

XpressDNA Blood Mini Kit

Protocol for isolation of high quality total genomic DNA from whole blood EDTA\Heparin\ Citrate\Fluoride treated vacutainer (fresh/frozen/stored).

Process Flow



Kit Contents

Components	Storage Conditions	Shipping Conditions
Blood Lysis Buffer 1	RT	RT
Solution A	RT	RT
Blood Lysis Buffer 2	RT	RT
Proteinase K	2 - 8 °C	RT
Proteinase K Buffer	2 - 8 °C	RT
Blood MagNa Mix	RT	RT
Blood Wash Buffer 1	RT	RT
Blood Wash Buffer 2	RT	RT
Blood Elution Buffer	RT	RT
MagNa Stand (optional)	RT	RT

^{*} RT denotes 15 - 25°C.

Materials not provided with the kit

- 1. 100% Ethanol to Wash Buffers as indicated on the bottle.
- 2. Water bath/heat block at 56°C.
- 3. Reconstitute Proteinase K with Proteinase K Buffer and store at $2 8^{\circ}$ C.

Important

Pay attention to standard lab practices and safety information before beginning the procedure. For more information, refer the appropriate Material Safety Data Sheet (MSDS) available from the product supplier or download from our website http://www.maggenome.com/

Technical Support

For any product related queries please write to us on info@maggenome.com, sales@maggenome.com, support@maggenome.com, support@maggenome.com,

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Protocol

	1. To a sterile 1.5 ml tube, add 400 μl of blood sample.		
	2. Add 1 ml of Blood Lysis Buffer 1 and invert mix the tube gently (4 – 5 times).		
	3. Add 50 µl of Solution A and invert mix the tube 4-5 times (Do not pipette mix).		
	4. Incubate the samples for 5 minutes at RT.		
	5. Centrifuge at 8000 rpm for 2 minutes at RT.		
n	6. Carefully discard the supernatant and add 1 ml of Blood Lysis Buffer 1 again.		
Blood Lysate Preparation	7. Resuspend the pellet by pipette mixing.		
Tieparation	8. Centrifuge the samples at 8000 rpm for 2 minutes at RT and discard the supernatant.		
	9. Add 500 μl of Blood Lysis buffer 2 to the pellet and resuspend by pipette mixing.		
	10. Add 20 μl of Proteinase K and resuspend by pipetting.		
	11. Incubate at 56°C for 10 minutes.		
	12. Transfer the lysate to a fresh tube.		
	(Note: Vortex the Blood MagNa Mix thoroughly before the next step)		
	13. Add 350 μl of Blood MagNa Mix and gently invert the tube 10 - 12 times to mix		
	properly. Do not vortex.		
DNA Binding	14. Incubate the samples at RT for 5 minutes.		
	15. Place the tube on a MagNa Stand until the solution becomes clear.		
	16. Carefully discard the supernatant without removing the tube from the MagNa Stand.		
	Ensure the magnetic nanoparticles are not disturbed.		
	17. To the magnetic nanoparticles, add 500 μl of Blood Wash Buffer 1, remove the		
	tube from the MagNa Stand and resuspend by thorough pipette mixing to ensure		
	complete dispersion of the particles. (Note: Use 200 µl pipette for better resuspension of the pellet.)		
	18. Place the tube back on the MagNa Stand for 30 - 60 seconds till the solution becomes		
	clear.		
	19. Discard the supernatant without removing the tube from the MagNa stand. Ensure the		
DNA Washing	magnetic nanoparticles are not disturbed.		
	20. Add 500 μl of Blood Wash Buffer 2 & gently invert mix the tube 5 – 6 times without		
	removing from the MagNa Stand (surface wash only).		
	21. Discard the supernatant without removing the tube from the MagNa Stand.		
	22. Repeat steps 20 - 21.		
	23. Air dry the magnetic nanoparticles without removing the tube from MagNa Stand for		
	10 - 15 minutes without over drying them.		



DNA Elution	24. After drying, remove the tube from the MagNa Stand.		
	25. Add 100 μl of Blood Elution buffer and resuspend the magnetic nanoparticles by		
	pipette mixing thoroughly.		
	26. Incubate at 56°C for 5 minutes with intermittent tapping.		
	27. Place the tube back on the MagNa Stand for 5 minutes or until the solution becomes		
	clear.		
	28. Carefully transfer the supernatant containing DNA to a sterile 1.5 ml tube, without		
	removing the tube from the MagNa Stand. Ensure the magnetic nanoparticles are not		
	disturbed.		
	29. Discard the magnetic nanoparticles in the appropriate hazard container.		

Note: In the elution step, if the Magnetic particles take more than 10 minutes for clearing, spin the tubes at 14,000 rpm for 5 minutes, place on MagNa Stand until solution clears and then collect the supernatant with pure DNA.



Troubleshooting Guide

Observation	Possible causes	Suggested Solution
	Incomplete Lysis	Proteinase K was not used at the suggested amount and for the specified time. Ensure proper re-suspension of the pellet while adding the Lysis buffer.
		Make sure that the incubation temperature and time for lysis is followed as per the protocol. In some cases when DNA content is very high, the
		lysate may appear slimy. Ensure thorough pipette mixing during lysis and before adding MagNa Mix in such circumstances.
Low DNA yield or Poor Quality	Incorrect reagent volumes were used	Use the exact volumes of reagents mentioned in the protocol.
	MagNa Mix was improperly handled	Resuspend the MagNa Mix by vortexing prior to use.
	Magnetic particle loss during binding or washing steps	Carefully remove the supernatant from the tube without removing the tube from the magnetic stand and without disturbing the MagNa particles.
	Improper elution	Completely resuspend the MagNa particles in elution buffer before incubation at 56°C for elution. Tap the tube few times during the 10 min incubation.
	Ethanol is not added to wash buffers	Add 100% ethanol to wash buffers before use as indicated on the bottles.
Poor performance of extracted DNA in downstream applications	Ethanol carryover	Air dry the MagNa particles after the washing steps to remove ethanol completely, but do not over dry the pellet.
	Salt carryover	Ensure that the correct amount of ethanol is added to the Wash Buffers and the two wash steps are performed with Wash Buffer 2.