

# MagMAX™ DNA Multi-Sample Ultra 2.0 Kit

With MagMAX™ DNA Cell and Tissue Extraction Buffer for high-throughput isolation of DNA from cultured cells

Catalog Number A45721

Pub. No. MAN0018808 Rev. A.0

**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](http://thermofisher.com/support).

## Product description

The Applied Biosystems™ MagMAX™ DNA Multi-Sample Ultra 2.0 Kit is developed for scalable, rapid purification of high-quality DNA from a variety of sample matrices. DNA purified with this kit can be used in a broad range of molecular biology downstream applications, such as sequencing, genotyping, and qPCR. This protocol describes automated isolation of DNA from cultured cells using the KingFisher™ Flex and the KingFisher™ Duo Prime instruments.

## Contents and storage

Reagents provided in the kit are sufficient for 100 reactions using up to  $1 \times 10^6$  cells per reaction.

Table 1 MagMAX™ DNA Multi-Sample Ultra 2.0 Kit

Component	Amount	Storage
Enhancer Solution	4.5 mL	15–30°C
Proteinase K	4.5 mL	
Binding Solution	45 mL	
DNA Binding Beads	4.5 mL	
Wash I Solution	110 mL	
Elution Solution <sup>[1]</sup>	12 mL	

<sup>[1]</sup> The amount of Elution Solution provided in the kit is sufficient for preparing 50 samples. If needed, additional Elution Solution can be purchased separately (Cat. No. A36582).

Table 2 MagMAX™ DNA Cell and Tissue Extraction Buffer

Component	Amount <sup>[1]</sup>	Storage
MagMAX™ DNA Cell and Tissue Extraction Buffer (Extraction Buffer)	60 mL	15–30°C

<sup>[1]</sup> Also available separately (Cat. No. A45469).

For 1,000 reactions use Cat. No. A36579 (DNA Binding Beads), A36580 (Wash I Solution), A36581 (Lysis/Binding Solution),

2 × A36582 (Elution Solution), and A45470 (MagMAX™ DNA Cell and Tissue Extraction Buffer).

## Required materials not supplied

Unless otherwise indicated, all materials are available through [thermofisher.com](http://thermofisher.com). MLS: Fisher Scientific ([fisherscientific.com](http://fisherscientific.com)) or other major laboratory supplier.

Item	Source
<b>Instrument</b>	
Magnetic particle processor (one of the following, depending on the quantity/volume of sample to be processed):	
• KingFisher™ Duo Prime Purification System	5400110
• For up to $1 \times 10^6$ cells: KingFisher™ Flex Purification System with 96 Deep-Well Head	5400630
• For up to $5 \times 10^6$ cells: KingFisher™ Flex Purification System with 24 Deep-Well Head	5400640
<b>Consumables</b>	
Deep-well plates:	
• For up to $1 \times 10^6$ cells: KingFisher™ Deepwell 96 Plate	95040450
• For up to $5 \times 10^6$ cells: KingFisher™ Flex 24 Deep-well plate	95040470
96-well standard plates (for use with KingFisher™ Flex only; tip comb placement and eluate storage):	97002540
KingFisher™ 96 KF microplates (200 µL)	
Tip comb, compatible with the magnetic particle processor used:	
• KingFisher™ Duo Prime 12-tip comb (for use with KingFisher™ Deepwell 96 Plate)	97003500
• KingFisher™ Duo Prime 6-tip comb (for use with KingFisher™ Flex 24 Deep-well plate)	97003510
• KingFisher™ 96 tip comb for DW magnets (KingFisher™ Flex protocol only)	97002534
• KingFisher™ Flex 24 Deep Well Tip Comb and plate (KingFisher™ Flex protocol only)	97002610
Elution strip (KingFisher™ Duo Prime protocol only; elution step):	97003520
• Elution strip	97003540
• KingFisher™ Duo Cap for elution strip	

Item	Source
<b>Equipment</b>	
Adjustable micropipette	MLS
Multichannel micropipette	MLS
<b>Reagents</b>	
PureLink™ RNase A (20 mg/mL) (RNase A)	12091021
Ethanol, 96–100% (molecular biology grade)	MLS
Nuclease-Free Water	AM9932
<b>Materials</b>	
MicroAmp™ Clear Adhesive Film	4306311

- (Optional): To prevent evaporation and contamination, cover the prepared processing plates with paraffin film or MicroAmp™ Clear Adhesive Film until they are loaded into the instrument.

