

Plasmid Miniprep Kit II

(BW-PD1213)

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Kit Contents

Catalog#	BW-PD1213 -00	BW-PD1213 -01	BW-PD1213 -02	BW-PD1213 -03
Preps	10	50	100	250
Mini Columns	10	50	100	250
2 mL Collection Tubes	10	50	100	250
Buffer A1	5 mL	25 mL	47 mL	120 mL
Buffer B1	5 mL	25 mL	47 mL	120 mL
Buffer N1	6 mL	30 mL	57 mL	150 mL
Buffer KB	6 mL	28 mL	52 mL	135 mL
DNA Wash Buffer*	3 mL	15 mL	2×15 mL	3×24 mL
Elution Buffer	1.8 mL	8 mL	17 mL	40 mL
RNase A (20 mg/mL)	25 µL	125 µL	235 µL	600 µL
User Manual	1	1	1	1

*Add 12 mL (BW-PD1213-00) or 60 mL (BW-PD1213-01) or 60 mL (BW-PD1213-02) or 96 mL (BW-PD1213-03) 96-100% ethanol to each DNA Wash Buffer bottle before use.

Introduction

Key to the kit is our proprietary DNA binding systems that allow the high efficient binding of DNA to the Mini Column while proteins and other contaminants are removed under certain optimal conditions. Nucleic acids are easily eluted with sterile water or Elution Buffer.

This kit is designed for fast and efficient purification of plasmid DNA from 5 to 12 mL of *E. coli* culture. The maximum binding capacity is 120 µg per column. The yield from 1 mL culture is typically around 8 to 12 µg.

The purified DNA is ready for downstream applications such as cloning/subcloning, RFLP, sequencing, and transfection of HEK293 cells.

Important Information

Plasmid Copy Numbers: The yield of plasmid DNA depends on the origin of the replication and the size of the plasmid. The protocols are optimized for high copy number plasmid purification. For low copy number plasmids, both the culture volume and the buffer volume need to be scaled up accordingly. Please contact our customer service for further information and refer to Table 1 for the commonly used plasmids.

Table 1 Commonly used plasmids and expected yield.

Plasmid	Origin	Copy Numbers	Expected Yield (µg per 1 mL)
pSC101	pSC101	5	0.1-0.2
pACYC	p15A	10-12	0.4-0.6
pSuperCos	pMB1	10-20	0.4-1
pBR322	pMB1	15-20	0.6-1
pGEM ^R	Muted pMB1	300-400	6-7
pBluescript ^R	ColE1	300-500	6-8
pUC	Muted pMB1	500-700	8-12

Host Strains: The strains used for propagating plasmid have significant influence on yield. The strains used for propagating plasmid have significant influence on yield. Host strains such as Top 10 and DH5α yield high-quality plasmid DNA. *EndA*⁺ strains such as JM101, JM110, HB101, TG1 and their derivatives, normally have low plasmid yield due to either endogenous endonucleases or high carbohydrates released during lysis. We recommend transform plasmid to an *endA*⁻ strain if the yield is not satisfactory.

Table 2 *endA* strains of *E.coli*.

<i>End A</i> ⁻ strains of <i>E.coli</i>							
DH5α	DH1	DH21	JM106	JM109	SK2267	SRB	XLO
TOP10	DH10B	JM103	JM107	SK1590	MM294	Stbl2 TM	XL1-Blue
BJ5182	DH20	JM105	JM108	SK1592	Select96 TM	Stbl4 TM	XL10-Gold
<i>End A</i> ⁺ strains of <i>E.coli</i>							
C600	JM110	RR1	ABLE [®] C	CJ236	KW251	P2392	BL21(DE3)
HB101	TG1	TB1	ABLE [®] K	DH12S TM	LE392	PR700	BL21(DE3) pLysS
JM101	JM83	TKB1	HMS174	ES1301	M1061	Q358	BMH71-18
All NM Strains				All Y Strains			

Optimal Cell Mass (OD₆₀₀ × mL of Culture): This procedure is designed for isolating plasmid grown in standard LB medium (Luria Bertani) for 12-16 hours to a density of OD₆₀₀ 2.0 to 3.0. If rich medium such as TB or 2× YT are used, make sure the cell density doesn't exceed 3.0 (OD₆₀₀). A high ratio of biomass over lysis buffers result in low DNA yield and purity. The Mini Column has an optimal biomass of 30-45. For example, if the OD₆₀₀ is 3.0, the optimal culture volume should be 10-15 mL. For over amount of cell numbers, either reduce the biomass or scale up the volume of Buffer A1, B1 and N1.

Culture Volume: Use a flask or tube 4 times bigger in volume than the culture medium to secure optimal condition for bacteria growth. Don't exceed the maximum culture volume suggested in the protocol. Incomplete lysis due to over amount of bacterial culture results in lower yield and purity.

Storage and Stability

Buffer A1 should be stored at 4°C once RNase A is added. All other materials can be stored at room temperature (15-25°C). The guaranteed shelf life is 12 months from the date of production.

Before Starting

Prepare all components and get all necessary materials ready by examining this user manual and become familiar with each step.

