# **applied**biosystems

# MagMAX<sup>™</sup> Viral/Pathogen II Nucleic Acid Isolation Kit INSTRUCTIONS FOR USE

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The customer is responsible for compliance with regulatory requirements that pertain to their procedures and uses of the instrument. The information in this guide is subject to change without notice.

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Revision	Date	Description
B.0	27 May 2021	<ul> <li>Added Cat. No. A5400630 for the KingFisher<sup>™</sup> Flex Magnetic Particle Processor with 96 Deep-Well Head.</li> </ul>
		Added stability information.
		(400-µL sample input volume only) For the Proteinase K digestion procedure, changed the incubation time to 10 minutes.
A.0	20 November 2020	New document.

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#### Intended use

The MagMAX<sup>™</sup> Viral/Pathogen II Nucleic Acid Isolation Kit is a magnetic bead technology-based nucleic acid purification kit intended for the isolation and purification of viral RNA and DNA from human nasopharyngeal swabs.

The kit is intended for use by qualified and trained clinical laboratory personnel specifically instructed and trained in the techniques of magnetic bead purification, either manual or automated, and *in vitro* diagnostic procedures.

#### **Product information**

The MagMAX<sup>™</sup> Viral/Pathogen II Nucleic Acid Isolation Kit (Cat. No. A48383) is specifically designed to recover RNA and DNA from viral particles contained in transport medium. The kit uses MagMAX<sup>™</sup> magnetic-bead technology to provide reproducible recovery of high-quality nucleic acid.

This product is intended for *in-vitro* diagnostic use and includes the following features:

- Automated workflow using the KingFisher<sup>™</sup> Flex Magnetic Particle Processor with 96 Deep-Well Head allows for 96 nasopharyngeal swab specimens to be processed in <30 minutes</li>
- · Protocol options support an automated workflow or a manual workflow
- Flexible protocol accommodates sample volume inputs from 200 to 400 µL of transport medium
- No need for carrier RNA
- Elution volume of 50 μL

## Contents and storage

The MagMAX $^{\text{\tiny M}}$  Viral/Pathogen II Nucleic Acid Isolation Kit contains sufficient reagents for 1,000 reactions with 400  $\mu$ L volume input or 2,000 reactions with 200  $\mu$ L volume input.

Component	Amount	Storage
Binding Solution	550 mL	
Wash Solution	1,000 mL	
Binding Beads	20 mL	15°C to 25°C
Proteinase K	10 mL	
Elution Buffer	100 mL	

## Required materials not supplied

Unless otherwise indicated, all materials are available through **thermofisher.com**. "MLS" indicates that the material is available from **fisherscientific.com** or another major laboratory supplier.

Catalog numbers that appear as links open the web pages for those products.

Item	Source
Automated nucleic acid extraction system and materials	
KingFisher <sup>™</sup> Flex Magnetic Particle Processor with 96 Deep-Well Head	5400630, A5400630
KingFisher <sup>™</sup> Flex 96 Deep-Well Heating Block	24075430
KingFisher <sup>™</sup> 96 Deep-Well Plate	95040450, A48305, <b>A48424</b> , 95040455
<ul> <li>96-well plate for the tip comb, one of the following:</li> <li>KingFisher<sup>™</sup> 96 KF microplate</li> <li>Tip Comb Presenting Plate for KF 96</li> <li>Nunc<sup>™</sup> MicroWell<sup>™</sup> 96-Well Microplate, Flat Bottom</li> <li>Nunc<sup>™</sup> MicroWell<sup>™</sup> 96-Well Microplate, barcoded</li> <li>ABgene<sup>™</sup> 96-Well Polypropylene Storage Microplate</li> <li>ABgene<sup>™</sup> 96-Well 1.2-mL Polypropylene Deepwell Storage Plate</li> <li>Nunc<sup>™</sup> F96 MicroWell<sup>™</sup> Black Polystyrene Plate</li> <li>Nunc<sup>™</sup> F96 MicroWell<sup>™</sup> White Polystyrene Plate</li> <li>KingFisher<sup>™</sup> 96 Deep-Well Plate</li> </ul>	<ul> <li>97002540</li> <li>267600</li> <li>167008</li> <li>269787</li> <li>AB0796</li> <li>AB1127</li> <li>137101</li> <li>136101</li> <li>95040450, A48305, A48424, 95040455</li> </ul>
KingFisher <sup>™</sup> 96 tip comb for DW magnets	97002534, A48438, A48414

