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Introduction

The E.Z.N.A.™ 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 the Omega Bio-tek's proprietary HiBind® matrix 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 Plasmid Maxi Kit combines the power of HiBind® technology with the time-tested consistency of alkaline-SDS lysis of bacterial cells to deliver high quality DNA. Omega Bio-tek's DNA Maxi-columns facilitate the binding, washing, and elution steps thus enabling multiple samples to be simultaneously processed. Yields vary according to plasmid copy number, *E.coli* strain, and conditions of growth, but 100-200 mL of overnight culture in LB medium typically produces 500-1000 µg high-copy plasmid DNA. Up to 500 mL culture may be processed when working with low-copy number plasmids. The product is suitable for automated fluorescent DNA sequencing, restriction endonuclease digestion, transfection of mammalian cells, and other manipulations.

The E.Z.N.A.™ High Performance (HP) Plasmid Purification System is the modified version of E.Z.N.A. plasmid isolation system which is designed specially for those applications when high quality plasmid is required such as transfection, auto-sequencing, etc. It is also suitable for isolating plasmid from bacterial hosts (such as EndoA+ strains) with high level of endonuclease activity. The plasmid from this system has much better stability for long term storage.

Storage and Stability

All E.Z.N.A.™ Plasmid isolation components are guaranteed for at least 12 months from the date of purchase when stored as follows: Solution I/RNase A at 4°C and OB protease mixtures at -20°C after received, all other material at 22-25°C. E.Z.N.A.™ HP

Kit Contents

Product Number	D7022-00	D7022-01	D7022-02
HiBind® DNA Maxi Columns	2	5	20
50 mL Collection tubes	2	5	20
Solution I	20 mL	70 mL	250 mL
Solution II	20 mL	70 mL	250 mL
Solution III	30 mL	100 mL	360 mL
Equilibration Buffer	7 mL	18 mL	65 mL
Buffer HB	15 mL	60 mL	220 mL
DNA Wash Buffer Concentrate	12 mL	25 mL	100 mL
Elution Buffer	6 mL	40 mL	160 mL
OB Protease Mixture	6 mg	16 mg	64 mg
RNase A, Concentrate	100 µL	300 µL	1.2 mL
Instruction Booklet	1	1	1

Before Starting

IMPORTANT	Add vial of RNase A to bottle of Solution I and store at 4°C.
	Dilute OB Protease with deionized water or TE Buffer as follows. Store aliquots at -20°C.
	D7022-00 Dissolve with 195 µl water or Buffer TE D7022-01 Dissolve with 520 µl water or Buffer TE D7022-02 Dissolve with 2.1 mL water or Buffer TE
	Dilute DNA Wash Buffer with absolute ethanol as follows
	D7022-00 Add 48 mL ~96-100% absolute ethanol D7022-01 Add 100 mL ~96-100% absolute ethanol D7022-02 Add 400 mL ~96-100% absolute ethanol

E.Z.N.A.™ HP Plasmid Maxi Spin Protocol

Materials supplied by user

- Laboratory centrifuge equipped with **swinging-bucket** rotor.
- High speed centrifuge capable of 5,000 x g
- Sterile 50 mL centrifuge tubes.

This Protocol is designed to isolate 500-1200 µg of high Copy-Number plasmids or 50-400 µg of low Copy-Number Plasmids from 200 mL overnight cultures using E.Z.N.A.™ High Performance Plasmid Maxi Kit. For increasing yield of low Copy-Number plasmid, proceed as "Low Copy-Number Plasmids protocol" on page 8.

■ Growth of bacterial culture

1. **Culture volume: Inoculate 100-200 mL LB/ampicillin (50 µg/mL) medium placed in a 1-4 liter culture flask with *E.coli* carrying desired plasmid and grow at 37°C with agitation for 12-16 h.** For best results use overnight culture as the inoculum. It is strongly recommended that an *endA* negative strain of *E.coli* be used for routine plasmid isolation. Examples of such strains include DH5α® and JM109®.

