysing Matrix D	100 x 2ml tubes	6913-100
Lysing Matrix D	500 x 2ml tubes	6913-500

#### II. Product Use Limitation & Warranty

The products presented in this instruction manual are for research or manufacturing use only. They are not to be used as drugs or medical devices in order to diagnose, cure, mitigate, treat or prevent diseases in humans or animals, either as part of an accepted course of therapy or in experimental clinical investigation. These products are not to be used as food, food additives or general household items. Purchase of MP Biomedicals products does not grant rights to reproduce, modify, or repackage the products or any derivative thereof to third parties. MP Biomedicals makes no warranty of any kind, expressed or implied, including merchantability or fitness for any particular purpose, except that the products sold will meet our specifications at the time of delivery. Buyer's exclusive remedy and the sole liability of MP Biomedicals hereunder shall be limited to, at our discretion, no replacement or compensation, product credits, refund of the purchase price of, or the replacement of materials that do not meet our specification. By acceptance of the product, Buyer indemnifies and holds MP Biomedicals harmless against, and assumes all liability for, the consequence of its use or misuse by the Buyer, its employees or others, including, but not limited to, the cost of handling. Said refund or replacement is conditioned on Buyer notifying MP Biomedicals. within thirty (30) days of receipt of product. Failure of Buyer to give said notice within thirty (30) days shall constitute a waiver by the Buyer of all claims hereunder with respect to said material(s).

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## Instruction Manual

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