## Guidelines for DNA Binding Bead Mix

- Vortex the DNA Binding Beads thoroughly, combine them with the Binding Solution in a nuclease-free tube, then invert the tube until homogeneous. This mixture can be stored for up to 1 day before aliquoting into the plates.
- Ensure that the beads stay fully mixed within the solution during pipetting.
- Avoid creating bubbles during mixing and aliquoting.

## General guidelines

- Perform all steps at room temperature (20–25°C) unless otherwise noted.
- Precipitates and high viscosity can occur if Enhancer Solution, Wash I Solution, and Binding Solution are stored when room temperature is too cold. If this occurs, warm them at 37°C, then gently mix to dissolve precipitates. Avoid creating bubbles.
- Yellowing of the Binding Solution and Wash I Solution is normal and will not affect buffer performance
- Per-plate volumes for reagent mixes are sufficient for one plate plus overage. To calculate volumes for other sample numbers, refer to the per-well volume and add 10% overage.

## Before first use of the kit

**Note:** RNase A is recommended for this protocol and must be purchased separately from the kit. See “Required materials not supplied” on page 1.

Prepare Wash II Solution: Make 80% ethanol from 100% absolute ethanol and nuclease-free water.

## Before each use of the kit

Vortex DNA Binding Beads to fully resuspend the beads before each use.

## Perform DNA purification using KingFisher™ Flex (96 deep-well format)

This method is recommended for samples that contain up to  $1 \times 10^6$  cultured cells.

- 1 **Set up the instrument**
  - a. Ensure that the instrument is set up for processing with the proper magnetic head (96 deep-well) for your application.
  - b. Ensure that the proper heat block (96 deep-well, not standard) is installed for your application.
  - c. Ensure that the proper program (MMX\_Ultra2\_Cell\_Tissue\_96\_Flex) has been downloaded from the product page and loaded onto the instrument.

- 2 **Set up the processing plates**

Set up the Wash, Elution, and Tip Comb Plates outside of the instrument according to the following table.

Plate ID	Plate position	Plate type	Reagent	Volume per well
<b>Up to <math>1 \times 10^6</math> cell input</b>				
Wash I Solution Plate	2	Deep Well	Wash I Solution	1,000 $\mu$ L
Wash II Solution Plate 1	3	Deep Well	Wash II Solution	1,000 $\mu$ L
Wash II Solution Plate 2	4	Deep Well	Wash II Solution	500 $\mu$ L
Elution Plate	5	Deep Well	Elution Solution	200 $\mu$ L
Tip Comb	6	Place a 96 Deep-well Tip Comb in a Standard Plate		

**Note:** Load the plates on the instrument immediately after the Sample Plate is prepared.

### 3 Prepare DNA Binding Bead Mix

Prepare DNA Binding Bead Mix according to the following table.

Component	Volume per well
Binding Solution	400 $\mu$ L
DNA Binding Beads	40 $\mu$ L
<b>Total DNA Binding Bead Mix</b>	<b>440 <math>\mu</math>L</b>

### 4 Prepare the Sample Plate

If needed, thaw the cell pellets at room temperature, then carefully remove as much media as possible from each pellet.

- To each cell pellet, add 500  $\mu$ L of Extraction Buffer for every  $1 \times 10^6$  cells.
- Vortex to dislodge the cell pellet from the tube.  
The cell pellet forms a clear, viscous, free-floating mass.
- Using a P1000 pipette, pipet up and down 10 times or until the cell pellet is completely dissolved.

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**IMPORTANT!** The sample can become viscous. It is critical to pipet up and down thoroughly to ensure complete mixing.

