applied biosystems

MagMAX™ *mir*Vana™ Total RNA Isolation Kit

High-throughput isolation of RNA (including small RNA) from tissue samples

Catalog Number A27828

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WARNING! Read the Safety Data Sheets (SDSs) and follow the handling instructions. Wear appropriate protective eyewear, clothing, and gloves. Safety Data Sheets (SDSs) are available from **thermofisher.com/support**.

Product description

The MagMAX[™] *mir* Vana[™] Total RNA Isolation Kit is designed for isolation of total RNA, including microRNA, from a wide variety of sample matrices. The kit uses MagMAX[™] magnetic-bead technology, ensuring reproducible recovery of high-quality RNA that is suitable for a broad range of applications, including TaqMan[®] miRNA Detection Assays.

Kit contents and storage

Table 1 MagMAX™ *mir*Vana™ Total RNA Isolation Kit (Cat. no. A27828, 96 reactions)

Contents	Amount	Storage
	Aillouilt	Storage
Box 1 of 2		
Proteinase K ^[1] , 50 mg/mL	0.48 mL	
Lysis/Binding Enhancer	0.96 mL	-25°C to -15°C
TURBO DNase™, 20 U/µL	0.2 mL	
Box 2 of 2		
Lysis Buffer	115 mL	
PK Digestion Buffer ^[1]	4.4 mL	
RNA Binding Beads ^[2]	2 mL	
Wash Solution 1 Concentrate ^[3]	20 mL	
Wash Solution 2 Concentrate ^[3]	60 mL	
Rebinding Buffer	4.8 mL	15°C to 30°C
MagMAX™ TURBO DNase™ Buffer	4.6 mL	
Elution Buffer	9.6 mL	
Processing Plate ^[1]	1	
Elution Plates	2	
Plate Covers	4	

^[1] Not used for RNA isolation from tissue samples.

Materials required but not supplied

Unless otherwise indicated, all materials are available from Life Technologies (**thermofisher.com**). MLS: Fisher Scientific (**fisherscientific.com**) or other major laboratory supplier.

Item	Source	
Magnetic particle processor, one of the foll	owing:	
KingFisher™ Flex Magnetic Particle Processor 96DW ^[1]	Cat. No. 5400630	
MagMAX™ Express-96 Deep Well Magnetic Particle Processor	_[2]	
KingFisher™ Duo Prime Magnetic Particle Processor ^[1]	Cat. No. 5400110	
Other equipment		
Thermo Scientific™ Compact Digital Microplate Shaker	Fisher Scientific 11-676-337	
Fisher Scientific™ Analog Vortex Mixer	Fisher Scientific 02-215-365	
PRO250 Homogenizer with 7 × 95 mm Saw Tooth Bottom Generator Probe	PRO Scientific 01-01250 and 02-07095	
Heating block, 37°C	MLS	
Adjustable micropipettors	MLS	
Multi-channel micropipettors	MLS	
Plates and combs ^[3]		
Deep Well Plates, one of the following:		
MagMAX™ Express-96 Deep Well Plates	Cat. No. 4388476	
KingFisher™ Flex Microtiter Deepwell 96 Plate, Sterile	Cat. No. 95040460	
Standard Well Plates, one of the following:		
MagMAX™ Express-96 Standard Plates	Cat. No. 4388475	
KingFisher™ 96 KF Microplate	Cat. No. 97002540	
One of the following tip combs, depending on the magnetic particle processor used:		
MagMAX™ Express-96 Deep Well Tip Combs	Cat. No. 4388487	
KingFisher™ 96 Tip Comb for DW Magnets	Cat. No. 97002534	
KingFisher™ Duo 12-Tip Comb, for Microtiter 96 Deepwell plate	Cat. No. 97003500	
Other consumables		
MicroAmp™ Clear Adhesive Film	Cat. No. 4306311	
Nonstick, RNase-free Microfuge Tubes (1.5 mL)	Cat. No. AM12450	
Nonstick, RNase-free Microfuge Tubes (2.0 mL)	Cat. No. AM12475	
5–mL Culture tubes	MLS	
Conical tubes (15 mL)	Cat. No. AM12500	
Aerosol-resistant pipette tips	MLS	
Reagent reservoirs	MLS	



^[2] Do not freeze the RNA Binding Beads.

