



INSTRUCTION MANUAL

ZymoPURE[™] II Plasmid Gigaprep Kit

Catalog Nos. D4204 (Patent Pending)

Highlights

- Fast, easy, reliable, ultra-pure transfection grade plasmid DNA Gigaprep using a spin-column.
- Innovative EndoZero[™] technology enables transfection in sensitive cells and *in vivo* research (≤ 0.025 EU/µg of plasmid DNA).
- State-of-the-art ZymoPURE binding technology guarantees highly concentrated plasmid DNA directly from a spin-column in ≤ 50 minutes. No gravity filtration or ethanol precipitation!

Contents

Product Contents	1
Product Specifications	1
Product Description	2
Procedure Overview	3
Buffer Preparation	4
Protocols	4-5
Troubleshooting	6
Ordering Information	7
Related Products	8-9

For Research Use Only

Version 1.1.0

Satisfaction of all Zymo Research products is guaranteed. If you are dissatisfied with this product, please call 1-888-882-9682.

Product Contents:

ZymoPURE [™] II Plasmid Gigaprep Kit (Kit Size)	D4204 (5 preps.)	Storage Temperature
ZymoPURE [™] P1 ¹ (Red)	2x 410 ml	4°C
ZymoPURE [™] P2 ² (Green)	2x 410 ml	Room Temp.
ZymoPURE [™] P3 (Yellow)	2x 410 ml	Room Temp.
ZymoPURE [™] Binding Buffer	2x 410 ml	Room Temp.
ZymoPURE [™] Wash 1	410 ml	Room Temp.
ZymoPURE [™] Wash 2 (Concentrate)	5x 23 ml	Room Temp.
ZymoPURE [™] Elution Buffer	30 ml	Room Temp.
Zymo-Spin [™] VI-P Columns	5	Room Temp.
600 ml Reservoir	5	Room Temp.
ZymoPURE [™] Giga Filter	5	Room Temp.
EndoZero [™] Spin-Columns	5	Room Temp.
15 ml Conical Reservoirs	5	Room Temp.
Instruction Manual	1	-

Note - Integrity of kit components is guaranteed for up to one year from date of purchase. Reagents are routinely tested on a lot-to-lot basis to ensure they provide maximal performance and reliability.

- ¹ ZymoPURE[™] P1 contains RNase A (100 μg/ml) and is stable at room temperature without loss in RNase activity, however, for long-term storage the product should be stored at 4-8° C.
- ² Caution: ZymoPURE[™] P2 Buffer contains NaOH. Please use proper safety precautions.

Specifications:

- **DNA Purity:** Eluted DNA is ultrapure, endotoxin-free, and well suited for transfection, transformation, sequencing, restriction endonuclease digestion, *in vitro* transcription, *in vivo* studies, and other sensitive applications.
 - Typical $Abs_{260/280} \ge 1.8$ and $Abs_{260/230} \ge 2.0$
 - o Endotoxin levels: ≤ 1 EU/µg of plasmid DNA using the Standard Protocol

≤ 0.025 EU/µg of plasmid DNA with optional EndoZero[™] Spin-Column

- **Plasmid DNA Yield:** Up to 10 mg per preparation (Actual yield is dependent on the plasmid copy number, culture growth conditions, and strain of *E. coli utilized*)
- Plasmid DNA Size: Up to 200 kb
- Recovery Volume: ≥ 2 ml of ZymoPURE[™] Elution Buffer or DNase free water
- Required Equipment: Vacuum/vacuum manifold and swinging bucket centrifuge.
 - Processing Time: 50 min

Notes:

This product is for research use only and should only be used by trained professionals. It is not for use in diagnostic procedures. Some reagents included with this kit are irritants. Wear protective gloves and eye protection. Follow the safety guidelines and rules enacted by your research institution or facility.

[™] Trademarks of Zymo Research Corporation.

Several ZymoPURE m product technologies are subject to U.S. and foreign patents or are patent pending.

pGL3[™] is a registered trademark of Promega Corporation.

Product Descript

The **ZymoPURE**[™] **II Plasmid Gigaprep Kit** features a simple spin-column based method for the purification of up to 10 mg of transfection grade plasmid DNA in less than 50 minutes. The eluted plasmid DNA is EndoZero and ready for immediate use in the most sensitive applications. The unique ZymoPURE methodology removes the need for slow gravity flow anion-exchange columns, alcohol precipitations, lengthy endotoxin removal incubations, and time-consuming centrifugation steps.