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- Transfer each sample to the appropriate well of a deep-well plate (Sample Plate).  
Pipet slowly to ensure that the entire sample is transferred, including the viscous portion.
- (Recommended) Add 10  $\mu$ L of RNase A to each sample, seal the plate with adhesive film, then mix using a plate shaker set at moderate speed for at least 5 minutes at room temperature.  
Alternatively, if a plate shaker is not available, pipet up and down 10 times to mix, then incubate for at least 5 minutes at room temperature.
- Invert the tube of DNA Binding Bead Mix several times to resuspend the beads, then add 440  $\mu$ L of DNA Binding Bead Mix to each sample.  
**Note:**
  - Remix DNA Binding Bead Mix frequently during pipetting to ensure that the beads are evenly distributed among the samples.
  - The mixture is viscous. Pipet slowly to ensure that the correct amount is added to each sample.
- Immediately proceed to process samples on the instrument (next section).

### 5 Process samples on the instrument

- Select the **MMX\_Ultra2\_Cell\_Tissue\_96\_Flex** program on the instrument.
- Start the run, then load the prepared plates in their positions when prompted by the instrument.
- At the end of the run, immediately remove the plate from the instrument, then transfer the eluate to a tube or plate for storage.

Store purified DNA on ice for immediate use, at  $-20^{\circ}\text{C}$  for up to 1 month, or at  $-80^{\circ}\text{C}$  for long-term storage.

## Perform DNA purification using KingFisher™ Flex (24 deep-well format)

This method is recommended for samples that contain up to  $5 \times 10^6$  cultured cells.

- 1 **Set up the instrument**
  - a. Ensure that the instrument is set up for processing with the proper magnetic head (24 deep-well) for your application.
  - b. Ensure that the proper heat block (24-well) is installed for your application.
  - c. Ensure that the proper program (**MMX\_Ultra2\_Cell\_Tissue\_24L\_Flex**) has been downloaded from the product page and loaded onto the instrument.

- 2 **Set up the processing plates** Set up the Wash, Elution, and Tip Comb Plates outside of the instrument according to the following table.

Plate ID	Plate position	Plate type	Reagent	Volume per well
<b>Up to <math>5 \times 10^6</math> cell input</b>				
Wash I Solution Plate	2	Deep Well	Wash I Solution	5 mL
Wash II Solution Plate 1	3	Deep Well	Wash II Solution	4 mL
Wash II Solution Plate 2	4	Deep Well	Wash II Solution	2 mL
Elution Plate	5	Deep Well	Elution Solution	1 mL
Tip Comb	6	Place a 24 Deep-well Tip Comb in a Standard Plate		

**Note:** Load the plates on the instrument immediately after the Sample Plate is prepared.

- 3 **Prepare DNA Binding Bead Mix** Prepare DNA Binding Bead Mix according to the following table.

Component	Volume per well
Binding Solution	2 mL
DNA Binding Beads	200 $\mu$ L
<b>Total DNA Binding Bead Mix</b>	<b>2.2 mL</b>

- 4 **Prepare the Sample Plate** If needed, thaw the cell pellets at room temperature, then carefully remove as much media as possible from each pellet.

- a. To each cell pellet, add 2.5 mL of Extraction Buffer for every  $5 \times 10^6$  cells.
- b. Vortex to dislodge the cell pellet from the tube.  
The cell pellet forms a clear, viscous, free-floating mass.
- c. Using a P1000 pipette, pipet up and down 10 times.
- d. Incubate the sample for 5–10 minutes at room temperature.
- e. Vortex, then use a P1000 pipette to pipet up and down 10 times or until the cell pellet is completely dissolved.

**IMPORTANT!** The sample can become viscous. It is critical to pipet up and down thoroughly to ensure complete mixing.

- f. Transfer each sample to the appropriate well of a deep-well plate (Sample Plate).  
Pipet slowly to ensure that the entire sample is transferred, including the viscous portion.

#### 4 Prepare the Sample Plate *(continued)*

- g. (Recommended) Add 50  $\mu\text{L}$  of RNase A to each sample, seal the plate with adhesive film, then mix using a plate shaker set at moderate speed for at least 10 minutes at room temperature.
- Alternatively, if a plate shaker is not available, pipet up and down 10 times to mix, then incubate for at least 10 minutes at room temperature.
- h. Invert the tube of DNA Binding Bead Mix several times to resuspend the beads, then add 2.2 mL of DNA Binding Bead Mix to each sample.