#### (continued)

Item	Source		
Manual nucleic acid extraction system and materials			
Magnetic Stand-96	AM10027, AM10050		
Compact Digital Microplate Shaker	88882005		
Incubator capable of reaching 65°C with slatted shelves	MLS		
KingFisher <sup>™</sup> 96 Deep-Well Plate	95040450, A48305, A48424, 95040455		
<ul> <li>Standard 96-well plate for the eluate, one of the following:</li> <li>KingFisher<sup>™</sup> 96 KF microplate</li> <li>MicroAmp<sup>™</sup> Fast Optical 96-Well Reaction Plate with Barcode, 0.1 mL</li> <li>MicroAmp<sup>™</sup> Fast Optical 96-Well Reaction Plate, 0.1 mL</li> <li>MicroAmp<sup>™</sup> Optical 96-Well Reaction Plate with Barcode, 0.2 mL</li> <li>MicroAmp<sup>™</sup> Optical 96-Well Reaction Plate, 0.2 mL</li> </ul>	<ul> <li>97002540</li> <li>4346906, 4366932</li> <li>4346907</li> <li>4306737, 4326659</li> <li>N8010560, 4316813</li> </ul>		
MicroAmp <sup>™</sup> Clear Adhesive Film	4306311		
Equipment			
Laboratory mixer, vortex or equivalent	MLS		
Single and multichannel adjustable pipettors (1.00 μL to 1,000.0 μL)	MLS		
Cold block or ice	MLS		
Reagents			
Fisher BioReagents <sup>™</sup> Ethanol, Absolute, Molecular Biology Grade <sup>[1]</sup> , or equivalent	BP2818100, BP2818500, BP28184		
Nuclease-free Water (not DEPC-Treated)	MLS		
(Optional) Extraction control, if required for your assay	See the assay guide for more information		
Tubes, plates, and other consumables			
MicroAmp <sup>™</sup> Clear Adhesive Film	4306311		
MicroAmp <sup>™</sup> Adhesive Film Applicator	4333183		
Sterile conical tubes for reagent preparation	MLS		
Sterile aerosol barrier (filtered) pipette tips	thermofisher.com/pipettetips		

<sup>[1]</sup> Available at fisherscientific.com.



# Extract nucleic acid from swabbased samples (automated method)

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Automated nucleic acid extraction is performed with the KingFisher<sup>™</sup> Flex Magnetic Particle Processor with 96 Deep-Well Head using a sample volume input of 200 µL or 400 µL. See your assay documentation for specific sample volume input recommendations.

#### Before you begin

- Ensure that you read and understand the information provided in this guide before you begin the extraction procedure.
- Review your assay documentation to determine if an extraction control is recommended to verify
  the efficacy of the nucleic acid preparation. Follow the extraction control guidelines provided in the
  assay documentation.
- Determine the number of required reactions based on the number of patient samples to be processed, plus one Negative Control per plate.
- Prepare fresh 80% Ethanol using Ethanol, Absolute, Molecular Biology Grade and Nuclease-free Water (not DEPC-Treated) for the required number of reactions, sufficient for 1 mL per reaction, plus 10% overage.
- Label the short side of each KingFisher<sup>™</sup> 96 Deep-Well Plate (4):

Label	Number of plates
Sample plate	1
Wash 1	1
Wash 2	1
Elution plate	1

Label the short side of the KingFisher<sup>™</sup> 96 KF microplate (1):

Label	Number of plates
Tip comb	1

**Note:** The following items can be used to hold the tip comb instead of the KingFisher<sup>™</sup> 96 KF microplate:

- Tip Comb Presenting Plate for KF 96
- Nunc<sup>™</sup> MicroWell<sup>™</sup> 96-Well Microplate, Flat Bottom
- Nunc<sup>™</sup> MicroWell<sup>™</sup> 96-Well Microplate, barcoded
- ABgene<sup>™</sup> 96–Well Polypropylene Storage Microplate
- ABgene<sup>™</sup> 96-Well 1.2-mL Polypropylene Deepwell Storage Plate
- Nunc<sup>™</sup> F96 MicroWell<sup>™</sup> Black Polystyrene Plate
- Nunc<sup>™</sup> F96 MicroWell<sup>™</sup> White Polystyrene Plate
- KingFisher<sup>™</sup> 96 Deep-Well Plate
- Mark the Negative Control well on the plate.