ZymoPURE[™] technology uses a modified alkaline lysis method and features novel binding chemistry, which enables the highest yields and concentration of plasmid DNA (up to 3 µg/µl) directly from a spin-column. Coupling ZymoPURE with the innovative **EndoZero**[™] **Spin-Columns**, to eliminate endotoxins, achieves EndoZero plasmid DNA (≤ 0.025 EU/µg of plasmid DNA), making it suitable for transfection, restriction endonuclease digestion, *in vivo* studies, bacterial transformation, PCR amplification, DNA sequencing, and other sensitive downstream applications.

As an added convenience, the **ZymoPURE**[™] **II Plasmid Gigaprep Kit** contains colored buffers that permit error-free visualization and identification of complete bacterial cell lysis and neutralization. Also, bottle top filters are included for rapid clearing of the lysate.

Simplest Workflow

EZ-Elute"

No Alcohol

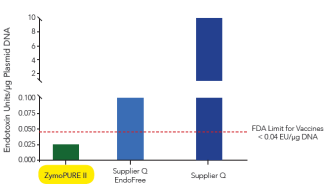
Precipitation!



Plasmid DNA concentration and yield from the ZymoPURE II Maxiprep kit compared to two separate kits from Supplier Q. Plasmid DNA (pGL3[®]) was isolated from 150 ml of JM109 E. *coli* culture grown overnight following the manufacturer's suggested protocol (in duplicate). One (1) µl of eluted plasmid DNA was visualized post agarose gel electrophoresis. M, ZR 1 kb DNA Marker (Zymo Research).

Highest Recovery

Lowest Endotoxin Levels



Manufacturers' stated endotoxin for two separate Anion-Exchange kits from Supplier Q compared to ZymoPURE II.

EZ-Load[™] No Gravity Flow!



EndoZero[™] Endotoxins < FDA limit for Vaccines

For **Technical Assistance**, please contact **Zymo** at 1-888-882-9682 or E-mail tech@zymoresearch.com.



Procedure Overview:

Bacterial cells are resuspended in ZymoPURE[™] P1 (red).

The solution will turn dark purple and viscous following the addition of ZymoPURE[™] P2 (green) indicating bacterial lysis is complete.

The solution will turn yellow and a precipitate will form after adding ZymoPURE[™] P3 (yellow) indicating neutralization is complete.

The neutralized lysate is loaded into the ZymoPURE[™] Giga Filter and clarified using a vacuum.

ZymoPURE[™] Binding Buffer is added to the cleared lysate and mixed thoroughly.



The mixture is loaded into the Zymo-Spin[™] VI-P Column using a vacuum manifold.



The Zymo-Spin[™] VI-P Column is washed using a vacuum manifold.



Ultra-pure plasmid DNA is eluted from the Zymo-Spin[™] VI-P Column using a centrifuge.



The eluted plasmid DNA is passed through the EndoZero[™] Column using a centrifuge.

Buffer Preparation:

- ✓ Add 88 ml of 95% ethanol to the 23 ml **ZymoPURE**^T Wash 2 (Concentrate) before use.
- ✓ The ZymoPURE[™] P2 and ZymoPURE[™] Binding Buffer may have precipitated. If this occurs, dissolve the precipitate by incubating the bottles at 30-37 °C for 10-20 minutes and mix by inversion. Do not microwave!

Before Starting:

✓ Centrifuge up to 2.5 liters of bacterial culture at ≥ 3,400 x g for 20 minutes to pellet the cells¹. Discard supernatant.

Protocol:

The following procedure should be performed at room temperature (15-30°C).

This product is compatible with any conventional vacuum-based manifold. The vacuum pump should be a single or double-staged unit capable of producing up to 400 mm Hg pressure at the vacuum manifold.

- 1. Add 150 ml of **ZymoPURE[™] P1 (Red)** to the bacterial cell pellet and resuspend completely by vortexing or pipetting.
- 2. Add 150 ml of **ZymoPURE[™] P2 (Green)** and immediately mix by gently inverting the tube 6 times. <u>Do not vortex!</u> Let sit at room temperature for 2-3 minutes².