**Note:**

- Remix DNA Binding Bead Mix frequently during pipetting to ensure that the beads are evenly distributed among the samples.
  - The mixture is viscous. Pipet slowly to ensure that the correct amount is added to each sample.
- i. Immediately proceed to process samples on the instrument (next section).

#### 5 Process samples on the instrument

- a. Select the **MMX\_Ultra2\_Cell\_Tissue\_24L\_Flex** program on the instrument.
- b. Start the run, then load the prepared plates in their positions when prompted by the instrument.
- c. At the end of the run, immediately remove the plate from the instrument, then transfer the eluate to a tube or plate for storage.

Store purified DNA on ice for immediate use, at  $-20^{\circ}\text{C}$  for up to 1 month, or at  $-80^{\circ}\text{C}$  for long-term storage.

### Perform DNA purification using KingFisher™ Duo Prime (96 deep-well format)

This method is recommended for samples that contain up to  $1 \times 10^6$  cultured cells.

#### 1 Set up the instrument

- a. Ensure that the instrument is set up for processing with the proper magnetic head (12 pin) and heat block for your application.
- b. Ensure that the proper program (**MMX\_Ultra2\_Cell\_Tissue\_96\_Duo**) has been downloaded from the product page and loaded onto the instrument.

#### 2 Set up the processing plates

Prepare the 96 deep-well plate according to the following table.

**Note:** The Sample Row will be prepared in a later step.

Row ID	Plate row	Reagent	Volume per well
<b>Plate layout</b>			
Elution Solution	A	Elution Solution	200 $\mu\text{L}$
Tip Comb	B	Tip Comb	N/A
—	C	Empty	
Wash II Solution	D	Wash II Solution	500 $\mu\text{L}$
Wash II Solution	E	Wash II Solution	1,000 $\mu\text{L}$
Wash I Solution	F	Wash I Solution	1,000 $\mu\text{L}$
—	G	Empty	
Sample	H	Sample	Varies

**Note:** Load the plate on the instrument immediately after the Sample Row is prepared.

### 3 Prepare DNA Binding Bead Mix

Prepare DNA Binding Bead Mix according to the following table.

Component	Volume per well
Binding Solution	400 $\mu$ L
DNA Binding Beads	40 $\mu$ L
<b>Total DNA Binding Bead Mix</b>	<b>440 <math>\mu</math>L</b>

### 4 Prepare the Sample Row

If needed, thaw the cell pellets at room temperature, then carefully remove as much media as possible from each pellet.

- To each cell pellet, add 500  $\mu$ L of Extraction Buffer for every  $1 \times 10^6$  cells.
- Vortex to dislodge the cell pellet from the tube.  
The cell pellet forms a clear, viscous, free-floating mass.
- Using a P1000 pipette, pipet up and down 10 times or until the cell pellet is completely dissolved.

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**IMPORTANT!** The sample can become viscous. It is critical to pipet up and down thoroughly to ensure complete mixing.

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- Transfer each sample to Row H (Sample Row) of the prepared deep-well plate.  
Pipet slowly to ensure that the entire sample is transferred, including the viscous portion.
- (Recommended) Add 10  $\mu$ L of RNase A to each sample, pipet up and down 10 times to mix, then incubate for at least 5 minutes at room temperature.
- Invert the tube of DNA Binding Bead Mix several times to resuspend the beads, then add 440  $\mu$ L of DNA Binding Bead Mix to each sample in Row H.  
**Note:**
  - Remix DNA Binding Bead Mix frequently during pipetting to ensure that the beads are evenly distributed among the samples.
  - The mixture is viscous. Pipet slowly to ensure that the correct amount is added to each sample.
- Immediately proceed to process samples on the instrument (next section).

### 5 Process samples on the instrument

- Select the **MMX\_Ultra2\_Cell\_Tissue\_96\_Duo** program on the instrument.
- Start the run, then load the prepared plates in their positions when prompted by the instrument.
- At the end of the run, immediately remove the plate from the instrument, then transfer the eluate to a tube or plate for storage.