Important Notes

☉RNase A: 20 mg/mL. It is stable for one year at room temperature (15-25°C). Spin down RNase A vial briefly. Buffer A1 should be stored at 4°C once RNase A is added.

☉Add 12 mL (BW-PD1213-00) or 60 mL (BW-PD1213-01) or 60 mL (BW-PD1213-02) or 96 mL (BW-PD1213-03) 96-100% ethanol to each DNA Wash Buffer bottle before use.

☉Buffer B1 precipitates below room temperature. It is critical to warm up the buffer at 37°C to dissolve the precipitates before use. Keep the cap tightly closed for Buffer B1 after use.

☉Ensure the availability of centrifuge capable of 12,000 rpm.

☉*Carry out all centrifugations at room temperature.*

Materials not Supplied

- ⊗ High speed microcentrifuge or vacuum manifold.
- ⊗ 96-100% ethanol.
- ⊗ 1.5 mL, 2 mL microcentrifuge tubes.
- ⊗ 15 mL conical tubes.

Safety Information

Buffer N1 contains chaotropic salts, wear gloves and protective eyewear when handling.

Buffer N1 and KB contains chaotropic salts, which may form reactive compounds when combines with bleach. Do not add bleach or acidic solutions directly to the waste.

Protocol (For spin)

For *1-4 mL* culture, reduce the volume of **Buffer A1**, **B1**, **N1** to **250 μ L**, **250 μ L** and **350 μ L**, respectively. And use the same volume of **DNA Wash Buffer** and **Elution Buffer**.

1. Inoculate **5-12 mL** LB containing appropriate antibiotic with a fresh colony. Incubate at 37°C for 14-16 hours with vigorous shaking.

Note: Prolonged incubation (>16 hours) is not recommended since the *E. coli* starts to lyse and the plasmid yield may be reduced.

Note: Do not grow the culture directly from the glycerol stock.

Note: This protocol is optimized for *E. coli* strain cultured in LB medium. When using TB or 2 x YT medium, special care needs to be taken to ensure the cell density doesn't exceed 3.0 (OD₆₀₀). Buffers need to be scaled up proportionally if over amount of cultures are being processed.

2. Harvest the bacterial culture by centrifuge for 1 minute at 12,000 rpm. Pour off the supernatant and blot the inverted tube on a paper towel to remove residue medium. Remove the residue medium completely.

Note: The culture can be centrifuged at 6,000 rpm in a 15 mL conical tube for 10 minutes if high speed centrifuge tubes are not available. Alternatively, the cultures can also be spin down in multiple 2.0 mL tubes.

3. Add **450 μ L Buffer A1** (*Add RNase A to Buffer A1 before use*) and completely resuspend bacterial pellet by vortexing or pipetting.

Note: Complete resuspension is critical for bacterial lysis and lysate neutralization.

4. Add **450 μ L Buffer B1**, mix gently by inverting the tube 10 times (*do not vortex*), and incubate at room temperature for 5 minutes. If necessary, continue inverting the tube until the solution becomes slightly clear.

Note: Do not incubate for more than 5 minutes.

Note: Buffer B1 precipitates (cloudy look) below room temperature. Warm up Buffer B1 at 37°C to dissolve precipitations before use.

5. Add **550 μ L Buffer N1**, mix completely by inverting the vial for 5-10 times.

Note: Incubating the lysate in ice for 1 min will improve the yield.

Note: It is critical to mix the solution well. If the mixture still appears conglobated, brownish or viscous, more mixing is required to completely neutralize the solution.

6. Centrifuge the lysate at 12,000 rpm for 10 minutes.

Note: If the lysate doesn't appear clean, reverse the tube angle, centrifuge for 5 more minutes and then transfer the clear lysate to Mini Column.

7. Insert a **Mini Column** into a **2 mL Collection Tube**.
8. Carefully transfer up to **700 µL** clear lysate from step **6** into the **Mini Column** with collection tube, avoid the precipitations, spin at 12,000 rpm for 1 minute, discard the flow-through and put the **Mini Column** back to the collection tube. Carefully transfer the remaining clear lysate to the **Mini Column** and centrifuge at 12,000 rpm for 1 minute and discard the flow-through in the collection tube. Put the **Mini Column** back to the collection tube.
9. **Optional:** Add **500 µL Buffer KB** into the **Mini Column**, centrifuge at 12,000 rpm for 1 minute and discard the flow-through. Put the column back to the collection tube.

Note: This step is NOT necessary if the plasmid is being purified from *endA*- strain such as DH5α and TOP10. Buffer KB is necessary for *endA*+ strains such as HB101, JM110, JM101 and their derived strains.

10. Add **600 µL DNA Wash Buffer** (*Add ethanol to DNA Wash Buffer before use*) into the **Mini Column**, centrifuge at 12,000 rpm for 1 minute and discard the flow-through.

Optional: Repeat step **10**.

11. Reinsert the **Mini Column**, *with the lid open*, into the collection tube and centrifuge for 2 minutes at 12,000 rpm.

Note: Residual ethanol can be removed more efficiently with the column lid open. It is critical to remove residual ethanol completely.