Cells are completely lysed when the solution appears clear, purple, and viscous.

3. Add 150 ml of **ZymoPURE[™] P3 (Yellow)** and mix gently but thoroughly by inversion. <u>Do not vortex!</u>

The sample will turn yellow when the neutralization is complete and a yellowish precipitate will form.

- 4. Place the ZymoPURE[™] Giga Filter onto a 33 mm or 45 mm-neck glass bottle and load the lysate into the ZymoPURE[™] Giga Filter. Ensure the ZymoPURE[™] Giga Filter is resting securely on top of the glass bottle and wait 10 minutes for the precipitate to float to the top.
- Connect the ZymoPURE[™] Giga Filter to a vacuum source and turn on the vacuum³ until approximately 375-400 ml of cleared lysate is recovered. <u>Save the cleared</u> <u>lysate</u>!
- 6. Add 150 ml **ZymoPURE[™] Binding Buffer** to the cleared lysate from step 5 and mix thoroughly by inverting the capped bottle 10 times.

(Continued on next page)

Notes:

¹ A vessel with a minimum volume of 500 ml is required to prepare the bacterial lysate.

² Do not allow the lysis reaction to proceed for more than 3 minutes. Excessive lysis can result in denatured plasmid DNA.

³ Gently pressing down on the top of the **ZymoPURE**[™] **Giga Filter** when the vacuum is applied will guarantee an airtight seal between the filter and neck of the glass bottle.

Notes:

¹ To achieve optimal performance, the vacuum pump should be able to apply at least 400 mm Hg pressure. If less pressure is applied, centrifuge the column prior to washing to remove any residual lysate remaining in the matrix.

² The **ZymoPURE[™] Elution Buffer** contains 10 mM Tris-HCl, pH 8.5 & 0.1 mM EDTA. If required, pure water can also be used to elute the DNA.

³ The DNA yield can be increased by pre-warming the **ZymoPURE[™] Elution Buffer** to 50 °C and/or increasing the incubation period up to 10 minutes prior to centrifugation.

⁴ For low-copy number plasmids or if higher concentration is desired, the plasmid DNA can be eluted in as little as 2 ml.

⁵ For plasmid preparations with expected yields of 5 mg or greater, use 5 ml of ZymoPURE™ Elution Buffer to elute the plasmid DNA.

 ⁶ This optional step will reduce endotoxin levels from
 ≤ 1 EU/µg of plasmid DNA to
 ≤ 0.025 EU/µg of plasmid DNA.

- 7. Securely attach the **600 ml Reservoir** to the top of the **Zymo-Spin[™] VI-P Column** and place onto a vacuum manifold.
- With the vacuum off, add the entire mixture from step 6 into the 600 ml Reservoir/Zymo-Spin[™] VI-P Column Assembly, and then turn on the vacuum¹ until all of the liquid has passed completely through the column.
- 9. <u>With the vacuum off</u>, add 80 ml of **ZymoPURE**[™] **Wash 1** to the 600 ml Reservoir. Turn on the vacuum until all of the liquid has passed completely through the column.
- 10. <u>With the vacuum off</u>, add 50 ml of **ZymoPURE**[™] **Wash 2** to the 600 ml Reservoir. Turn on the vacuum until all of the liquid has passed completely through the column. <u>Repeat this wash step</u>.
- 11. Remove and discard the 600 ml Reservoir and place the Zymo-Spin^T VI-P Column in a 50 ml conical tube. Centrifuge at \geq 3,400 x *g* for 10 minutes in order to remove any residual wash buffer.
- 12. Transfer the column into a clean 50 ml conical tube and add 3 ml of **ZymoPURE**[™] **Elution Buffer**^{2,3,4,5} directly to the column matrix. Wait 5 minutes, and then centrifuge at ≥ 3,400 x g for 5 minutes.
- 13. Optional: For EndoZero Plasmid DNA⁶, ensure the clear Luer Lock cap attached to the EndoZero[™] Spin-Column is finger tight and securely attach the 15 ml Conical Reservoir to the top of the EndoZero[™] Spin-Column. Place the assembly into a clean 50 ml conical tube and add the entire eluate from Step 12 into the 15 ml Conical Reservoir/EndoZero[™] Spin-Column Assembly. Centrifuge at 2,000 *x g* for 10 minutes in a centrifuge. Store the eluted plasmid DNA at ≤ -20°C.

