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Introduction

The EZN.A[™] Mag-Bind[®] family of products is an innovative system that radically simplifies extraction and purification of nucleic acids from a variety of sources. Key to the system is Omega Bio-Tek's proprietary Mag-Bind[®] Particle that avidly, but reversibly, binds DNA or RNA under certain optimal conditions allowing proteins and other contaminants to be removed. Nucleic acids are easily eluted with deionized water or low salt buffer.

The E.Z.N.A.[™] Mag-Bind[®] Plasmid Mega Kitcombines the power of Mag-Bind[®] technology with the time-tested consistency of alkaline-SDS lysis of bacterial cells to deliver high qualityplasmid DNA. Cultured bacterial cells are pelleted by centrifugation, cells are then suspended and lysed in a alkaline-SDS buffer. By addition of neutralization buffer, genomic DNA, proteins are removed. The cleared cell lysate is mixed with magnetic particles on which the DNA binds. With two wash steps, the purified DNA was eluted with lower salt bufferorwater. Yields vary according to plasmid copynumber, E.colistrain, and conditions of growth, but 1L of overnight culture in LB medium typically produces 5-10 mg high-copy plasmid DNA. The purified plasmid can be used directly for automated fluores cent DNA sequencing, such as with BigDye sequencing chemistry, as well as for other standard molecular biology techniques including restriction enzyme digestion.

Storage and stability

All Mag-Bind[®] Plasmid isolation components are guaranteed for at least 12 months from the date of purchase when stored as follows: Solution I/RNaseAat4°Creceived, all other material at 22-25°C.

Kit Contents

Product Number	M1259-01	M1259-02
Purification times	5 Preps	20 Preps
Mag-Bind [®] Particle Solution	3 mL	11 mL
MGC Binding Buffer	2 x 200 mL	5 x200 mL
Solution I	450 mL	2 x 1000 mL
Solution II	450 mL	2 x 1000 mL
Neutralization Buffer	450 mL	2 x 1000 mL
SPM Wash Buffer Concentrate	200 mL	3 x 200 mL
RNase A Concentrate	3 mL	3 x 7 mL
Instruction Booklet	1	1

Before Starting

Briefly examine this booklet and become familiar with each step. Prepare all components and have the necessary materials ready before starting.

IMPORTANT	1. Add vial of RNase A to a bottle of Solution I and Store at 4° C.		
	 SPM Wash Buffer Concentrate has to be diluted with absolute ethanol (~96-100%)as follows: 		
	M1259-01 Add 800 mL ~ 96%-100% ethanol		
	M1259-02 Add 800 mL ~96%-100% ethanol per bottle		
	 MGC Binding Buffer has to be diluted with absolute ethanol (~96-100%) as follows: 		
	M1259-01 Add 800 mL ~96%-100% ethanol per bottle		
	M1259-02 Add 800 mL ~96%-100% ethanol per bottle		
	Store diluted SPM Wash Buffer & MGC Binding Bufefr at room temperature.		

E.Z.N.A.[™] Mag-Bind[®] Plasmid Mega Spin Protocol

Materials Supplied By User

- Centrifuge Capable of 15,000 x g
- Tubes or vessel capable of 15,000 x g
- 500mL centrifuge tube.
- Sterile deionized water (or TE buffer)
- Absolute (96%-100%) ethanol
- waterbath or heat block preset to 70°C
- Pipettor
- Isolate a single colony from freshlystreaked selective antibiotic plate and inoculate a starter culture of 2-5mL LB medium containing proper antibiotic. Incubate st37°C for about 8 hours with vigorously shaking. Dilute 1.5-3.0 mL starter culture into a 1000-1500 mL selective LB/antibiotic(s) medium and grow at 37°C with agitation for 12-16 h. The culture density should reach 3-4 x 10° per mL.

It is strongly recommended that an endAnegative strain of *E.coli* be used for routine plasmid isolation. Examples of such strains include DH5 α° and JM109°.