# Extract nucleic acid—Automated method (200-µL sample input volume)

#### Set up the instrument (200-µL sample input volume)

1. Ensure that the KingFisher<sup>™</sup> Flex Magnetic Particle Processor with 96 Deep-Well Head is set up with the KingFisher<sup>™</sup> Flex 96 Deep-Well Heating Block.

**IMPORTANT!** Failure to use the proper magnetic head and heat block results in lower yields and potential harm to the instrument.

2. Ensure that the MVP\_2Wash\_200\_Flex program has been downloaded from the MagMAX<sup>™</sup> Viral/Pathogen II Nucleic Acid Isolation Kit product page at www.thermofisher.com and loaded onto the instrument.

#### 2

#### Prepare the processing plates (200-µL sample input volume)

**Note:** During the wash steps, the Wash Solution may develop inert white or brown particulates that float in solution. This is not a cause for concern and does not negatively affect performance.

Prepare the processing plates according to the following table. Cover the plates with a temporary seal (such as MicroAmp<sup>™</sup> Clear Adhesive Film), then store at room temperature for up to 1 hour while you set up the sample plate.

Plate ID	Plate position	Plate type	Reagent	Volume per well
Wash 1 Plate	2		Wash Solution	500 μL
Wash 2 Plate	3	KingFisher <sup>™</sup> 96 Deep-Well Plate	80% Ethanol	500 μL
Elution Plate	4		Elution Buffer	50 μL
Tip Comb Plate	5	Place a KingFisher <sup>™</sup> 96 tip comb f KF microplate or	or DW magnets in a Kir equivalent plate <sup>[1]</sup>	ngFisher <sup>™</sup> 96

<sup>[1]</sup> See "Before you begin" on page 8 for equivalent plates.

**Note:** Open reagent bottles, or reagents poured in reagent reservoirs, have been shown to be stable for ≤2 hours at room temperature.

#### Prepare Binding Bead Mix (200-µL sample input volume)

Prepare the required amount of Binding Bead Mix on each day of use.

- 1. Vortex the Binding Beads to ensure that the bead mixture is homogeneous.
- 2. For the number of required reactions, prepare the Binding Bead Mix according to the following table:

Component	Volume per well <sup>[1]</sup>
Binding Solution	265 μL
Binding Beads	10 μL
Total volume per well	275 μL

<sup>[1]</sup> Include 10% overage when making the Binding Bead Mix for use with multiple reactions.

**Note:** The Binding Bead Mix has been shown to be stable for ≤8 hours at room temperature.

3. Mix well by inversion, then store at room temperature.

#### Prepare sample plate

 Invert the Binding Bead Mix 5 times gently to mix, then add 275 µL to each sample well and the Negative Control well in the Sample Plate (KingFisher<sup>™</sup> 96 Deep-Well Plate).

**Note:** Remix the Binding Bead Mix by inversion frequently during pipetting to ensure even distribution of beads to all samples or wells. The Binding Bead Mix is viscous, so pipet slowly to ensure that the correct amount is added. DO NOT reuse pipette tips to add Binding Bead Mix to the samples, as the high viscosity will cause variations in the volumes added.

- 2. Add 200 µL of sample to each sample well.
- 3. Add 200 µL of Nuclease-free Water (not DEPC-Treated) to the Negative Control well.
- 4. Add the following components to each well, including the Negative Control well.
  - Add 5 µL of Proteinase K.
  - (Optional) If using an extraction control, add the required volume. For more information about using an extraction control, see the assay documentation.

**IMPORTANT!** Add the components only to the top layer of the solution in each well. Do not push the pipette tip into the binding mix layer.

**Note:** The Proteinase K and extraction control can be pre-mixed on each day of use, then kept on ice. Add the combined required volume of Proteinase K and extraction control to each well of the Sample Plate.

For example, if your assay recommends 5  $\mu$ L of the extraction control per reaction, add 10  $\mu$ L of pre-mixed Proteinase K and extraction control to each well.