^[3] Final volume; see "Before first use: prepare Wash Solutions" on page 2.

Item	Source
Reagents	
Isopropanol, 100% (molecular grade or higher)	MLS
Ethanol, 200 proof (absolute)	MLS
2-Mercaptoethanol	MLS
(Optional) Chloroform	MLS

 $^{^{[1]}}$ See "If needed, download the KingFisher" Flex or Duo program" on page 2

Sample collection and storage

- Process tissue immediately, store them in RNAlater[™] Stabilization Solution, or freeze them in liquid nitrogen and store at -80°C.
- For ease of processing, we recommend pre-weighing and storing the tissue in pieces of incremental sizes, according to the following guidelines:
 - Increments of ≤50 mg for tissues containing low or normal level of cellular RNase.
 - Increments of ≤30 mg for tissues containing high level of cellular RNase, such as spleen or pancreas.

Important procedural guidelines

- Perform all steps at room temperature (20–25°C) unless otherwise noted.
- When mixing samples by pipetting up and down, avoid creating bubbles.
- Cover the plate during the incubation and shaking steps to prevent spill-over and cross-contamination. The same Plate Cover can be used throughout the procedure, unless it becomes contaminated.
- If you use a titer plate shaker other than the Thermo Scientific
 Compact Digital Microplate Shaker, verify that:
 - The plate fits securely on your titer plate shaker.
 - The recommended speeds are compatible with your titer plate shaker. Ideal speeds should allow for thorough mixing without splashing.
- Volumes for reagent mixes are given per well. We recommend that you prepare master mixes for larger sample numbers. To calculate volumes for master mixes, refer to the per-well volume and add 5% overage.
- Lysed samples can be stored in Lysis Binding Mix at -20°C for up to 4 days before adding the Binding Beads Mix. Thaw frozen samples to room temperatures before use.

If needed, download the KingFisher™ Flex or Duo program

The program required for this protocol is not pre-installed on the KingFisher[™] Flex Magnetic Particle Processor 96DW or on the KingFisher[™] Duo Prime Magnetic Particle Processor.

- On the MagMAX[™] mirVana[™] Total RNA Isolation Kit web page, scroll down to the **Product Literature** section.
- 2. Right-click on the appropriate program for your instrument:
 - A27828_FLEX_Tissue_Cells for KingFisher[™] Flex Magnetic Particle Processor 96DW.
 - A27828_DUO_Tissue_cells for KingFisher[™] Duo Prime Magnetic Particle Processor.
- 3. select Save as Target to download to your computer.
- **4.** Refer to the manufacturer's documentation for instructions for installing the program on the instrument.

Before first use: prepare Wash Solutions

Prepare the Wash Solutions from the concentrates:

- Add 10 mL of isopropanol to Wash Solution 1 Concentrate, mix, and store at room temperature.
- Add 48 mL of ethanol to Wash Solution 2 Concentrate, mix, and store at room temperature.

Before each use: prepare TURBO DNase™ Solution and Binding Beads Mix

 Prepare the TURBO DNase[™] Solution as indicated in the following table, mix, and store on ice until use.

Component	Volume per well
MagMAX™ TURBO DNase™ Buffer	48 μL
TURB0 DNase™	2 μL
Total TURBO DNase™ Solution	50 μL

 Prepare the Binding Beads Mix as indicated in the following table, mix, and store on ice until use.

Component	Volume per well
RNA Binding Beads	10 μL
Lysis/Binding Enhancer	10 μL
Total Binding Beads Mix	20 μL

^[2] Not available for sale.