Troubleshooting Guide:

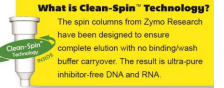
Problem	Possible Causes and Suggested Solutions
Low DNA Yield	
Culture growth conditions	 Poor aeration of culture. The optimal culture volume to air volume ratio is 1:5 o less. For best aeration, use baffled culture flasks, or a vented or gas-permeable sea on the culture vessel. The culture was overgrown, undergrown, contaminated, or antibiotics were omitted from the growth medium. Use a fresh culture for optimal performance. Ar OD₆₀₀ of 0.2-0.35 is the optimal optical density of a tenfold dilution of the culture.
Cell density is too high	 Too much culture used. Lysis and neutralization will be incomplete and the Zymo PURE[™] Giga Filter may clog during filtration. More culture does not always equa more plasmid. Incomplete lysis and neutralization are two of the most commor causes of failed plasmid preps and both are caused by too much culture being used. Incomplete lysis: After addition of ZymoPURE[™] P2, the solution should change from opaque pink to a clear viscous purple, indicating complete lysis. Different <i>E coli</i> strains often require different growth conditions and may vary in their susceptibility to alkaline lysis. Incomplete neutralization: The solution should not be viscous following neutralization and the yellowish precipitate should appear fluffy and readily float to the surface. Make sure the neutralization is complete prior to filtration. Invert the tube an additional 2-3 times after the sample turns yellow following the addition o ZymoPURE[™] P3.
Lysate Clarification	 Less than 375-400 ml of cleared lysate was recovered from the ZymoPURE[™] Giga Filter. For optimal performance, add 150 ml of ZymoPURE[™] Binding Buffer to approximately 375-400 ml of clarified lysate.
ZymoPURE P2 and ZymoPURE Binding Buffer precipitated	 Both buffers may have precipitated during shipping. To completely resuspend the buffers, incubate the bottles at 30-37 °C for 10 minutes and mix by inversion. DO NOT MICROWAVE.
Wash buffer	 Ensure that ethanol has been added to the ZymoPURE[™] Wash 2. Ensure that the bottle cap is screwed on tightly after each use to preven evaporation of the ethanol.
DNA elution	 Incomplete elution: For large size plasmids (> 10 kb), add ZymoPURE[™] Elution Buffer and incubate the column for 5-10 minutes before centrifugation. Also, pre warm the ZymoPURE[™] Elution Buffer to 50 °C prior to elution.
Low DNA Quality	
DNA does not perform well	 Incomplete neutralization: Incomplete neutralization generates poor qualit supernatant. Ensure that neutralization is complete by inverting the sample at additional 2-3 times after the addition of ZymoPURE[™] P3 and extending the incubation. Ethanol contamination in eluate. Centrifuge the Zymo-Spin[™] VI-P column a indicated in the protocol prior to adding the ZymoPURE[™] Elution Buffer.
RNA in eluate	 Ensure that ZymoPURE[™] P1 has been stored at 4°C. RNase A can be purchased separately if necessary.
Genomic DNA in eluate	 Improper handling (Sample was vortexed or handled too roughly). Genomic DNA contamination is usually caused by excessive mechanical shearing during the lysis and neutralization steps. Also, prolonged lysis or incomplete mixing of lysis o neutralization buffers may contribute to genomic DNA contamination in your sample. Overgrown culture. Overgrown or old cultures may contain more genomic DNA contamination than fresh cultures.

Ordering Information

Product Description	Kit Size	Catalog No.
ZymoPURE [™] II Plasmid Gigaprep Kit	5 preps.	D4204

For Individual Sale	Amount	Catalog No.
ZymoPURE [™] P1 (Red)	150 ml 210 ml 410 ml	D4200-1-150 D4200-1-210 D4200-1-410
ZymoPURE [™] P2 (Green)	150 ml 210 ml 410 ml	D4200-2-150 D4200-2-210 D4200-2-410
ZymoPURE [™] P3 (Yellow)	150 ml 210 ml 410 ml	D4200-3-150 D4200-3-210 D4200-3-410
ZymoPURE [™] Binding Buffer	150 ml 210 ml 410 ml	D4200-4-150 D4200-4-210 D4200-4-410
ZymoPURE [™] Wash 1	20 ml 55 ml 410 ml	D4200-5-20 D4200-5-55 D4200-5-410
ZymoPURE [™] Wash 2 (Concentrate)	10 ml 23 ml	D4200-6-10 D4200-6-23
ZymoPURE [™] Elution Buffer	6 ml 12 ml 30 ml	D4200-7-6 D4200-7-12 D4200-7-30
Zymo-Spin [™] VI-P	5	C1044-5
600 ml Reservoir	5	C1033-5
ZymoPURE [™] Giga Filter	1	C1038-1
EndoZero [™] Spin-Columns	5	C1051-5
15 ml Conical Reservoir	5 25	C1031-5 C1031-25