2. Harvest the bacterial cells by centrifugation at 6,000 x g for 15 minutes at 4° C.

Note: If you want to stop the protocol and continue later, discard the medium and freeze the cell pellet at -20 $^\circ\text{C}.$

- Discard supernatantinto a waste container. Drythe pellet byplacing centrifuge tube upside-down on a paper towel to remove excess media. Add 80 mL of Solution I/RNase A to the bacterial pellet. Resuspend cells completely by shaking or pipetting. Complete resuspension of cell pellet is vital for obtaining good plasmid yields.
- 4. Add 80 mL Solution II and mix by gently shaking and rotating the tube for 1 minute to obtain a cleared lysate. A 2-3 minutes incubation at room temperature maybenecessary. Avoid vigorous mixing as doing so will shear chromosomal DNA and lower plasmid purity. (Store Solution II tightly capped when not in use.)

Note: do not incubate the lysate over 5 minutes since it can cause permanently denature plasmid.

- 5. Add 80 mL of Chilled (4°C) Neutralization Buffer and mix by gently shaking and rotating for 1 minute until a flocculent white precipitate forms.
- 6. Centrifuge at \geq 15,000 x gfor 30 minutes at 4° C. Remove the supernatant contains plasmid DNA promptly to a new tube.
- 7. Carefully transfer the supernatant and clear it again by filtering through filter paper (Whatman #1 or autoclaved coffee filter) into a vessel or tube.
- 8. Divide the cleared cell lysate into 2 of 500 mL centrifuge bottles.

 Add 250 μl of Mag-Binds[®] Particles Solution into each tube and follow by addition of equal volume of MGC Binding Buffer diluted with absolute ethanol. Mix well by inverting the tube few times.

NOTE: The Mag-Binds[®] Particles will settle and bead together at the bottom of their container after several hours. Please check container before use. If beading has occurred, gentlyshake or vortex container until particles have been re-dispersed in solution. (IMPORTANT)

- 10. Incubate for 10 minutes at room temperature, mixing few times by inverting the bottle.
- 11. Centrifuge at 6000 x g for 10 minutes to pellet the magnetic particles. Discard the supernatant.
- 12. Add 50 mL of SPM Wash Buffer diluted with absolute ethanol into each tube. Resuspend the magnetic particles by pipetting or vortexing.
- 13. Combine the magnetic particles into one tube. Centrifuge at 6000 x g for 10 minutes to pellet the magnetic particles. Discard the supernatant.

NOTE: For better washing efficiency, Mag-Binds[®] particles should be fully resuspended. Resuspension can be performed by pipetting or by vortexing.

- 14. Add 40 mL of SPM Wash Buffer diluted with absolute ethanol. Resuspend the Mag-Binds[®] particles by vortexing. Transfer the suspended Mag-Bind particles into a 50 mLcentrifuge tube.
- 15. **Centrifuge at 6000 x g for 10 minutes to pellet the magnetic particles**. Discard the supernatant and remove any liquid by invert the tube on a absorbent paper.
- 16. Air dry the Mag-Binds[®] particles pellet for 5-10 minutes at room temperature.
- 17. Elute DNA: Resuspend the Mag-Binds[®] particles pellet with 2-5 mL Elution Bufferor TE buffer. Incubate at 60°C for 10 minutes.
- 18. Centrifuge at 6000 x g for 10 minutes to pellet the Mag-Binds[®] particles.
- 19. Transfer the supernatant containing the purified plasmid into a clean 50 mL centrifuge tube.

Mag-Bind[®] Plasmid Mega Magnetic Protocol

- 1. Culture the bacteria and prepare the cell lysate by following step 1-7 from Centrifugation protocol on page 4.
- 2. Add 500 µl of of Mag-Binds[®] Particles Solution into each tube and follow by addition of equal volume of MGC Binding Buffer. Mix well by inverting the tube few times.
- 3. Incubate for 10 minutes at room temperature, mixing few times by inverting the bottle.
- 4. Magnetize the magnetic particles with a magnet until liquid is cleared.
- 5. Add 70 mL of SPM Wash Buffe diluted with absolute ethanolr. Invert the tube few times to mix throughly.
- 6. Magnetize the magnetic particles with a magnet until liquid is cleared.
- 7. Discard the cleared supernatant. Wash the beads again by repeating step 5-6.
- 8. Aspirate the cleared supernatant and airdrythe magnetic pellet atroom temperature for 5-10 minutes.
- 9. Elute DNA: Resuspend the Mag-Binds[®] particles pellet with 2-5 mL Elution Buffer or TE buffer. Incubate at 60°C for 10 minutes.
- 10. Centrifuge at 6000 xg for 10 minutes to pellet the Mag-Binds[®] particles. Transfer the eluted DNA to a clean tube.