^[3] KingFisher™ Duo Combi Pack (Cat. no. 97003530) includes plates and combs for the KingFisher™ Duo Prime Magnetic Particle Processor.

Perform RNA extraction from tissues samples

Lyse the samples

Lyse the tissue samples

- a. Determine the size of tissue (in mg) to be homogenized.
- b. Determine the amount of Lysis Binding Mix that is needed to homogenize the tissue, according to the following ratios:
 - For tissues containing low to normal levels of cellular RNase (for example, brain, heart, or liver), use $20 \mu L$ of Lysis Binding Mix for 1 mg of tissue (1:20 ratio).

For example, use 200 µL of Lysis Binding Mix for 10 mg of tissue.

• For tissues containing high levels of cellular RNase (for example, spleen or pancreas), use 40 μ L of Lysis Binding Mix for 1 mg of tissue (1:40 ratio).

For example, use 400 µL of Lysis Binding Mix for 10 mg of tissue.

Note: Most mechanical homogenizers require a minimum volume of 200 μ L. Therefore, we recommend processing no less than 10 mg when homogenizing.

c. Prepare sufficient Lysis Binding Mix, according to the following table.

Component	Volume for 100 µL of Lysis Binding Mix
Lysis Buffer	100 μL
2-Mercaptoethanol	0.7 μL
Total Lysis Binding Mix	~100 µL

- d. Add tissue to the prepared Lysis Binding Mix.
- e. Homogenize the tissue sample using standard homogenization procedures.

IMPORTANT! Make sure that the tissue homogenization is complete to ensure maximal RNA recovery. For the tissue input amounts listed previously, we recommend homogenizing for 30 seconds to fully break up the sample and ensure maximal RNA recovery and quality.

Proceed to the appropriate procedures:

- "Isolate RNA using the MagMAX™ Express-96 Deep Well Magnetic Particle Processor or the KingFisher™ Flex Magnetic Particle Processor 96DW".
- "Isolate RNA using the KingFisher™ Duo Prime Magnetic Particle Processor".

Isolate RNA using the MagMAX™ Express-96 Deep Well Magnetic Particle Processor or the KingFisher™ Flex Magnetic Particle Processor 96DW

Bind the RNA to the RNA Binding Beads If samples were frozen at the previous step, thaw them completely to room temperature before proceeding.

a. Vortex the lysates and transfer 100 μL to a separate well in a MagMAX Express-96 Deep Well Plate.

Table 2 Recommended tissue mass per 100 µL of Lysis Binding Mix

	Levels of cellular RNase		
Samples	Low to normal (for example, liver)	High (for example, spleen)	
Frozen tissue	up to 10 mg	up to 2.5 mg	
Tissue stored in RNA <i>later</i> ™ Stabilization Solution	up to 5 mg	up to 2.5 mg	

Note: Most mechanical homogenizers require a minimum of 200 μ L. Therefore, we recommend processing no less than 10 mg of tissue when homogenizing.

b. (Optional) Add $10 \mu L$ of chloroform to each well.

IMPORTANT! The addition of chloroform is required for samples containing high level of RNase (spleen, pancreas).

c. Cover the plate and shake as indicated.

Time	Speed
5 minutes	1150 rpm (Speed 10) ^[1]

 $^{^{[1]}}$ Setting for Lab-Line $^{\!\scriptscriptstyle\mathsf{M}}$ shaker.

During the agitation, proceed to "Set up the processing plates" on page 4.

d. Add 100 µL of isopropanol to each sample, cover the plate, and shake as indicated.

Time	Speed
2 minutes	950 rpm (Speed 7) ^[1]

^[1] Setting for Lab-Line $^{\text{\tiny{M}}}$ shaker.

Bind the RNA to the **RNA Binding Beads** (continued)

e. Add 20 µL of the prepared Binding Beads Mix to each sample and shake as indicated.