Optimal growth conditions of bacteria is vital of obtaining maximal plasmid DNA yields. The best conditions are achieved by picking a single isolated colony from a freshly transformed or freshly plate to inoculate a 2-5mL starter culture containing the appropriate antibiotic. Incubate for ~8 hours at 37°C with vigorous shaking (~300rpm). Then used to inoculate appropriate volume of Pre-warmed liquid growth medium with antibiotic. Grow at 37°C for 12-16 hr with vigorous shaking(~300rpm).Using a flask or vessel with a volume of a least 3-4 times the volume of the culture and dilute the starter culture 1/500 to 1/1000 into growth medium.

Following overnight bacterial growth, an OD₆₀₀ of 1.5~2.0 indicates a well-grown culture. For the best result determination of OD₆₀₀ for each culture is recommended. it is important to dilute the bacterial culture (10 to 20 fold) to enable photometric measurement in the linear range between 0.1 and 0.5 OD₆₀₀. We recommend a bacterial density of between 2.0 and 3.0 at OD₆₀₀. When using nutrient-rich media, care should be taken ensure that the cell density does not exceed an OD₆₀₀ of 3.0.

If using a frozen glycerol stock as inoculum, streak it onto an agar plate containing the appropriate antibiotic for single colony isolation. Then picking a single colony and inoculate the 2-5mL starter culture as described above.

■ Lyse bacterial cells with alkaline-SDS Solution

2. **Pellet up to 100-200 mL bacteria in appropriate vessels by centrifugation at 3,500-5,000 × g for 10 min at room temperature.**
3. Decant or aspirate medium and discard. To ensure that all traces of the medium are removed, use a clean paper towel to blot excess liquid from the wall of the vessel. **To the bacterial pellet add 12 mL Solution I/RNase A.** Resuspend cells

completely by vortexing or pipetting up and down.
Complete resuspension of cell pellet is vital for obtaining good yield.

4. Transfer cell suspension to a 50 mL centrifuge tube capable of withstanding at least 12,000 xg (screw-cap polycarbonate or Corex® glass tubes will suffice). **Add 12.0 mL Solution II and 100 µL OB Protease Mixture, gently mix by inverting and rotating tube 7-10 times to obtain a cleared lysate.** Incubate 3-5 minutes at room temperature.

Avoid vigorous mixing as this will shear chromosomal DNA and lower plasmid purity. Prolonged incubation may lead to nicking of plasmid DNA. (Store Solution II tightly capped when not in use.)

5. **Add 17 mL Solution III, cover, and gently mix by inverting tube several times until a flocculent white precipitate forms.** Centrifuge at $\geq 12,000 \times g$ for 10 minutes at 4°C to pellet the cellular debris and genomic DNA.

Note: The Buffers must be mixed thoroughly. If the mixture appears still viscous, brownish and conglobated, more mixing is required to completely neutralize the solution. Complete neutralization of the solution is vital of obtaining good yields. Increasing centrifugation speed is helpful to completely remove the precipitated bacterial cell material. A tightly packed cell debris pellet indicates efficient lysis.

6. **Take a HiBind® DNA Maxi column pre-inserted in a 50 ml collection tube. Add 3 ml Equilibration Buffer into the column. Let the column sit for 4 minutes at room temperature to equilibrate the membrane. Centrifuge at 3000 x g for 3 minutes.**

Note: Step 7 to 13 should be performed in swinging-bucket rotor for maximal plasmid DNA yields. And all centrifugation steps must be carried out at room temperature.

■ Purify Plasmid DNA with HiBind® DNA Maxi Column

7. **Carefully aspirate and add 20 mL of the clear supernatant to a clean HiBind® DNA Maxi column assembled in an 50 mL collection tube, making sure that no cellular debris is carried over.** The Maxi column has a maximum capacity of 20 mL. Centrifuge at 2,000-3,000 x g for 2 min at room temperature to completely pass lysate through column. Discard the flow-through liquid and add the remaining of cleared lysate to the column. Centrifuge as above and repeat until the entire sample has been passed through. Discard the flow-through and reuse the collection tube in Step 7.

