



Genemed Synthesis, Inc.
6203 Wood Lake Center Dr., Bldg. 2, San Antonio, TX 78244, USA
Toll free (800) 344-5337; Phone: (210) 745-5988; Fax (210) 745-5992
Email: info@genemedsyn.com, Website: www.genemedsyn.com

Product Data Sheet

Cat # V-200-50	DNA PLASMID EXTRACTION SYSTEM	Size: 50 Preps
Cat # V-200-250	DNA PLASMID EXTRACTION SYSTEM	Size: 250 Preps

Plasmid DNA Extraction System

GSI Plasmid DNA Extraction System provides a simple, fast and cost-effective method to purify plasmid DNA without phenol/chloroform extraction. It is based on binding of DNA to silica-based membranes in chaotropic salts. An average yield of 1 to 40 µg of plasmid DNA can be expected from 1 to 5 ml overnight bacterial culture.

Downstream Application

- * Restriction digestion
- * Radioactive and fluorescent sequencing
- * Transformation
- * Ligation
- * PCR, RAPD

Product Contents

Preps	4
MX1 Buffer (RNaseA added)	1.5ml
MX2 Buffer	2ml
MX3 Buffer	3ml
WN Buffer	1ml
WS Buffer	1ml
Elution Buffer	0.5ml
Mini Plus™ Column	4
Collection Tube	4
Protocol	1

Shipping & Storage

The sample of Mini Plus™ Plasmid DNA Extraction System is shipping and storage at ambient temperature up to 6 months.

If precipitate form by freezing temperature on any buffer, warm up at 37°C to redissolve.

Protocol

❖ Please read the following notes before starting the procedures.

Important Notes

- Add 4 ml of 98 - 100 % ethanol into WN Buffer bottle when first open.
- Add 4 ml of 98 - 100 % ethanol into WS Buffer bottle when first open.
- Buffers provided in this system contain irritants. Appropriate safety apparels such as gloves and lab coat should be worn.
- If precipitation forms in MX2 Buffer, incubate the buffer at 55 °C for 10 minutes to redissolve the salt precipitates.
- Do not shake MX2 Buffer, SDS in MX2 will lead to serious foaming.
- All procedures should be done at room temperature (20 - 25 °C).
- All centrifugation steps are done at 7,000 x g - 10,000 x g (9,000 rpm – 13,000 rpm) in a microcentrifuge, if not notice.
- For long-term storage of the eluted plasmid, TE buffer should be used for elution. Since EDTA in TE may affect downstream applications, Elution Buffer (provided) or ddH₂O (pH 7.0 - 8.5) is preferred for DNA elution immediately used for further enzymatic reactions.

I. Protocol for Spin Method:

1. **Grow 1 to 5 ml plasmid-containing bacterial cells in LB medium with appropriate antibiotic(s) overnight (12-16 hours) with vigorous agitation.**
2. **Pellet the cells by centrifuging for 1 - 2 minutes. Decant the supernatant and remove all medium residue by pipet.**
3. **Add 200 µl of MX1 Buffer to the pellet, resuspend the cells completely by vortexing or pipetting.**



Genemed Synthesis, Inc.

6203 Wood Lake Center Dr., Bldg. 2, San Antonio, TX 78244, USA
Toll free (800) 344-5337; Phone: (210) 745-5988; Fax (210) 745-5992
Email: info@genemedsyn.com, Website: www.genemedsyn.com

No cell clumps should be visible after resuspension of the pellet.

4. **Add 250 μ l of MX2 Buffer and gently mix (invert the tube 4-6 times) to lyse the cells until the lysate becomes clear. Incubate at room temperature for 1 - 5 minutes.**

Do not vortex, vortexing will shear genomic DNA.

5. **Add 350 μ l of MX3 Buffer to neutralize the lysate, then immediately and gently mix the solution.** The white precipitate should be formed.

