





- Wear appropriate skin and eye protection throughout the extraction procedure
- Lysis buffer BLM, mag particle suspension BLM and Wash buffer BLM 1 contain high concentrations of salts and detergents. **Note:** In case of accidental contact, thoroughly rinse or flush the affected areas with water
- Prepared Wash buffer BLM 2 contains up to 70 % organic solvents. Keep away from naked flames.

Kit component	GHS symbol	Hazard phrases	Precaution phrases
Lysis buffer BLM	 Warning	H302/H315/H319	P280/P305+P351+P338/P362/ P301+P312/P332+P313
Protease	 Danger	H315/H319/H334/ H335	P261/P305+P351+P338/ P342+P311
mag particle-suspension BLM	 Danger	H314	P260/P303+P361+P353/P305+P351+ P338/P310/P405
Wash buffer BLM 1	 Danger	H302+H312+H332/ H314/H412	P260/P303+P361+P353/ P305+P351+P338/P310/P405
Wash buffer BLM 2 (concentrate)	-	-	-
Elution buffer BLM	-	-	-

SDS (Safety data sheet) are available at our "Genomics Resource Center" on our webpage www.lgcgroup.com/genomics.



www.lgcgroup.com/genomics

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40420/4.00-2016-07



mag midi kit

Catalogue number **40402** and **40420**

(For research use only. Not for use in diagnostic procedures.)

Description

mag™ kits use magnetic separation for the preparation of nucleic acids.

Superparamagnetic particles coated with mag surface chemistry are used to capture nucleic acids from a sample. The nucleic acid/particle complex is subsequently washed to remove impurities. The nucleic acid is then eluted from the particles and ready for use in downstream processes.



Kit uses

mag midi kits are used to extract DNA from whole blood, blood products and tissue lysates. The method was developed and optimised using 50 µL of whole blood. The following anticoagulants have been tested and found to be compatible with mag nucleic acid extraction kits:

- EDTA
- Heparin
- Citrate

For additional information or advice on protocols please contact our application specialists via email: info.de@lgcgroup.com or Tel: +49 (0)30 5304 2200.

	Colour	Cat. 40402	Cat. 40420
Lysis buffer BLm	Blue	15 mL	100 mL
Protease	Grey	12 mg	120 mg
mag particle suspension BLm	White	1,7 mL	18 mL
Wash buffer BLm 1	Red	30 mL	250 mL
Wash buffer BLm 2 (concentrate)	Yellow	18 mL	150 mL
Elution buffer BLm	Black	15 mL	200 mL

Additional required reagents:

- Ultra pure sterile water
- Ethanol
- Acetone

Additional buffers can be purchased separately, catalogue numbers available on request

Storage

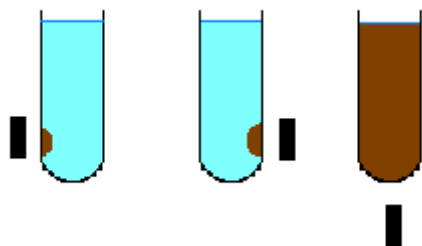
Kit components should be used within 12 months of delivery and stored under the recommended conditions. Please refer to the kit box label for the expiry date.

Room temperature	-20 °C
Lysis buffer BLm	Protease
mag particle suspension BLm	
Wash buffer BLm 1	
Wash buffer BLm 2	
Elution buffer BLm	

Problem	Possible cause	Corrective action
PCR inhibition	Incomplete buffer removal	Ensure all the buffer is removed before adding the next buffer. Check and if necessary adjust the liquid handling parameters for automated systems
Low yield	Poor protease activity	Prepare the protease as detailed in the 'Reagent preparation' section, aliquot into several tubes and store -20 °C. Remove and thaw aliquots as required. Do not use protease which has been kept at room temperature for an extended period of time
	Inefficient binding	Ensure that the lysate, ethanol and mag particles are mixed thoroughly
	Wash buffer BLm 2 acetone composition <70 %	Ensure that the Wash buffer BLm 2 bottle is closed tightly when not in use to prevent acetone evaporation
Coloured eluates	Incomplete buffer removal	Ensure all the buffer is removed before adding the next buffer. Check and if necessary adjust the liquid handling parameters for automated systems
	Heavily stained sample material	Contact our technical specialists for advice
Particles present in eluates	Aspirating too fast	Reduce the speed at which supernatants are removed
	Loose pellet	Increase separation time to allow time for a tighter pellet to form
	Disrupting pellet during aspiration	Position tip further away from pellet whilst removing supernatants
Low ratio between A₂₆₀ and A₂₈₀	Acetone carryover in eluate	Acetone has a maximum UV absorbance at 268 nm and a A ₂₆₀ /A ₂₈₀ of 1.53. If this phenomenon occurs prolong the drying time to ensure all the acetone evaporates

Using sep™ boxes (cont'd.)

- The magnets can be placed in three positions in relation to the sample – left, right and underneath (away from the sample)



- For effective re-suspension of particle pellets it is recommended to move the magnets from the left to right positions using the 'cycle mode'. See sep box operating manual for more details
- For efficient elution of the nucleic acids from the particles it is recommended to use the 'cycle mode' during the elution incubation period.

Presence of precipitates

Salt precipitates can form in Lysis buffer BLm, mag particle suspension BLm and Wash buffer BLm 1 at low temperatures. Check for the presence of precipitates prior to use and if required re-dissolve them by incubating the reagents at 37 °C.