12. Carefully transfer the **Mini Column** into a clean 1.5 mL tube and add **100-150 µL** sterile ddH₂O or **Elution Buffer** into the center of the column and let it stand for 1 minute. Elute the DNA by centrifugation at 12,000 rpm for 1 minute.

Optional: Reload the eluate into the column and centrifuge again to improve the recovery.

Note: The first elution normally yields around 70% of the plasmid DNA. Add the eluted DNA back to the column for another elution yields 20-30% of the DNA.

Note: The DNA is ready for downstream applications such as cloning/subcloning, RFLP, library screening, *in vitro* translation, sequencing, transfection of robust cells such as HEK293 cells.

Note: It's highly recommended to remove the endotoxin (PD1212) if the DNA is used for endotoxin-sensitive cell lines, primary cultured cells or microinjection.

13. The DNA concentration can be calculated as follows,

Concentration (µg/mL)=OD₂₆₀×50×dilution factor.

Protocol (For spin/vacuum)

1. Set up the vacuum manifold according to manufacturer's instruction and connect the **Mini Column** to the manifold.
2. Carry out step **1-6** on page **6** in previous protocol.
3. Carefully transfer the clear lysate to the **Mini Column** and turn on the vacuum to allow the lysate pass through the column.

4. **Optional:** Add **500 μ L Buffer KB** into the **Mini Column** and allow the lysate pass through the column by vacuum.

Note: This step is NOT necessary if the plasmid is being purified from *endA*- strain such as DH5 α and TOP10. Buffer KB is necessary for *endA*+ strains such as HB101, JM110, JM101 and their derived strains.

5. Add **600 μ L DNA Wash Buffer** to the **Mini Column** and allow the vacuum to draw the liquid through the manifold. Turn off the vacuum.

Optional: Repeat step **5**.

6. Turn on the vacuum, dry the empty column for 5 minutes.

7. Carefully transfer the **Mini Column** into a clean 1.5 mL tube and add **100-150 μ L** sterile ddH₂O or **Elution Buffer** into the center of the column and let it stand for 1 minutes. Elute the DNA by centrifugation at 12,000 rpm for 1 minute.

Optional: Reload the eluate into the column and centrifuge again to improve the recovery.

Note: We recommended Elution Buffer instead of ddH₂O for stability of plasmid DNA. If ddH₂O is applied, please make sure the pH is no less than 7.0 (7.0-8.5 is preferred). NaOH could be used to adjust the pH of ddH₂O. For long term storage, please use Elution Buffer.

Note: The DNA is ready for downstream applications such as cloning, RFLP, library screening, *in vitro* translation, sequencing, transfection of robust cells such as HEK293 cells.

Purification of Low-Copy-Number Plasmid/Cosmid

The yield of low copy number plasmid is normally around 0.1-1 µg/mL of overnight culture. For isolating low copy number or medium copy number plasmid DNA, use the following guideline:

- ☉ Culture volume: Use **2×volumes** of the high copy number culture.
- ☉ Use **2×volumes** of the **Buffer A1**, **Buffer B1** and **Buffer N1**. Additional buffers can be purchased from Biomiga.
- ☉ Use **same volume** of **DNA Wash Buffer** and **Elution Buffer**.

Trouble Shooting Guide

Problems	Possible Reasons	Suggested Improvements
Low yield	Poor cell lysis.	Resuspend pellet thoroughly by vortexing and pipetting prior to adding Buffer B1. Make fresh Buffer B1 if the cap had not been closed tightly. (Buffer B1: 0.2 M NaOH and 1% SDS).
Low yield	Bacterial culture overgrown or not fresh.	Grow bacterial 12-16 hours. Spin down cultures and store the pellet at -20°C if the culture is not purified the same day. Do not store culture at 4°C overnight.
Low yield	Low copy number plasmid.	Increase culture volume and the volume of Buffer A1,B1,N1 as instructed on page 9.
No DNA	Plasmid lost in host <i>E. coli</i> .	Prepare fresh culture.
Genomic DNA contamination	Over-time incubation after adding Buffer B1.	Do not vortex or mix aggressively after adding Buffer B1. Do not incubate more than 5 minutes after adding Buffer B1.
RNA contamination	RNase A not added to Buffer A1.	Add RNase A to Buffer A1.
Plasmid DNA floats out of wells while running in agarose gel	Ethanol traces were not completely removed from column.	Make sure that no ethanol residue remains in the silicon membrane before elute the plasmid DNA. Re-centrifuge or vacuum again if necessary.

Limited Use and Warranty

This product is intended for *in vitro* research use only. Not for use in human.

This product is warranted to perform as described in its labeling and in BEIWO's literature when used in accordance with instructions. No other warranties of any kind expressed or implied, including, without limitation, implied warranties of merchantability or fitness for a particular purpose, are provided by BEIWO. BEIWO's sole obligation and purchaser's exclusive remedy for breach of this warranty shall be, at the option of BEIWO, to replace the products, BEIWO shall have no liability for any direct, indirect, consequential, or incidental damage arising out of the use, the results of use, or the inability to use it product.

For technical support or learn more product information, please contact us or visit our website.



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