Time	Speed
5 minutes	950 rpm (Speed 7) ^[1]

 $^{^{[1]}}$ Setting for Lab-Line $^{\text{\tiny{M}}}$ shaker.

Set up the processing plates While the samples are incubating, set up the Wash, DNase, Elution, and Tip Comb Plates outside the instrument as described in the following table.

Table 3 Processing plates

Plate ID	Plate position ^[1]	Plate type	Reagent	Volume per well
Wash Plate 1	2	Standard	Wash Solution 1	150 μL
Wash Plate 2	3	Standard	Wash Solution 2	150 μL
DNase Plate ^[2]	4	Standard	TURBO DNase™ Solution	50 μL
Wash Plate 3	5	Standard	Wash Solution 2	150 μL
Wash Plate 4	6	Standard	Wash Solution 2	150 μL
Elution Plate	7	Standard	Elution Buffer	50-100 μL ^[3]
Tip Comb	8	Deep Well or standard	Place a MagMAX™ Express-96 Deep Well Tip Comb in a MagMAX™ Express-96 Deep Well Plate or in a MagMAX™ Express-96 Standard Plate.	

^[1] Position on the instrument

RNA

- Wash, rebind, and elute the a. Ensure that the instrument is set up for processing with the deep well magnetic head and select the program on the instrument.
 - AM1830DW on MagMAX[™] Express-96 Deep Well Magnetic Particle Processor
 - A27828_FLEX_Tissue_Cells on KingFisher[™] Flex Magnetic Particle Processor
 - b. Start the run and load the prepared processing plates in their positions when prompted by the instrument (see Table 3).
 - c. Load the sample plate (containing lysate, isopropanol, and Binding Beads Mix) at position 1 when prompted by the instrument.
 - **d.** When prompted by the instrument (30–35 minutes after the initial start):
 - 1. Remove the DNase Plate from the instrument.
 - 2. Add 50 μ L of Rebinding Buffer and 100 μ L of isopropanol to each sample well. Add Rebinding Buffer and isopropanol immediately after the prompt, to prevent excessive drying of any beads that are still captured on the Tip Comb.

IMPORTANT! Do not pre-mix the Rebinding Buffer and isopropanol. Add them separately to the samples.

- 3. Load the DNase Plate back onto the instrument, and press Start.
- e. At the end of the run (approximately 45 minutes after the initial start), remove the Elution Plate from the instrument and seal immediately with a new MicroAmp[™] Clear Adhesive Film.
 - (*Optional*) Eluates can be transferred to a storage plate after collection.
 - If excess bead residue is seen in the wells, place the Elution Plate on the Magnetic Stand-96 to capture any residue prior to downstream use of the RNA.

IMPORTANT! Do not allow the purified samples to sit uncovered at room temperature for more than 10 minutes, to prevent evaporation and contamination.

The purified samples are ready for immediate use. Alternatively, store the covered Elution Plate:

- On ice for up to 8 hours.
- At -20°C or -80°C for long-term storage.

f. Proceed immediately to "Wash, rebind, and elute the RNA" on page 4.

 $^{^{[2]}}$ The instrument prompts the user to add 50 μ L of Rebinding Buffer and 100 μ L of isopropanol to the DNase Plate after the DNase treatment step.

Use 50 μ L for lower-yielding samples such as brain or heart or 100 μ L for higher-yielding samples such as liver or spleen.

Isolate RNA using the KingFisher™ Duo Prime Magnetic Particle Processor

Bind the RNA to the RNA Binding Beads

If samples were frozen at the previous step, thaw them completely to room temperature before proceeding.

a. Vortex the lysates and transfer 100 μL to a separate well of row H of a MagMAX^{$^{\text{IM}}$} Express-96 Deep Well Plate.