Store purified DNA on ice for immediate use, at  $-20^{\circ}\text{C}$  for up to 1 month, or at  $-80^{\circ}\text{C}$  for long-term storage.

## Perform DNA purification using KingFisher™ Duo Prime (24 deep-well format)

This method is recommended for samples that contain up to  $5 \times 10^6$  cultured cells.

- 1 Set up the instrument
  - a. Ensure that the instrument is set up for processing with the proper magnetic head (6 pin) and heat block for your application.
  - b. Ensure that the proper program (**MMX\_Ultra2\_Cell\_Tissue\_24L\_Duo**) has been downloaded from the product page and loaded onto the instrument.

- 2 Set up the processing plates
- Prepare the 24 deep-well plates according to the following table.

**Note:** The Sample Row will be prepared in a later step.

Row ID	Plate row	Reagent	Volume per well
<b>Plate 1 layout</b>			
Sample	A	Sample	Varies
Wash I Solution	B	Wash I Solution	5 mL
Wash II Solution	C	Wash II Solution	4 mL
Wash II Solution	D	Wash II Solution	2 mL
<b>Plate 2 layout</b>			
Elution Solution	A	Elution Solution	1 mL
Tip Comb	B	Tip Comb	N/A
—	C	Empty	
—	D	Empty	

**Note:** Load the plates on the instrument immediately after the Sample Row is prepared.

- 3 Prepare DNA Binding Bead Mix

Prepare DNA Binding Bead Mix according to the following table.

Component	Volume per well
Binding Solution	2 mL
DNA Binding Beads	200 $\mu$ L
<b>Total DNA Binding Bead Mix</b>	<b>2.2 mL</b>

- 4 Prepare the Sample Row
- If needed, thaw the cell pellets at room temperature, then carefully remove as much media as possible from each pellet.

- a. To each cell pellet, add 2.5 mL of Extraction Buffer for every  $5 \times 10^6$  cells.
- b. Vortex to dislodge the cell pellet from the tube.  
The cell pellet forms a clear, viscous, free-floating mass.
- c. Using a P1000 pipette, pipet up and down 10 times.
- d. Incubate the sample for 5–10 minutes at room temperature.
- e. Vortex, then use a P1000 pipette to pipet up and down 10 times or until the cell pellet is completely dissolved.

**IMPORTANT!** The sample can become viscous. It is critical to pipet up and down thoroughly to ensure complete mixing.

#### 4 Prepare the Sample Row *(continued)*

- f. Transfer each sample to Row A (Sample Row) of the prepared Plate 1.  
Pipet slowly to ensure that the entire sample is transferred, including the viscous portion.
- g. (Recommended) Add 50 µL of RNase A to each sample, pipet up and down 10 times to mix, then incubate for at least 10 minutes at room temperature.
- h. Invert the tube of DNA Binding Bead Mix several times to resuspend the beads, then add 2.2 mL of DNA Binding Bead Mix to each sample.  
**Note:**
  - Remix DNA Binding Bead Mix frequently during pipetting to ensure that the beads are evenly distributed among the samples.
  - The mixture is viscous. Pipet slowly to ensure that the correct amount is added to each sample.
- i. Immediately proceed to process samples on the instrument (next section).

#### 5 Process samples on the instrument

- a. Select the **MMX\_Ultra2\_Cell\_Tissue\_24L\_Duo** program on the instrument.
- b. Start the run, then load the prepared plates in their positions when prompted by the instrument.
- c. At the end of the run, immediately remove the plate from the instrument, then transfer the eluate to a tube or plate for storage.

Store purified DNA on ice for immediate use, at –20°C for up to 1 month, or at –80°C for long-term storage.

## Quantitate the gDNA samples

To most accurately quantitate gDNA samples that are isolated from cultured cells, we recommend using a NanoDrop™ spectrophotometer. Quantitation can also be performed using qPCR and the Applied Biosystems™ TaqMan™ RNase P Detection Reagents Kit (Cat. No. 4316831).

## Limited product warranty

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For descriptions of symbols on product labels or product documents, go to [thermofisher.com/symbols-definition](http://thermofisher.com/symbols-definition).

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Revision	Date	Description
A.0	7 October 2019	New document for new sample type.

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