**Note:** The Proteinase K and extraction control mix has been shown to be stable for ≤8 hours on ice.

#### Process the samples (200-µL sample input volume)

- Select the MVP\_2Wash\_200\_Flex on the KingFisher<sup>™</sup> Flex Magnetic Particle Processor with 96 Deep-Well Head.
- 2. Start the run, then load the prepared plates into position when prompted by the instrument.

Note: Once prepared, the sample plates must be loaded on the instrument within 30 minutes.

3. After the run is complete (~22 minutes after start), immediately remove the Elution Plate from the instrument, then cover the plate with MicroAmp<sup>™</sup> Clear Adhesive Film.

**IMPORTANT!** To prevent evaporation, seal the plate containing the eluate immediately.

The samples are eluted in 50  $\mu$ L of Elution Buffer (see "Prepare the processing plates (200- $\mu$ L sample input volume)" on page 10).



#### Note:

- If bead carry over is observed, place elution plate on a magnetic stand to pellet the beads, then
  pipette the eluate to a new 96-well plate for use in real-time PCR. Review real-time PCR results to
  determine if re-extraction is required.
- To ensure reliable performance of the KingFisher<sup>™</sup> Flex Magnetic Particle Processor, perform preventive maintenance as instructed by the manufacturer.

Place the Elution Plate on ice for immediate use in real-time RT-PCR.

# Extract nucleic acid—Automated method (400-µL sample input volume)

#### Set up the instrument

Ensure that the KingFisher<sup>™</sup> Flex Magnetic Particle Processor with 96 Deep-Well Head is set up with the KingFisher<sup>™</sup> Flex 96 Deep-Well Heating Block.

**IMPORTANT!** Failure to use the proper magnetic head and heat block results in lower yields and potential harm to the instrument.

2. Ensure that the MVP\_2Wash\_400\_Flex program has been downloaded from the MagMAX<sup>™</sup> Viral/Pathogen II Nucleic Acid Isolation Kit product page at www.thermofisher.com and loaded onto the instrument.

#### Prepare the processing plates

**Note:** During the wash steps, the Wash Solution may develop inert white or brown particulates that float in solution. This is not a cause for concern and does not negatively affect performance.

Prepare the processing plates according to the following table. Cover the plates with a temporary seal (such as  $MicroAmp^{T}$  Clear Adhesive Film), then store at room temperature for up to 1 hour while you set up the sample plate.

Plate ID	Plate position	Plate type	Reagent	Volume per well
Wash 1 Plate	2		Wash Solution	1,000 µL
Wash 2 Plate	3	KingFisher <sup>™</sup> 96 Deep-Well Plate	80% Ethanol	1,000 µL
Elution Plate	4		Elution Buffer	50 μL
Tip Comb Plate	5	Place a KingFisher <sup>™</sup> 96 tip comb for DW magnets in a KingFisher <sup>™</sup> 96  KF microplate or equivalent plate <sup>[1]</sup>		

<sup>[1]</sup> See "Before you begin" on page 8 for equivalent plates.

**Note:** Open reagent bottles, or reagents poured in reagent reservoirs, have been shown to be stable for  $\leq 2$  hours at room temperature.

#### Prepare Binding Bead Mix (400-µL sample input volume)

Prepare the required amount of Binding Bead Mix on each day of use.

- 1. Vortex the Binding Beads to ensure that the bead mixture is homogeneous.
- 2. For the number of required reactions, prepare the Binding Bead Mix according to the following table:

Component	Volume per well <sup>[1]</sup>
Binding Solution	530 μL
Binding Beads	20 μL
Total volume per well	550 μL

<sup>[1]</sup> Include 10% overage when making the Binding Bead Mix for use with multiple reactions.

**Note:** The Binding Bead Mix has been shown to be stable for ≤8 hours at room temperature.

3. Mix well by inversion, then store at room temperature.

#### Prepare sample plate

1. Invert the Binding Bead Mix 5 times gently to mix, then add 550 μL to each sample well and the Negative Control well in the Sample Plate (KingFisher<sup>™</sup> 96 Deep-Well Plate).