Table 4 Recommended tissue mass per 100 µL of Lysis Binding Mix

	Levels of cellular RNase	
Samples	Low to normal (for example, liver)	High (for example, spleen)
Frozen tissue	up to 10 mg	up to 2.5 mg
Tissue stored in RNA <i>later</i> ™ Stabilization Solution	up to 5 mg	up to 2.5 mg

Note: Most mechanical homogenizers require a minimum of 200 μ L. Therefore, we recommend processing no less than 10 mg of tissue when homogenizing.

b. (Optional) Add 10 µL of chloroform to each well.

IMPORTANT! The addition of chloroform is required for samples containing high level of RNase (spleen, pancreas).

c. Cover the plate and shake as indicated.

Time	Speed	
5 minutes	1150 rpm (Speed 10) ^[1]	

^[1] Setting for Lab-Line™ shaker.

d. Add 100 μL of isopropanol to each sample, cover the plate, and shake as indicated.

Time	Speed	
2 minutes	950 rpm (Speed 7) ^[1]	

^[1] Setting for Lab-Line™ shaker.

e. Add 20 µL of the prepared Binding Beads Mix to each sample and shake as indicated.

Time	Speed	
5 minutes	950 rpm (Speed 7) ^[1]	

^[1] Setting for Lab-Line™ shaker.

Set up the processing plate Add processing reagents as indicated in the following table.

Table 5 Volume of processing reagents and plate location

Row ID	Plate row ^[1]	Reagent	Volume per well
Elution	A	Elution Buffer	50-100 μL ^[2]
Tip Comb	В	Place a KingFisher™ Duo 12-Tip Comb in Row B.	
Wash 4	С	Wash Solution 2	150 μL
Wash 3	D	Wash Solution 2	150 μL
DNase ^[3]	Е	TURBO DNase™ Solution	50 μL
Wash 2	F	Wash Solution 2	150 μL
Wash 1	G	Wash Solution 1	150 μL

^[1] Row on the MagMAX™ Express-96 Deep Well Plate.

Wash, rebind, and elute the RNA

- a. Ensure that the instrument is set up for processing with the deep well 96–well plates and select the program A27828_DUO_Tissue_cells on the instrument.
- b. Start the run and load the prepared processing plate when prompted by the instrument (see "Wash, rebind, and elute the RNA" on page 5).
- ${\bf c.}~$ When prompted by the instrument (30–35 minutes after the initial start):
 - 1. Remove the plate from the instrument.
 - 2. Add 50 μ L of Rebinding Buffer and 100 μ L of isopropanol to each sample well in Row E. Add Rebinding Buffer and isopropanol immediately after the prompt, to prevent excessive drying of any beads that are still captured on the Tip Comb.

IMPORTANT! Do not pre-mix the Rebinding Buffer and isopropanol. Add them separately to the samples.

3. Load the plate back onto the instrument, and press Start.

 $^{^{[2]}}$ Use 50 μ L for lower-yielding samples such as brain or heart or 100 μ L for higher-yielding samples such as liver or spleen.

 $^{^{[3]}}$ The instrument prompts the user to add 50 μ L of Rebinding Buffer and 100 μ L of isopropanol to the DNase Plate after the DNase treatment step.

Wash, rebind, and elute the RNA (continued)

- **d.** At the end of the run (approximately 45 minutes after the initial start), remove the Elution Plate from the instrument and transfer the eluted RNA (Row A) to an Elution Plate.
- e. Seal immediately with a new MicroAmp[™] Clear Adhesive Film.

IMPORTANT! Do not allow the purified samples to sit uncovered at room temperature for more than 10 minutes, to prevent evaporation and contamination.

The purified samples are ready for immediate use. Alternatively, store the covered Elution Plate:

- On ice for up to 8 hours.
- At -20°C or -80°C for long-term storage.

Limited product warranty

Life Technologies Corporation and/or its affiliate(s) warrant their products as set forth in the Life Technologies' General Terms and Conditions of Sale at www.thermofisher.com/us/en/home/global/terms-and-conditions.html. If you have any questions, please contact Life Technologies at www.thermofisher.com/support.

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