

Manual

MagnifiQ[™] 16 Genomic DNA instant kit

Kit for automated, magnetic isolation of genomic DNA in the 16 samples per plate format. Contains ready-to-use, reagent-filled plates and all necessary consumables.

catalog #	size	compatible devices *
604A-16U-64	64 isolations	Auto-Pure 32A
604A-16V-64	64 isolations	Auto-Pure Mini Auto-Pure S32
604A-16U-256	256 isolations	Auto-Pure 32A
604A-16V-256	256 isolations	Auto-Pure Mini Auto-Pure S32

* Compatible devices

The kit has been tested with specific Allsheng brand isolation devices. This does not preclude it from working with other devices. If your device is not listed, please contact us at info@aabiot.com.

For research use only.

Guarantee

A&A Biotechnology provides a guarantee on this product.

The company does not guarantee the correct performance of this kit in the event of:

- not adhering to the supplied protocol
- use of not recommended equipment or materials
- use of other reagents than recommended or which are not a component of the product
- use of expired or improperly stored product or its components



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Advantages

• MagnifiQ[™] 16 Genomic DNA instant kit does not require initial preparation of buffers. Just add samples to the plate and get extracted material within approximately half an hour.

Specification

• It enables isolation of different samples with universal kit and automated extraction programme.

	sample size		
Bacteria G-, G+ (cultures)	up to 2 x 10 ⁸	protocol time	~ 30 min.
Yeast (cultures)	up to 1 ml	elution volume	100 µl 1
<u>Cell cultures</u>	up to 1 x 10 ⁶	elution solution	Tris buffer
Blood fresh or frozen,	up to 200 µl	binding capacity	30 µg DNA
<u>serum, plasma</u>			gPCR, RT-gPCR,
Animal tissue	up to 20 mg	downstream applications	sequencing
<u>Swab</u>	1 pc		
Feces	20 - 50 mg		
Feces (sample stored in conservation solution)	250 - 500 μl		

Sample type

¹ The elution volume prepared on the plate is 100 µl. To obtain a smaller elution volume, subtract the appropriate amount of elution solution from the wells of columns 6 and 12 on the XP-G plate. Attention! Do not reduce the elution volume below 50 µl. To obtain a larger elution volume, add the appropriate amount of elution solution from the wells of columns 6 and 12 on the XP-G plate. Attention! Do not reduce the elution volume elution volume add the appropriate amount of elution solution from the wells of columns 6 and 12 on the XP-G plate. Attention! Do not increase the elution volume above 300 µl.

Description

MagnifiQ[™] 16 Genomic DNA instant kit is designed for DNA isolation from various types of biological materials. The isolated material is perfect for further analyzes and tests by qPCR and RT-PCR methods and for sequencing.

The MagnifiQ[™] product series is based on the automated isolation of nucleic acids with use of magnetic beads. This method significantly shortens working time and reduces risk of mistake in comparison to manual methods.

Contents

	(604A-16U-64	604A-16U-256		
component	quantity	cat#	quantity	cat#	storage
XP-G - extraction plate	4 pcs	K-P96U22XGD	16 pcs	K-P96U22XGD	15-25 °C
Proteinase K	3 ml	K-PRK-3	12 ml	K-PRK-12	15-25 °C
Tris buffer	30 ml	K-TRIS-30	115 ml	K-TRIS-115	4-8 °C*
LTE 2X buffer	15 ml	K-LTE2X-15	55 ml	K-LTE2X-55	15-25 °C
LSDE buffer	35 ml	K-LSDE-35	140 ml	K-LSDE-140	15-25 °C
tip comb 8	4 x 2 pcs	K-C8U-2	16 x 2 pcs	K-C8U-2	15-25 °C
protective film	4 pcs	K-MQF-4	16 pcs	K-MQF-16	15-25 ℃

* Proteinase K can be stored at 15-25 °C for up to 12 months.