8. **Add 10 mL Buffer HB to the HiBind® Maxi column and centrifuge as above.** This step ensures that residual protein contamination is removed and must be included for downstream applications requiring high quality DNA. Discard flow-through liquid and reuse the collection tube in the next step.

9. **Wash the column by adding 15 mL of DNA Wash Buffer diluted with ethanol. Centrifuge as above and discard flow-through.**

use. See label for directions. If refrigerated, DNA Wash Buffer must be brought to room temperature before use.

10. **Optional Step:** Repeat wash step with another 10 mL DNA Wash Buffer. Centrifuge as above and discard fluid.
11. **Centrifuge the empty column for 10 min at maxi speed (no more than 6,000 x g) to dry the column matrix.** Remove any traces of ethanol from the column's inner surface or O-ring using a pipette.

DO NOT skip this step - it is critical for removing traces of ethanol that may otherwise interfere with downstream applications.

■ Elution Plasmid DNA From HiBind® DNA Maxi column

Optional: For maximal yield and high concentration of plasmid, see alternative protocol of elution on page 7. For fast elution, proceed step 12-13

12. **Further Drying The Column (Optional).** Choose either of the methods below to further dry the column before eluting DNA (only if necessary):
 - A. Place the column into a vacuum container to dry the ethanol for 10 minutes. Then, remove the column and place into a vacuum chamber at room temperature. Any device connected to a vacuum source may be used. Seal the chamber and apply vacuum for 15 min. Remove the column and proceed to Step 12.
 - B. Bake the column in a vacuum oven or incubator at 65°C for 10 minutes. Remove the column and proceed to Step 12.
13. **Place column into a clean 50 mL centrifuge tube. Add 2-3 mL (depending on desired concentration of final product) Elution Buffer (10mM Tris-HCl, pH 8.5) or water directly onto the column matrix.** Allow column to sit for 2 min at room temperature. Centrifuge at maxi speed (no more than 6,000 xg) for 2 min to elute DNA. This represents approximately 70-80% of bound DNA. An optional second elution will yield any residual DNA, though at a lower concentration. Alternatively, a second elution may be performed using the first eluate to maintain a high DNA concentration. Also, preheating the water to 70°C prior to elution may significantly increase yields.

Note: The plasmid DNA obtained using this protocol performs well in PCR, restriction digests, lipid mediated transfection and transformation. The expected concentration of plasmid is vary between different copy number vector. However, the concentration of high copy-number plasmid is 150-400ug/mL. Some residual ethanol is present, but does not interfere with these downstream applications. One may get high concentration and absolutely remove ethanol with optional elution step as following.

~~Note: DNA Wash Buffer Concentrate must be diluted with absolute ethanol before~~

Alternative Elution Step from Column

1. Place HiBind™ DNA Maxi column into a clean 50 mL centrifuge tube. Add 6 mL Elution Buffer (or water) directly onto the column matrix. Allow column to sit 2 min at room temperature. Centrifuge at maxi speed (no more than 8,000 x g) to elute DNA. Preheating the water to 70°C and allowing the column to soak for 2 min at room temperature before elution may significantly increase yields.
2. Carefully transfer eluted plasmid to a new 50 mL centrifuge tube for precipitation. Add 1/10 volume of NaCl and equal volume of isopropanol. Vortex to mix and centrifuge at >15,000 × g for 30 min at 4°C. Carefully decant the supernatant.
3. Wash DNA pellet once with 4 mL ice-cold 70% ethanol and centrifuge at > 15,000 × g for 10 min. Carefully decant the supernatant without disturbing the pellet.
4. Air-dry the pellet for 5-10 min.
5. Finally resuspend DNA pellet in 0.5-1 mL (depending on desired concentration of final product) TE Buffer or water.