**Note:** Remix the Binding Bead Mix by inversion frequently during pipetting to ensure even distribution of beads to all samples or wells. The Binding Bead Mix is viscous, so pipet slowly to ensure that the correct amount is added. DO NOT reuse pipette tips to add Binding Bead Mix to the samples, as the high viscosity will cause variations in the volumes added.

- 2. Add 400 μL of sample to each sample well.
- 3. Add 400 µL of Nuclease-free Water (not DEPC-Treated) to the Negative Control well.
- 4. Add the following components to each well, including the Negative Control well.
  - Add 10 µL of Proteinase K.
  - (Optional) If using an extraction control, add the required volume. For more information about using an extraction control, see the assay documentation.

**IMPORTANT!** Add the components only to the top layer of the solution in each well. Do not push the pipette tip into the binding mix layer.

**Note:** The Proteinase K and extraction control can be pre-mixed on each day of use, then kept on ice. Add the combined required volume of Proteinase K and extraction control to each well of the Sample Plate.

For example, if your assay recommends 10  $\mu$ L of the extraction control per reaction, add 20  $\mu$ L of pre-mixed Proteinase K and extraction control to each well.



**Note:** The Proteinase K and extraction control mix has been shown to be stable for ≤8 hours on ice.

#### Process the samples (400-µL sample input volume)

- Select the MVP\_2Wash\_400\_Flex on the KingFisher<sup>™</sup> Flex Magnetic Particle Processor with 96 Deep-Well Head.
- 2. Start the run, then load the prepared plates into position when prompted by the instrument.

Note: Once prepared, the sample plates must be loaded on the instrument within 30 minutes.

3. After the run is complete (~24 minutes after start), immediately remove the Elution Plate from the instrument, then cover the plate with MicroAmp<sup>™</sup> Clear Adhesive Film.

**IMPORTANT!** To prevent evaporation, seal the plate containing the eluate immediately.

The samples are eluted in 50 µL of Elution Buffer (see "Prepare the processing plates" on page 12).

#### Note:

- If bead carry over is observed, place elution plate on a magnetic stand to pellet the beads, then pipette the eluate to a new 96-well plate for use in real-time PCR. Review real-time PCR results to determine if re-extraction is required.
- To ensure reliable performance of the KingFisher<sup>™</sup> Flex Magnetic Particle Processor, perform preventive maintenance as instructed by the manufacturer.

Place the Elution Plate on ice for immediate use in real-time RT-PCR.



# Extract nucleic acid from swabbased samples (manual method)

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Manual nucleic acid extraction can be performed using a sample volume input of 200  $\mu$ L or 400  $\mu$ L. See your assay documentation for specific sample volume input recommendations.

#### Before you begin

- Ensure that you read and understand the information provided in this guide before you begin the extraction procedure.
- Review your assay documentation to determine if an extraction control is recommended to verify
  the efficacy of the nucleic acid preparation. Follow the extraction control guidelines provided in the
  assay documentation.
- Determine the number of required reactions based on the number of patient samples to be processed, plus one Negative Control per plate.
- Prepare fresh 80% Ethanol using Ethanol, Absolute, Molecular Biology Grade and Nuclease-free Water (not DEPC-Treated) for the required number of reactions, plus 10% overage.

Sample input volume	Volume of 80% Ethanol per reaction
200 μL	0.75 mL
400 μL	1.5 mL

- Mark the Negative Control well on the plate.
- Note the following for all kit components and 80% Ethanol: Open reagent bottles, or reagents poured in reagent reservoirs, have been shown to be stable for ≤2 hours at room temperature.

# Extract nucleic acid—Manual method (200-µL sample input volume)

#### Prepare Binding Bead Mix (200-µL sample input volume)

Prepare the required amount of Binding Bead Mix on each day of use.

- 1. Vortex the Binding Beads to ensure that the bead mixture is homogeneous.
- 2. For the number of required reactions, prepare the Binding Bead Mix according to the following table:

Component	Volume per well <sup>[1]</sup>
Binding Solution	265 μL
Binding Beads	10 μL
Total volume per well	275 μL

<sup>[1]</sup> Include 10% overage when making the Binding Bead Mix for use with multiple reactions.

Note: The Binding Bead Mix has been shown to be stable for ≤8 hours at room temperature.

3. Mix well by inversion, then store at room temperature.

#### Digest with Proteinase K (200-µL sample input volume)

This section provides volumes for the sample plate. Your plate layout will depend on the number of samples you run.