		604A-16V-64	604A-16V-256		
component	quantity	cat#	quantity	cat#	storage
XP-G - extraction plate	4 pcs	K-P96V22XGD	16 pcs	K-P96V22XGD	15-25 ℃
Proteinase K	3 ml	K-PRK-3	12 ml	K-PRK-12	4-8 °C*
Tris buffer	30 ml	K-TRIS-30	115 ml	K-TRIS-115	15-25 ℃
LTE 2X buffer	15 ml	K-LTE2X-15	55 ml	K-LTE2X-55	15-25 °C
LSDE buffer	35 ml	K-LSDE-35	140 ml	K-LSDE-140	15-25 ℃
tip comb 8	4 x 2 pcs	K-C8U-2	16 x 2 pcs	K-C8U-2	15-25 °C
protective film	4 pcs	K-MQF-4	16 pcs	K-MQF-16	15-25 ℃

* Proteinase K can be stored at 15-25 °C for up to 12 months.

Additional equipment and reagents

Necessary

- 1.5 ml Eppendorf tubes (sample lysis)
- automated pipette
- pipette tips

Optional

- RNAse (10 µl per sample), <u>cat # 1006-10</u>
- 2.2 ml plates (sample lysis)
- swing-out rotor centrifuge for 96-well plates
- protective film (lysis in a 96 deep-well plate)
- thermoblock
- vortex

Important notes

The following material preparation protocols apply to the procedure carried out in 1.5 ml Eppendorf tubes. If the material preparation is to be carried out in a 96 deep-well plate see the <u>Additional Information</u>.

Material preparation

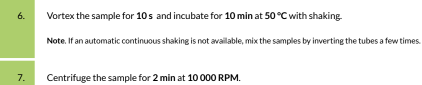
Bacteria G-, G+ (cultures)

Additional reagents you will need:

- BacBreaker bacteria lysis enzyme mix (20 µl per sample), cat # K-BACB-15A
- BS suspension buffer (200 µl per sample), <u>cat # K-BS-30</u>

Option:

- Lysostaphin (5 µl per sample), cat # 1007-3; For Staphylococcus aureus we recommend using lysostaphin.
- Transfer the bacterial culture sample containing 2 x 10⁸ bacteria to the 1.5 ml Eppendorf tube (not included). Centrifuge for 3 min at 10 000 RPM. Discard the supernatant.
 Suspend the bacterial pellet in 200 µl of BS buffer.
 Add 20 µl BacBreaker enzyme mix. Optional RNA removal. Add 10 µl of RNAse (cat # 1006-10). Attention. For lysis of *Staphylococcus* bacteria, add 5 µl of lysostaphin.
 Vortex the sample for 10 s and incubate for 10 min at 50 °C. Attention. For lysis of *Staphylococcus* bacteria with lysostaphin, mix and incubate for 10 min at 37 °C.
 - 5. Add 200 µl LTE 2X and 20 µl Proteinase K.



8. Attention. In the isolation protocol, use the supernatant as the sample.

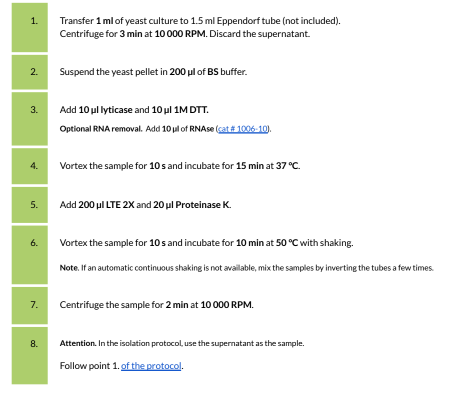
Follow point 1. of the protocol.