Protocol 2. HP Plasmid Maxi Kit Vacuum/Spin Protocol

1. Prepare cleared cell lysate by following step 1-5 of Spin procedure on page 4-5.
2. Transfer cleared cell lysate into the HiBind® DNA Maxi column, be careful not to overfill the column, apply the vacuum to allow all sample pass through the column. Repeat transfer the lysate into the column until the entire sample has been passed through.
3. Add 10.0 mL HB buffer to the column and apply the vacuum to draw the liquid through the column.
4. Wash the column: add 15 mL of DNA wash buffer (pre-diluted with absolute ethanol) into the column and allow it pass through the column.
5. Wash the column again with 10 mL DNA wash buffer by repeating step 4. Keep the vacuum on for another 10 minutes after the liquid pass through the column. (This step to ensure the removal of residue ethanol).
6. Proceed Elution Step as Spin protocol Fast Elution Step on Page 6 or Alternative elution step on page 7.

Protocol 3: Low Copy-Number Plasmids Protocol


Low copy plasmids generally give 0.1-1 µg DNA per mL overnight culture. For isolation of plasmid from low copy-number plasmids (0.1-1 µg/mL culture) or low-midi copy-number plasmids (1-2 µg/mL culture) bacteria, the method can be modified to essentially increase the yield if necessary.

~~Start with 400-500 mL bacterial culture, centrifuge for 10 min at 3,500-5,000 xg in a~~

centrifuge tube. Proceed to Step 3 (Page 4) and double the volumes of Solution I, II, III and OB Protease Mixture. Continue as above using only one HiBind® DNA Maxi column. There is no need to increase the volumes of Buffer HB and DNA Wash Buffer used. The Buffer of Solution I, II, III and OB Protease Mixture can be purchase separately.

Note: This method is not recommended for high copy number plasmids because above 500 mL culture, the HiBind® DNA Maxi column quickly becomes saturated. In this situation we recommend processing of multiple samples from the same culture.

Protocol 4: Short Protocol For Experienced Users



1. Pellet cells from 100-200 mL (high copy) or 400-500 mL (low copy) overnight culture.
2. Resuspend cells in 12 mL (high copy) or 24 mL (low copy) Solution I/RNase A.
3. Add 12 mL (high copy) or 24 mL (low copy) Solution II and 100 µl/200 µl OB Protease mixture. Mix gently but thoroughly by inverting 4-6 times to obtain cleared lysate. incubate at RT for 10 min.
4. Add 17 mL (high copy) or 32 mL (low copy) Solution III and mix well to form white precipitate.
5. Centrifuge at maximum (at least 12,000 x g) speed 10 min at 4°C.
6. Transfer cleared lysate to a HiBind® DNA Maxi column placed in a 50 mL collection tube. Centrifuge 1 min at 2000-3000 x g for 2 min. Discard liquid. Repeat this step until the entire sample has been passed through.
7. Wash column with 10 mL Buffer HB. Centrifuge as above. Discard liquid.
8. Using same collecting tube, wash column with 15 mL DNA Wash Buffer diluted with ethanol. Centrifuge as above.
9. Optional: Wash column a second time with 10 mL DNA Wash Buffer. Centrifuge as above.
10. Centrifuge the empty column for 10 min at max speed to dry.
11. Elute plasmid with 2-3 mL Elution Buffer or water. And Centrifuge at max speed for 2 min.
12. Remove the column from collection tube and store the eluted DNA at -20°C.

Yield and quality of DNA

Determine the absorbance of an appropriate dilution (10- to 20-fold) of the sample at 260 nm and then at 280 nm. The DNA concentration is calculated as follows:

$$\text{DNA concentration} = \text{Absorbance}_{260} \times 50 \times (\text{Dilution Factor}) \mu\text{g/mL}$$

The ratio of $(\text{Absorbance}_{260})/(\text{Absorbance}_{280})$ is an indication of nucleic acid purity. A value greater than 1.8 indicates > 90% nucleic acid. Alternatively, quantity (as well as quality) can sometimes best be determined by agarose gel/ethidium bromide electrophoresis by comparison to DNA samples of known concentrations. Typically, the majority of the DNA eluted is in monomeric supercoil form, though concatamers may also be present.