Preparation of tissue lysate

Perform overnight protease incubation at 55 °C to create the tissue lysate. Add 50 µL of Lysis buffer BLm to a fresh tube. Transfer 50 µL of the tissue lysate to the tube containing the Lysis buffer BLm. Mix thoroughly. Continue from step 4 of the protocol.

Protease

Prepare the Protease by adding the appropriate amount of pure water to the vial of Protease. For kit catalogue number 40402 add 600 µL. For kit catalogue number 40420 add 6 mL. When not in use store the Protease at -20 °C.

Lysis mix

To reduce the number of pipetting steps a lysis mix can be prepared at the start of the process. Thaw the Protease thoroughly. Add 5 µL of Protease to 50 µL of Lysis buffer BLm for the number of samples to be processed. The table below gives some example calculations including a 10 % wastage factor. Mix thoroughly. Use the lysis mix within 30 minutes.

Number of samples	Vol. of Lysis buffer BLm	Vol. of Protease
1	55 µL	5.5 µL
20	1.1 mL	110 µL
96	5.3 mL	528 µL

mag particle suspension BLm

The mag particles are suspended in a specially formulated buffer which avoids rapid sedimentation or clogging of particles during handling. Mix the suspension thoroughly before use to fully re-suspend the particles.

Wash buffer BLm 2

Prepare the Wash buffer BLm 2 according to the instructions on the bottle label. For kit catalogue number 40402 add 14 mL of acetone to each bottle. For kit catalogue number 40420 add 350 mL of acetone. Ensure the lid is closed tightly when the bottle is not in use to avoid evaporation.

Manual protocol

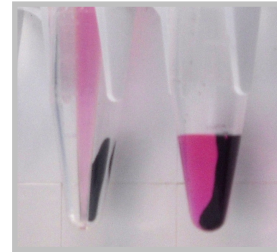
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1. Ensure the blood samples are well mixed prior to starting the protocol
2. Add 50 μL of **Lysis buffer BLm** and 5 μL of **Protease** to 50 μL of blood. Mix thoroughly, set pipette volume to 80 μL and pipette up and down 5 times
3. Incubate at 55 $^{\circ}\text{C}$ for 10 minutes then allow to cool down to room temperature
4. Add 50 μL of **ethanol** to each sample
5. Ensure the **mag particle suspension BLm** is fully re-suspended. Add 16 μL to each sample. Mix thoroughly, set pipette volume to 120 μL and pipette up and down 5 times
6. Incubate for 2 minutes at room temperature to allow sufficient time for binding to occur
7. Bring magnet into contact with the sample tubes. Wait for 1 minute at room temperature to allow the mag particles to form a pellet
8. Remove the supernatant and discard. Ensure as much of the supernatant is removed as possible without dislodging the particle pellet
9. Move the magnet away from the sample tubes
10. Add 170 μL of **Wash buffer BLm 1** and re-suspend the pellet. Mix thoroughly, set pipette volume to 150 μL and pipette up and down 5 times or until pellet is fully re-suspended
11. Incubate at room temperature for 10 minutes, agitating the sample during the time period. Use a shaker or vortex periodically
12. Bring magnet into contact with the sample tubes. Wait for 1 minute at room temperature to allow the mag particles to form a pellet
13. Remove the supernatant and discard. Ensure as much of the supernatant is removed as possible without dislodging the particle pellet
14. Repeat steps 9 to 13 with 175 μL of **Wash buffer BLm 2**
15. Repeat steps 9 to 13 a second time with 175 μL of **Wash buffer BLm 2**
16. Dry the pellet at 55 $^{\circ}\text{C}$ for 6 minutes. Sample tubes must be left open to allow evaporation to occur
17. Add 63 μL of **Elution buffer BLm** and re-suspend the pellet. Mix thoroughly, set pipette volume to 50 μL and pipette up and down 5 times or until pellet is fully re-suspended
18. Incubate at 55 $^{\circ}\text{C}$ for 10 minutes, agitating the sample during the time period. Use a heated shaker or vortex periodically
19. Bring magnet into contact with the sample tubes. Wait for 3 minutes at room temperature to allow the mag particles to form a pellet
20. Remove the eluate and place into a new sample tube. To avoid particle transfer it is recommended to transfer only 50 μL of the eluate.

Tips for manual protocol

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For manual testing of the protocol or if no magnet is available it is recommended to spin tubes for 10 seconds to enable the magnetic particles to form a pellet.



When removing supernatants it is important to remove as much of the liquid as possible without dislodging the particle pellet. With magnets used for manual protocols the particle pellet forms on the back wall of the sample tube. When placing the pipette tip inside the tube be sure to aim the end of the tip to the front wall of the sample tube to avoid disrupting the particle pellet.

To remove as much liquid as possible it is recommended to aspirate once, let any liquid run down the walls of the tube and then aspirate a second time to remove these remnants of liquid.

Tips for automated protocol

Follow the manual protocol as specified overleaf in respect to volumes. Tips on automated mixing are given below:

Mixing with automated liquid handling system

- Set mixing volume to be between 50 % to 80 % of the volume to be mixed (instrument dependent)
- For each mixing step aspirate and dispense between 5 and 10 times depending on the efficiency of the liquid handler
- Keep mix aspirate and dispense speeds low with Lysis buffer BLm to avoid frothing
- Increase aspirate and dispense speeds when re-suspending pellets in wash buffers to ensure complete re-suspension.

Using sep™ boxes

- sep boxes are computer driven magnetic particle collectors with active cooling and heating functionality
- Depending on the sep box used the volumes specified in the manual protocol may need to be changed to be within their maximum working volume. **Note:** sep 96 x 0.2 has a maximum working volume of 180 μL .