Yeast (cultures)

Additional reagents you will need:

- Lyticase (10 µl per sample), cat # 1018-10
- DTT RTU (10 µl 1M solution per sample), cat # 2010-10P
- BS suspension buffer (200 µl per sample), cat # K-BS-30

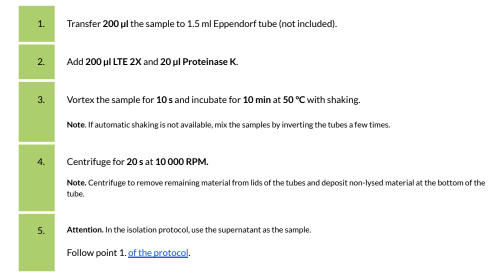
Prepare **1M DTT** solution. Add **1** ml of sterile water (not included) to a vial containing DTT powder to obtain 1M DTT solution. Mix or vortex until complete dissolution of DTT powder. Store a clear solution at -20 °C.



Cell cultures

- Transfer the cell culture sample containing 1 x 10⁶ cells to 1.5 ml Eppendorf tube (not included). Centrifuge for 3 min at 10 000 RPM. Discard the supernatant.
- 2. Suspend the cell pellet in $200 \,\mu$ l of Tris buffer.
- Add 200 μl LTE 2X and 20 μl Proteinase K. Optional RNA removal. Add 10 μl of RNAse (cat # 1006-10).
 Vortex the sample for 10 s and incubate for 10 min at 50 °C with shaking. Note. If an automatic continuous shaking is not available, mix the samples by inverting the tubes a few times.
 Follow point 1. of the protocol.

Blood: fresh or frozen, plasma, serum



Animal tissue

1.	Transfer up to 20 mg of fragmented animal tissue to 1.5 ml Eppendorf tube (not included).
	Note. The tissue should be fragmented by cutting into pieces or homogenization.
2.	Add 400 μl LSDE buffer and 40 μl Proteinase K.
	Optional RNA removal. Add 10 μl of RNAse (<u>cat # 1006-10</u>).
3.	Vortex the sample for $10 s$ and incubate until complete lysis at $50 {}^\circ \! C$ with shaking.
	Note. If an automatic continuous shaking is not available, mix the samples by inverting the tubes a few times.
	Information. The lysis step can last from 1 to 12 hours. For maximum efficiency, lysis should be carried out until the tissue is completely dissolved in the lysis solution.
4.	Centrifuge the sample for 2 min at 10 000 RPM .
5.	Attention. In the isolation protocol, use the supernatant as the sample.
	Follow point 1. of the protocol.

Swabs with transport medium

No additional material preparation is required.

Dry swabs

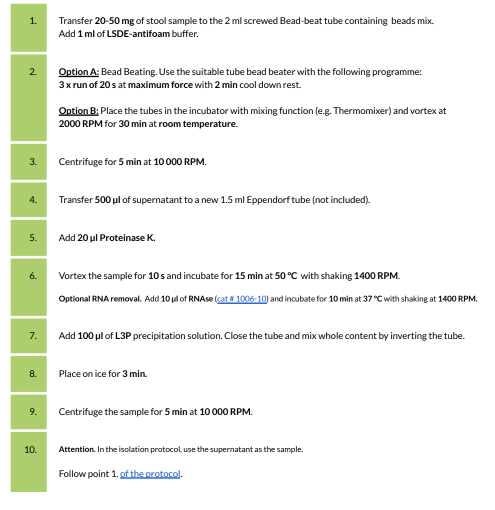
1.	Break or cut off part of the swab with the collected sample and place it in a 1.5 ml Eppendorf tube (not included). Note. The portion of the swab with the collected sample should fit completely into the tube.
2.	Add 500 μl of LSDE buffer and 20 μl Proteinase K. Note. Part of the swab with the sample should be completely immersed in the buffer. Optional RNA removal. Add 10 μl of RNAse (<u>cat # 1006-10</u>).
3.	Vortex the sample for 10 s and incubate for 10 min at 50 °C with shaking. Note. If an automatic continuous shaking is not available, mix the samples by inverting the tubes a few times.
4.	Attention. For the isolation process, take the entire volume of the sample, but not more than 400 μ l. Follow point 1. <u>of the protocol</u> .