Plasmid Copy-Number and Yield

The Yield and quality of the plasmid DNA depend on a number of factors including plasmid copy number, size of insert, host strain, culture volume, culture medium and binding capacity of kit. In these factors, The copy number of vector, culture volume and binding capacity of kit are most important. Copy number of plasmid is vary from one copy to several hundred copies per cell as dictated by their origin of replication. But very large plasmids are often maintained at very low copy number per cell.

Plasmid	Replicon	Copy Number	Expected Yield of 200mL culture
pUC vector	pMB1	500-700	700-900 μg
pBR322 and its derivatives	pMB1	15-20	40-80 μg
PACYC and its derivatives	p15A	10-12	20-100 μg
pSC101 and its derivatives	pSC101	~5	20-80 μg
pBluescript	ColE14	300-500	400-600 μg
ColE14	ColE14	15-20	20-80 μg
pGEM	pMB1	300-700	600-800 μg

Troubleshooting Guide

Problem	Likely Cause	Suggestions
Low DNA yields	Poor cell lysis	Only use LB or YT medium containing ampicillin. Do not use more than 500 mL. Cells may not be dispersed adequately prior to addition of Solution II. Vortex/ pipet cell suspension to completely disperse bacterial clumps. Increase incubation time with Solution II to obtain a clear lysate. Solution II if not tightly closed, may need to be replaced. Prepare as follows: 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 to yield and quality.
	Low copy-number plasmid used	Such plasmids may yield as little as 0.1 μg DNA from a 1 mL overnight culture. Increase culture volume to 500 mL.
No DNA eluted.	DNA Wash Buffer Concentrate not diluted with absolute ethanol.	Prepare Wash Buffer Concentrate 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 walls with viscous lysate.
Optical densities do not agree with DNA yield on agarose gel.	Trace contaminants eluted from column increase A_{260} .	Make sure to wash column as instructed in steps 7-9. Alternatively, rely on agarose gel/ethidium bromide electrophoresis for quantitation.
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 agarose gel, does not freeze, or smells of ethanol.	Ethanol traces not completely removed from column following wash steps.	Centrifuge column as instructed in step 10 or vacuum as indicated to dry. A swinging-bucket rotor is recommended for centrifugation. Alternatively, precipitate the eluted DNA with isopropanol as indicated in step .

Ordering Information

Product No.	Product Name	Description
D6942-01/02 D6943-01/02	Plasmid Mini Kit I	Isolation of up to 30µg plasmid in 15 minutes
D6945-01/02	Plasmid Mini Kit II	Isolation of up to 70µg plasmid in 15 minutes
D7042-01/02 D7043-01/02	HP Plasmid Mini Kit I	Isolation of up to 30µg plasmid from end A+ bacterial in 25 minutes
D7045-01/02	HP Plasmid Mini Kit II	Isolation of up to 70µg plasmid from end A+ bacterial in 25 minutes
D6948-01/02	Endo-free Plasmid Kit I	Isolation of up to 30µg endotoxin free plasmid
D6950-01/02	Endo-free Plasmid Kit II	Isolation 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 ≥200µg plasmid with midi column
D6905-03/04	Fastfilter Plasmid Midi kit	Isolation of ≥200µg plasmid under 30 min
D6915-01/03/04	Endo-free Fastfilter Plasmid Midi kit	Isolation of up to 200µg endotoxin-free plasmid in less than 60 minutes
D6922-01/02	Plasmid Maxi Kit	Isolation ≥200µg plasmid with maxi column
D6924-01/03/04	Fastfilter Plasmid Maxi kit	Isolation of ≥ 1.5 mg plasmid under 30 min.
D6926-01/03/04	Endo-free Fastfilter Plasmid Maxiprep kit	Isolation 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	Isolation of plasmid in 96 well format with lysate clearance plate
D1900-01	E-Z 96 M13 Isolation Kit	Isolation of M13 DNA in 96 well format