1. Invert the Binding Bead Mix 5 times gently to mix, then add 275 μL to each sample well and Negative Control well.

**Note:** Remix the Binding Bead Mix by inversion frequently during pipetting to ensure even distribution of beads to all samples or wells. The Binding Bead Mix is viscous, so pipet slowly to ensure that the correct amount is added. DO NOT reuse pipette tips to add Binding Bead Mix to the samples, as the high viscosity will cause variations in the volumes added.

- 2. Add 200 µL of sample to each sample well of a KingFisher<sup>™</sup> 96 Deep-Well Plate.
- 3. Add 200 µL of Nuclease-free Water (not DEPC-Treated) to the Negative Control well.
- 4. Add the following components to each well, including the Negative Control well.
  - Add 5 µL of Proteinase K.
  - (Optional) If using an extraction control, add the required volume. For more information about using an extraction control, see the assay documentation.

**IMPORTANT!** Add the components only to the top layer of the solution in each well. Do not push the pipette tip into the binding mix layer.

**Note:** The Proteinase K and extraction control can be pre-mixed on each day of use, then kept on ice. Add the combined required volume of Proteinase K and extraction control to each well of the Sample Plate.

For example, if your assay recommends 5  $\mu$ L of the extraction control per reaction, add 10  $\mu$ L of pre-mixed Proteinase K and extraction control to each well.

**Note:** The Proteinase K and extraction control mix has been shown to be stable for ≤8 hours on ice.

- 5. Seal the plate with MicroAmp<sup>™</sup> Clear Adhesive Film, then shake the sealed plate at 1,050 rpm for 2 minutes.
- 6. Incubate the sealed plate at 65°C for 5 minutes (ensure the bottom of the plate is uncovered), then shake the plate at 1,050 rpm for 5 minutes.
- 7. Place the sealed plate on the magnetic stand for 10 minutes or until all of the beads have collected.

#### Wash the beads (200-µL sample input volume)

**Note:** During the wash steps, the Wash Solution may develop inert white or brown particulates that float in solution. This is not a cause for concern and does not negatively affect performance.

1. Keeping the plate on the magnet, carefully remove the cover, then discard the supernatant from each well.

**IMPORTANT!** Avoid disturbing the beads.

- 2. Remove the plate from the magnetic stand, then add 500 µL of Wash Solution to each sample.
- 3. Reseal the plate, then shake at 1,050 rpm for 1 minute.
- 4. Place the plate back on the magnetic stand for 2 minutes, or until all the beads have collected.
- 5. Keeping the plate on the magnet, carefully remove the cover, then discard the supernatant from each well.

**IMPORTANT!** Avoid disturbing the beads.

- 6. Repeat step 2 to step 5 using 500 µL of 80% Ethanol.
- 7. Repeat step 2 to step 5 using 250 µL of 80% Ethanol.
- 8. Dry the beads by shaking the plate (uncovered) at 1,050 rpm for 2 minutes.

#### Elute the nucleic acid (200-µL sample input volume)

- 1. Add 50 μL of Elution Buffer to each sample, then seal the plate with MicroAmp<sup>™</sup> Clear Adhesive Film.
- 2. Shake the sealed plate at 1,050 rpm for 5 minutes.
- 3. Place the plate in an incubator at 65°C for 10 minutes.
- 4. Remove the plate from the incubator, then shake the plate at 1,050 rpm for 5 minutes.
- 5. Place the sealed plate on the magnetic stand for 3 minutes or until clear to collect the beads against the magnets.
- 6. Keeping the plate on the magnet, carefully remove the seal, transfer the eluates to a fresh standard (not deep-well) 96-well plate, then seal the plate with MicroAmp<sup>™</sup> Clear Adhesive Film.

**IMPORTANT!** To prevent evaporation, seal the plate containing the eluate immediately after the transfers are complete.

**Note:** If bead carry over is observed, extend the time on the magnetic stand to further pellet the beads, then pipette the eluate to a new 96-well plate for use in real-time PCR. Review real-time PCR results to determine if re-extraction is required.

Place the plate on ice for immediate use in real-time RT-PCR.