Feces (microbiome including G+, G- bacteria)

Additional reagents you will need:

- Bead-beat tubes (2 ml screwed Bead-beat tubes containing beads mix), cat # K-PKCM-50
- L3P precipitation solution (100 µl per sample), cat # K-L3P-60
- LSDE buffer (additional 500 µl per sample), cat # K-LSDE-500
- antifoam (10 µl per sample), cat# K-AYS-1

Before starting the process, mix the LSDE buffer with antifoam. Prepare the LSDE-antifoam mix by combining 1 ml of LSDE buffer with 10 μ l of antifoam per sample. Prepare a volume sufficient for the number of isolated samples with a 10% excess. Mix well before use.



Feces (sample stored in conservation solution StoolSave[™] DNA Protection kit)

Additional reagents you will need:

For stool samples stored in the StoolSave[™] DNA Protection kit (cat # 006-10):

- Bead-beat tubes (2 ml screwed Bead-beat tubes containing beads mix), cat # K-PKCM-50
- L3P precipitation solution (100 µl per sample), cat # K-L3P-60

For stool samples stored in another preservation solution:

- Bead-beat tubes (2 ml screwed Bead-beat tubes containing beads mix), cat # K-PKCM-50
- L3P precipitation solution (100 µl per sample), cat # K-L3P-60
- LSDE buffer (additional 250 µl per sample),cat # K-LSDE-500
- antifoam (10 µl per sample), cat# K-AYS-1

1.	Feces stored in conservation solution StoolSave [™] DNA Protection kit: Transfer 500 μl of stool sample suspended in conservation/ transportation solution to the 2 ml screwed Bead-beat tube containing zirconia beads. Add 500 μl of LSDE buffer.
	Feces stored in another preservation solution:Before starting the process, mix the LSDE buffer with antifoam. Prepare the LSDE-antifoam mix by combining 750 μl of LSDE buffer with 10 μl of antifoam per sample. Prepare a volume sufficient for the number of isolated samples with a 10% excess. Mix well before use.Transfer 250 μl of stool sample suspended in conservation/ transportation solution to the 2 ml screwed Bead-beat tube containing zirconia beads. Add 750 μl of LSDE-antifoam buffer.
2.	<u>Option A</u> : Bead Beating. Use the suitable tube bead beater with the following programme: 3 x run of 20 sec at maximum force with 1 min cool down. <u>Option B</u> : Place the tubes in the incubator with mixing function (e.g. Thermomixer) and vortex at 2000 RPM for 30 min at room temperature.
3.	Centrifuge for 5 min at 10 000 RPM .
4.	Transfer 500 μI of supernatant to a new 1.5 ml Eppendorf tube (not included).
5.	Add 20 μl Proteinase K.
6.	Vortex the sample for 10 s and incubate for 15 min at 50 °C with shaking 1400 RPM. Optional RNA removal. Add 10 µl of RNAse (cat # 1006-10) and incubate for 10 min at 37 °C with shaking at 1400 RPM.

- 7. Add **100** µl of L3P precipitation solution. Close the tube and mix whole content by inverting the tube.
- 8. Place on ice for **3 min**.
- 9. Centrifuge the sample for **5 min** at **10 000 RPM**.

10. Attention. In the isolation protocol, use the supernatant as the sample.

Follow point 1. of the protocol.