Troubleshooting Guide

Problem	Likely Cause	Suggestions
Low DNA yields	Poor cell lysis	Do not use more than 1.5L with high copy plasmids.
		Cells may not be dispersed adequately prior to addition of Solution II. Vortex cell suspension to completely disperse.
		Increase incubation time with Solution II to obtain a clear lysate.
		Solution II if not tightly closed, may need to be replaced. Prepare as follow s: 0.2 N NaOH, 1% SDS.
	Bacterial culture overgrown or not fresh.	Do not incubate cultures for more than 16 hr at 37°C. Storage of cultures for extended periods prior to plasmid isolation is detrimental.
	Low copy-number plasmidused	Such plasmids may yield as little as $0.1 \mu g$ DNA from a 1 mL overnight culture.
	Lost Mag-Bind Particles during operation	careful remove the supernatant when aspirating the supernatant during process.
No DNA eluted.	SPM Wash Buffer or MGC Binding Buffer is not diluted with absolute ethanol.	Prepare SPM Wash Buffer and MGC Binding Buffer as instructed on the label.
High molecular weight DNA contamination of product.	Over mixing of cell lysate upon addition of Solution II.	Do not vortex or mix aggressively after adding Solution II. Adequate mixing is obtained by simply inverting and rotating tube to cover w alls w ith viscous lysate.
Optical densities do not agree with DNA yield on agarose gel.	Trace contaminants eluted from column increase A ₂₈₀ .	Make sure to wash Mag-Bind pellet as instructed. Alternatively, rely on agarose gel/ethidium bromide electrophoresis for quantization.
RNA visible on agarose gel.	RNase A not added to Solution I.	Add 1 vial of RNase to each bottle of Solution I.
Plasmid DNA floats out of well while loading agarosegel	Ethanol not completely removed before elution.	Increase air dry time before elution step
Plasmid DNA will not perform in downstream application	Traces of ethanol remain on column prior to elution.	The DNA plate must be washed with absolute ethanol and dried before elution. Ethanol precipitation may be required follow ing elution.

Ordering Information

Product No.	Product Name	Description		
E.Z.N.A.™ Plasmid Mini System				
D6942-01/02 D6943-01/02	Plasmid Mini Kit I	lsolation of up to 30µg plasmid in 15 minutes		
D6945-01/02	Plasmid Mini Kit II	lsolation of up to 70µg plasmid in 15 minutes		
D7042-01/02 D7043-01/02	HP Plasmid Mini Kit I	lsolation of up to 30µg plasmid from end A+ bacterial in 25 minutes		
D7045-01/02	HP Plasmid Mini Kit II	lsolation of up to 70µg plasmid from end A+ bacterial in 25 minutes		
D6948-01/02	Endo-free Plasmid Kit I	lsolation of up to 30μg endotoxin free plasmid		
D6950-01/02	Endo-free Plasmid Kit II	lsolation of up to 70μg endotoxin free plasmid		
D3476-01/02 D3376-01/02	Yeast Plasmid Kit	Isolation of plasmid from yeast		
D6900-01/02	M13 isolation kit	Isolation of M13 DNA from culture		
E.Z.N.A.™ Plasmid Midi/Maxi Isolation System				
D6904-01/02	Plasmid Midi Kit	Isolation of $\ge 200 \mu g$ plasmid w ith midi column		
D6905-03/04	Fastfilter Plasmid Midi kit	Isolation of $\ge 200 \mu g$ plasmid under 30 min		
D6915- 01/03/04	Endo-free Fastfilter Plasmid Midi kit	lsolation of up to 200μg endotoxin-free plasmid in less than 60 minutes		
D6922-01/02	Plasmid Maxi Kit	Isolation \ge 200µg plasmid w ith maxi column		
D6924- 01/03/04	Fastfilter Plasmid Maxi kit	Isolation of \ge 1.5 mg plasmid under 30 min.		
D6926- 01/03/04	Endo-free Fastfilter Plasmid Maxiprep kit	lsolation of up to 1.5 mg endotoxin-free plasmid in less than 60 minutes		
E-Z 96 [®] Plasmid Isolation System				
D1097-01/02	E-Z 96 [®] Fastfilter Plasmid Isolation Kit	lsolation of plasmid in 96 w ell format w ith lysate clearance plate		
D1900-01	E-Z 96 M13 Isolation Kit	Isolation of M13 DNA in 96 w ell format		