# Extract nucleic acid—Manual method (400-µL sample input volume)

#### Prepare Binding Bead Mix (400-µL sample input volume)

Prepare the required amount of Binding Bead Mix on each day of use.

- 1. Vortex the Binding Beads to ensure that the bead mixture is homogeneous.
- 2. For the number of required reactions, prepare the Binding Bead Mix according to the following table:

Component	Volume per well <sup>[1]</sup>
Binding Solution	530 μL
Binding Beads	20 μL
Total volume per well	550 μL

<sup>[1]</sup> Include 10% overage when making the Binding Bead Mix for use with multiple reactions.

Note: The Binding Bead Mix has been shown to be stable for ≤8 hours at room temperature.

3. Mix well by inversion, then store at room temperature.

#### Digest with Proteinase K (400-µL sample input volume)

This section provides volumes for the sample plate. Your plate layout will depend on the number of samples you run.

1. Invert the Binding Bead Mix 5 times gently to mix, then add 550 μL to each sample well and Negative Control well.

**Note:** Remix the Binding Bead Mix by inversion frequently during pipetting to ensure even distribution of beads to all samples or wells. The Binding Bead Mix is viscous, so pipet slowly to ensure that the correct amount is added. DO NOT reuse pipette tips to add Binding Bead Mix to the samples, as the high viscosity will cause variations in the volumes added.

- 2. Add 400 µL of sample to each sample well of a KingFisher<sup>™</sup> 96 Deep-Well Plate.
- 3. Add 400 µL of Nuclease-free Water (not DEPC-Treated) to the Negative Control well.
- 4. Add the following components to each well, including the Negative Control well.
  - Add 10 μL of Proteinase K.
  - (Optional) If using an extraction control, add the required volume. For more information about using an extraction control, see the assay documentation.

**IMPORTANT!** Add the components only to the top layer of the solution in each well. Do not push the pipette tip into the binding mix layer.

**Note:** The Proteinase K and extraction control can be pre-mixed on each day of use, then kept on ice. Add the combined required volume of Proteinase K and extraction control to each well of the Sample Plate.

For example, if your assay recommends 10  $\mu$ L of the extraction control per reaction, add 20  $\mu$ L of pre-mixed Proteinase K and extraction control to each well.

**Note:** The Proteinase K and extraction control mix has been shown to be stable for ≤8 hours on ice.

- 5. Seal the plate with MicroAmp<sup>™</sup> Clear Adhesive Film, then shake the sealed plate at 1,050 rpm for 2 minutes.
- 6. Incubate the sealed plate at 65°C for 10 minutes (ensure the bottom of the plate is uncovered), then shake the plate at 1,050 rpm for 5 minutes.
- 7. Place the sealed plate on the magnetic stand for 10 minutes or until all of the beads have collected.

#### Wash the beads (400-µL sample input volume)

**Note:** During the wash steps, the Wash Solution may develop inert white or brown particulates that float in solution. This is not a cause for concern and does not negatively affect performance.

1. Keeping the plate on the magnet, carefully remove the cover, then discard the supernatant from each well.

IMPORTANT! Avoid disturbing the beads.

- 2. Remove the plate from the magnetic stand, then add 1 mL of Wash Solution to each sample.
- 3. Reseal the plate, then shake at 1,050 rpm for 1 minute.
- 4. Place the plate back on the magnetic stand for 2 minutes, or until all the beads have collected.
- 5. Keeping the plate on the magnet, carefully remove the cover, then discard the supernatant from each well.

**IMPORTANT!** Avoid disturbing the beads.

- 6. Repeat step 2 to step 5 using 1 mL of 80% Ethanol.
- 7. Repeat step 2 to step 5 using 500 µL of 80% Ethanol.
- 8. Dry the beads by shaking the plate (uncovered) at 1,050 rpm for 2 minutes.

#### Elute the nucleic acid (400-µL sample input volume)

- 1. Add 50 μL of Elution Buffer to each sample, then seal the plate with MicroAmp<sup>™</sup> Clear Adhesive Film.
- 2. Shake the sealed plate at 1,050 rpm for 5 minutes.
- 3. Place the plate in an incubator at 65°C for 10 minutes.
- 4. Remove the plate from the incubator, then shake the plate at 1,050 rpm for 5 minutes.
- 5. Place the sealed plate on the magnetic stand for 3 minutes or until clear to collect the beads against the magnets.
- 6. Keeping the plate on the magnet, carefully remove the seal, transfer the eluates to a fresh standard (not deep-well) 96-well plate, then seal the plate with MicroAmp<sup>™</sup> Clear Adhesive Film.