Protocol

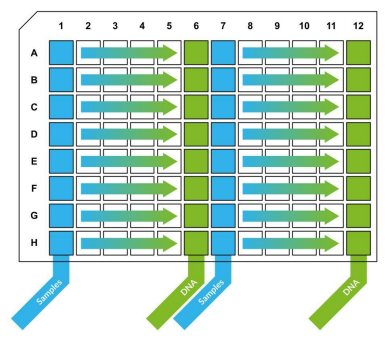
Protocol files

device	protocol name	protocol file	installation		
			1.	Create folder "items" on a USB drive and copy the protocol file to it.	
		aabiot.com/protocols/magnifig	2.	Insert the USB drive into a USB slot in the device.	
Auto-Pure Mini	MQ-UND-MI	/MI/MQ-UND-MI.txt	3.	On a device screen, go to Settings > System > Transfer >Import.	
			4.	Select the protocol and tap "Import".	
Auto-Pure Mini (QR code)	MQ-UND-MI		1. 2.		
			1.	Create folder "items" on a USB drive and copy the protocol file to it.	
	MQ-UND-32A	aabiot.com/protocols/magnifig /32A/MQ-UND-32A.txt	2.	Insert the USB drive into a USB slot in the device.	
Auto-Pure 32A			3.	On a device screen, go to Settings > Im.&Export > Import.	
			4.	Select the protocol and tap "Import."	
	MQ_UND_S32	aabiot.com/protocols/magnifig /S32/MQ_UND_S32.txt	1.	Create folder "im_export_protocols" on a USB drive and copy the protocol file to it.	
Auto-Pure S32			2.	Insert the USB drive into a USB slot in the device.	
			3.	On a device screen, go to Protocols >Import.	
			4.	Select the protocol and tap "Import".	

Extraction protocol

- 1. Carefully centrifuge the **XP-G** plate for **1 min** at **2000 RPM**.
- 2. Gently remove the foil from the **XP-G** plate.
- 3. Add 400 µl of samples to the wells in columns 1 and 7 of the XP-G plate.
- 4. Place one or two **XP-G** plates in the extraction device.
- 5. Place the appropriate number of tip combs 8 in the extraction device.
- 6. Run the protocol on your device.
- 7. After the program is over, remove the combs and then remove **XP-G** plate from the extraction device and seal it with **protective film**. The extracted DNA is located in columns **6** and **12**.

Note. For longer storage of extracted material, transfer it from the plate to appropriate tubes and store at 4 °C.



Additional information

Preparation of material in a 96 deep-well plate

Lysis of the material in a 96 deep-well plate should be carried out according to the respective procedure for 1.5 ml Eppendorf tubes in the Material Preparation section. The following changes should be made:

- Incubation parameters
 Increase the incubation temperature by 5 °C and extend the time by 10 min with a minimum of 1000 RPM continuous shaking speed. Secure tight sealing to prevent any well-to-well cross contamination!
- Centrifugation parameters Centrifuge the plate for **5 min** at **1 000 x g**. For feces sample preparation use specific centrifugation conditions to ensure formation of compact pellets in the deep-well plate.

Safety information

	Proteinase K
DANGER	H315 Causes skin irritation. H319 Causes serious eye irritation. H334 May cause allergy or asthma symptoms or breathing difficulties if inhaled. H335 May cause respiratory irritation. P261 Avoid breathing dust. P305+P351+P338 If in eyes: rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing. P342+P311 If experiencing respiratory symptoms call a Poison Center or doctor/physician.
^	LTE 2X
WARNING	H302 Harmful if swallowed. H315 Causes skin irritation. H319 Causes serious eye irritation. P305+P351+P338 If in eyes: rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
	XP-G - extraction plate
I DANGER	H225 Highly flammable liquid and vapor. H302+H312+H332 Harmful if swallowed, in contact with skin or if inhaled. H314 Causes severe skin burns and eye damage. H412 Harmful to aquatic life with long lasting effects. P210 Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking. P273 Avoid release to the environment. P280 Wear protective gloves/protective clothing/eye protection/face protection/hearing protection. P301+P312+P330 If swallowed: Call a poison center/doctor/ if you feel unwell. P303+P361+P353 If on skin (or hair): Take off immediately all contaminated clothing. Rinse skin with water or shower. P304+P340 If inhaled: Remove person to fresh air and keep comfortable for breathing. P305+P351+P338 If in eyes: rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.



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