6. **Centrifuge at 10,000 x g (13,000 rpm) for 5 - 10 minutes, meanwhile place a Mini Plus™ Column onto a Collection Tube.**

7. **Transfer the supernatant carefully into the column.**

8. **Centrifuge at 7,000 x g (9,000 rpm) for 30 - 60 seconds. Discard the flow-through.**

9. **Wash the column once with 0.5 ml WN Buffer by centrifuging at 7,000 x g (9,000 rpm) for 30 - 60 seconds. Discard the flow-through.**

10. **Wash the column once with 0.7 ml WS Buffer by centrifuging at 7,000 x g (9,000 rpm) for 30 - 60 seconds. Discard the flow-through.**

11. **Centrifuge the column at 10,000 x g (13,000 rpm) for another 3 minutes to remove residual ethanol.**

It is important to remove ethanol residue, residual ethanol may inhibit subsequent enzymatic reactions.

12. **Place the column onto a new 1.5-ml centrifuge tube. Add 50 μ l of Elution Buffer (provided) onto the center of the membrane.**

For effective elution, make sure that the elution solution is dispensed onto the center of the membrane and is completely absorbed.

13. **Stand the column for 2 - 3 minutes and centrifuge at 10,000 x g (13,000 rpm) for 2 - 3 minutes to elute DNA.**

14. **Store plasmid DNA at 4 °C or -20 °C.**

II. Protocol for Vacuum Method:

1. Grow 1 to 5 ml plasmid-containing bacterial cells in LB medium with appropriate antibiotic(s) overnight (12-16 hours) with vigorous agitation.

2. Pellet the cells by centrifuging for 1 - 2 minutes. Decant the supernatant and remove all medium residue by pipet.

3. Add 200 μ l of MX1 Buffer to the pellet, resuspend the cells completely by vortexing or pipetting.

No cell clumps should be visible after resuspension of the pellet.

4. Add 250 μ l of MX2 Buffer and gently mix (invert the tube 4-6 times) to lyse the cells until the lysate becomes clear. Incubate at room temperature for 1 - 5 minutes.

Do not vortex, vortexing will shear genomic DNA.

5. Add 350 μ l of MX3 Buffer to neutralize the lysate, then immediately and gently mix the solution.

The white precipitate should be formed.

6. Centrifuge at 10,000 x g (13,000 rpm) for 5 - 10 minutes, meanwhile insert the tip of a Mini Plus™ Column into the luer-lock of a vacuum manifold (e.g. Promega's Vac-man*).

7. Transfer the supernatant carefully into the column.

8. Apply vacuum to draw all the liquid into the manifold.

9. Wash the column once with 0.5 ml WN Buffer by re-applying vacuum to draw all the liquid.

10. Wash the column once with 0.7 ml WS Buffer by re-applying vacuum to draw all the liquid.

It is important to remove ethanol residue, residual ethanol may inhibit subsequent enzymatic reactions.

11. Place the column onto a Collection Tube. Centrifuge the

12. column at 10,000 x g (13,000 rpm) for another 3 minutes to remove residual ethanol.

It is important to remove ethanol residue, residual

All buffers need to be mixed well before use.



Genemed Synthesis, Inc.

6203 Wood Lake Center Dr., Bldg. 2, San Antonio, TX 78244, USA
Toll free (800) 344-5337; **Phone:** (210) 745-5988; **Fax** (210) 745-5992
Email: info@genemedsyn.com, **Website:** www.genemedsyn.com

ethanol may inhibit subsequent enzymatic reactions.

13. Place the column onto a new 1.5 ml centrifuge tube. Add 50 μ l of Elution Buffer (provided) onto the center of the membrane.
For effective elution, make sure that the elution solution is dispensed onto the center of the membrane and is completely absorbed.
14. Stand the column for 2 - 3 minutes and centrifuge at 10,000 x g (13,000 rpm) for 2 - 3 minutes to elute DNA.
15. Store plasmid DNA at 4 °C or -20 °C.

* Vac-man is a trademark of Promega Corporation