**IMPORTANT!** To prevent evaporation, seal the plate containing the eluate immediately after the transfers are complete.

**Note:** If bead carry over is observed, extend the time on the magnetic stand to further pellet the beads, then pipette the eluate to a new 96-well plate for use in real-time PCR. Review real-time PCR results to determine if re-extraction is required.

Place the plate on ice for immediate use in real-time RT-PCR.

## Safety





**WARNING! GENERAL SAFETY.** Using this product in a manner not specified in the user documentation may result in personal injury or damage to the instrument or device. Ensure that anyone using this product has received instructions in general safety practices for laboratories and the safety information provided in this document.

- Before using an instrument or device, read and understand the safety information provided in the user documentation provided by the manufacturer of the instrument or device.
- Before handling chemicals, read and understand all applicable Safety Data Sheets (SDSs) and use appropriate personal protective equipment (gloves, gowns, eye protection, and so on). To obtain SDSs, see the "Documentation and Support" section in this document.

#### **Chemical safety**



**WARNING! GENERAL CHEMICAL HANDLING.** To minimize hazards, ensure laboratory personnel read and practice the general safety guidelines for chemical usage, storage, and waste provided below. Consult the relevant SDS for specific precautions and instructions:

- Read and understand the Safety Data Sheets (SDSs) provided by the chemical manufacturer before you store, handle, or work with any chemicals or hazardous materials. To obtain SDSs, see the "Documentation and Support" section in this document.
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing).
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with sufficient ventilation (for example, fume hood).
- Check regularly for chemical leaks or spills. If a leak or spill occurs, follow the manufacturer cleanup procedures as recommended in the SDS.
- · Handle chemical wastes in a fume hood.
- Ensure use of primary and secondary waste containers. (A primary waste container holds the immediate waste. A secondary container contains spills or leaks from the primary container.
   Both containers must be compatible with the waste material and meet federal, state, and local requirements for container storage.)
- After emptying a waste container, seal it with the cap provided.
- Characterize (by analysis if needed) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
- Ensure that the waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.
- **IMPORTANT!** Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.

#### Biological hazard safety



WARNING! BIOHAZARD. Biological samples such as tissues, body fluids, infectious agents, and blood of humans and other animals have the potential to transmit infectious diseases. Conduct all work in properly equipped facilities with the appropriate safety equipment (for example, physical containment devices). Safety equipment can also include items for personal protection, such as gloves, coats, gowns, shoe covers, boots, respirators, face shields, safety glasses, or goggles. Individuals should be trained according to applicable regulatory and company/ institution requirements before working with potentially biohazardous materials. Follow all applicable local, state/provincial, and/or national regulations. The following references provide general guidelines when handling biological samples in laboratory environment.

- Laboratory biosafety manual, fourth edition. Geneva: World Health Organization; 2020 (Laboratory biosafety manual, fourth edition and associated monographs)
   www.who.int/publications/i/item/9789240011311
- U.S. Department of Health and Human Services, Biosafety in Microbiological and Biomedical Laboratories (BMBL), 6th Edition, HHS Publication No. (CDC) 300859, Revised June 2020 https://www.cdc.gov/labs/pdf/CDC-BiosafetymicrobiologicalBiomedicalLaboratories-2020-P.pdf



**WARNING!** Potential Biohazard. If you use the kit with the automated nucleic extraction workflow, the surface of the KingFisher<sup>™</sup> purification system may be considered a biohazard. Use appropriate decontamination methods when working with biohazards.

## Documentation and support

#### Related documentation

Document	Publication Number
Thermo Scientific <sup>™</sup> KingFisher <sup>™</sup> Flex User Manual	N07669

#### **Customer and technical support**

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- Worldwide contact telephone numbers
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  - User guides, manuals, and protocols
  - Certificates of Analysis
  - Safety Data Sheets (SDSs; also known as MSDSs)

**Note:** For SDSs for reagents and chemicals from other manufacturers, contact the manufacturer.

#### Limited product warranty

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