Purify DNA from PCR & other sources

DNA Clean & Concentrator[™] (DCC[™])

- ✓ Recovery of ultra-pure DNA that is free of salts and contaminants.
- ✓ Small (≥6 µl) elution volume.
- ✓ DNA is ideal for ligation, PCR, Next-Gen sequencing, etc.

Product	Size (Cat. No.)
DNA Clean & Concentrator [™] -5	50 Preps. (D4013) 200 Preps. (D4014)
ZR-96 DNA Clean & Concentrator [™] -5	2 x 96 Preps. (D4023) 4 x 96 Preps. (D4024)
Genomic DNA Clean & Concentrator™	25 Preps. (D4010) 100 Preps. (D4011)



DCC[™]-5 compared to Supplier Q .

Boost DNA recoveries from agarose gels to >80%

23 kb

2 kb 500 bp

Zymoclean[™] Gel DNA Recovery

- ✓ Rapid (15 min.) recovery of ultra-pure DNA from agarose gels in ≥6 μ l.
- ✓ Ultra-pure DNA ideal for DNA ligation, sequencing, etc.
- ✓ Format also available for large DNA >20 kb.

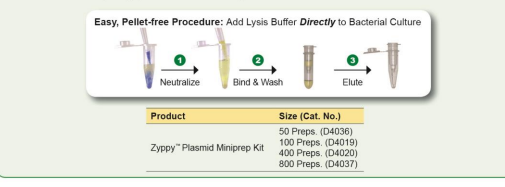
Product	Size (Cat. No.)
Zymoclean [™] Gel DNA Recovery Kit	50 Preps. (D4001) 200 Preps. (D4002)
Zymoclean [™] Large Fragment DNA Recovery Kit	25 Preps. (D4045) 100 Preps. (D4046)

DNA fragments recovered from an agarose gel using the Zymoclean[™] Gel DNA Recovery Kit. Lanes: M: DNA Ladder; 1-5: individual ladder DNA fragments.

Recover transfection-quality plasmid DNA directly from culture

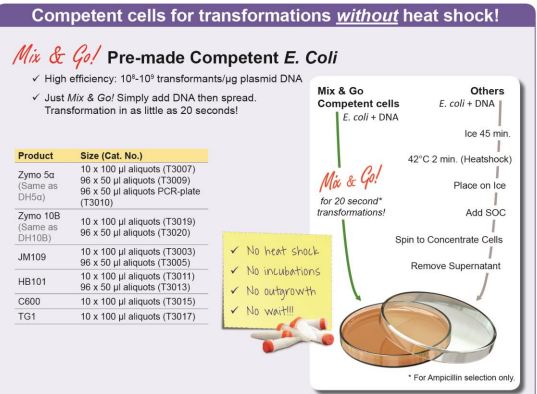
Zyppy[™] Plasmid Prep Kits

- ✓ The fastest, simplest method available for purifying high quality plasmid DNA from E. coli.
- ✓ Pellet-Free[™] procedure omits conventional cell-pelleting and resuspension steps.
- ✓ Transfection quality plasmid DNA directly from culture in under 15 minutes.



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OTHER INNOVATIVE PRODUCTS FROM ZYMO RESEARCH...



The fastest method for complete bisulfite conversion of DNA

EZ DNA Methylation-Lightning[™] Kits

- ✓ The next generation of bisulfite conversion technology by the most cited provider in the industry
- ✓ Guarantees high conversion efficiencies of cytosine (>99.5%)
- \checkmark Maintains the highest template integrity following bisulfite conversion
- Recovered DNA is ideal for PCR, MSP, array, bisulfite, and next-generation sequencing.

