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#### Feb 25, 2022

# Quick Protocol for Monarch® Plasmid Miniprep Kit (NEB #T1010) V.4

New England Biolabs<sup>1</sup>

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New England Biolabs (NEB)

Tech. support phone: +1(800)632-7799 email: info@neb.com

Isabel Gautreau New England Biolabs

This is the quick version of the Monarch® Plasmid DNA Miniprep Kit Protocol (NEB #T1010). For the full protocol, please click <u>here</u>.

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https://www.neb.com/protocols/2015/12/08/quick-protocol-for-monarch-plasmid-miniprep-kit-t1010

New England Biolabs 2022. Quick Protocol for Monarch® Plasmid Miniprep Kit (NEB #T1010). **protocols.io** 

https://dx.doi.org/10.17504/protocols.io.bg9qjz5w Isabel Gautreau

Monarch Kit, DNA, Plasmid, Lysis

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For detailed protocol and more information, visit <a href="https://www.neb.com/T1010">www.neb.com/T1010</a>

The full protocol is available here.

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The video protocol is available <u>here</u>.

#### **General Guidelines:**

Yield and quality of plasmid DNA is affected by plasmid copy number, plasmid size, insert toxicity, host strain, antibiotic selection, growth media and culture conditions. For standard cloning strains of  $\it E. coli$ , we recommend using a single colony from a freshly streaked selective plate to inoculate a standard growth media, such as LB (Luria-Bertaini) media. Cultures are typically grown at 37°C and 200–250 RPM in vessels that allow some aeration (Erlenmeyer flasks or culture tubes on a roller drum) and harvested after 12–16 hours as the culture transitions from logarithmic growth to stationary phase. This is the time at which the plasmid DNA content is highest. While cultures in LB often saturate with a final  $OD_{600}$  between 3–6, growth to saturation often leads to cell lysis. As a result, plasmid yields and quality are reduced and the likelihood of co-purifying unwanted host chromosomal DNA increases. Use of rich media, such as 2X YT or TB, produces higher biomass in a shorter time period. If chosen for growth, adjustments to the culture times and amount of cells used in the prep should be made to correct for these differences, and to avoid overloading the matrix and reducing DNA yield and quality.

Α	В	С	D
PLASMID	REPLICON	COPY	CLASSIFICATION
		NUMBER	
pUC and its derivatives	pMB1*	> 75	High copy
pBR322 and its derivatives	pMB1	15-20	Low copy
pACYC and its derivatives	p15A	10-12	Low copy
pSC101	pSC101	~5	

<sup>\*</sup>pUC and its derivatives lack the *Rop*gene and contain a point mutation in the RNAII transcript. These changes result in higher copy number during routine growth with many sources reporting levels as high as 500 copies per cell.

#### **Antibiotics for Plasmid Selection**

ANTIBIOTIC	CONCENTRATION OFSTOCK SOLUTION	STORAGETEMP.	WORKING CONCENTRATION
Ampicillin	100 mg/ml (H2O)	-20°C	50-200 μg/ml
Carbenicillin	100 mg/ml (H2O)	-20°C	20-200 μg/ml
Chloramphenicol	34 mg/ml (ethanol)	-20°C	25-170 μg/ml
Kanamycin	10 mg/ml (H2O)	-20°C	10-50 μg/ml
Streptomycin	10 mg/ml (H2O)	-20°C	10-50 μg/ml
Tetracycline	5 mg/ml (ethanol)	-20°C	10-50 μg/ml

#### **MATERIALS**

Monarch® Plasmid Miniprep Kit New England

**Biolabs Catalog #T1010** 

For hazard information and safety warnings, please refer to the SDS (Safety Data Sheet).

- All centrifugation steps should be carried out at <a>\$\@\$16000 x g</a> (~<a>\$\@\$13.000 rpm</a>).
- Add ethanol to Monarch Plasmid Wash Buffer 2 prior to use (4 volumes of ≥ 95% ethanol per volume of Monarch Plasmid Wash Buffer 2).
- For 50-prep kit add 

  24 mL of ethanol to 

  6 mL of Monarch Plasmid Wash Buffer
- 2. For 250-prep kit add **□120 mL** of ethanol to **□30 mL** of Monarch Plasmid Wash Buffer 2
- If precipitate has formed in Lysis Buffer (B2), incubate at § 30 °C § 37 °C, inverting periodically to dissolve.
- Store Plasmid Neutralization Buffer (B3) at § 4 °C after opening, as it contains RNase
   A.

## 1

Pellet  $\Box 1$  mL –  $\Box 5$  mL (not to exceed 15 OD units) bacterial culture by centrifugation at 316000 x g for 00:00:30. Discard supernatant.

■1.5 mL of culture is sufficient for most applications. Ensure cultures are not overgrown (12-16 hours is ideal).

### 2

Resuspend pellet in 200 µL Plasmid Resuspension Buffer (B1) (pink).

Vortex or pipet to ensure cells are completely resuspended. There should be no visible clumps.



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Add  $\blacksquare$ 200  $\mu$ L Plasmid Lysis Buffer (B2) (green), gently invert tube 5–6 times, and incubate at 8 Room temperature for  $\bigcirc$  00:01:00. Do not vortex.

Care should be taken not to handle the sample roughly and risk shearing chromosomal DNA, which will co-purify as a contaminant. Avoid incubating longer than one minute to prevent irreversible plasmid denaturation.



Add  $\Box 400~\mu L$  Plasmid Neutralization Buffer (B3) (yellow), gently invert tube until neutralized, and incubate at & Room temperature for @00:01:00. Do not vortex.

Be careful not to shear chromosomal DNA by vortexing or vigorous shaking. Firmly inverting the tube promotes good mixing, important for full neutralization.



Centrifuge lysate at **316000 x g** for 2-5 minutes.

Spin time should not be less than 2 minutes. Careful handling of the tube will ensure no debris is transferred and the 2 minute recommended spin can be successfully employed to save valuable time. For culture volumes > 1 ml, we recommend a 5 minute spin to ensure efficient RNA removal by RNase A. Also, longer spin times will result in a more compact pellet that lower the risk of clogging the column.



Carefully transfer supernatant to the spin column and centrifuge for © 00:01:00 . Discard



flow-through.







Discarding the flow-through is optional. The collection tube is designed to hold 800  $\mu$ l of flow-through fluid and still allow the tip of the column to be safely above the top of the liquid. Empty the tube whenever necessary to ensure the column tip and flow-through do not make contact.







Add ⊒400 µL Plasmid Wash Buffer 2 and centrifuge for ⊙00:01:00.

9 Transfer column to a clean 1.5 ml microfuge tube.

Use care to ensure that the tip of the column does not come into contact with the flow-through. If there is any doubt, re-spin the column for 00:01:00.

### 10





Add  $\geq$   $\square 30~\mu L$  DNA Elution Buffer to the center of the matrix. Wait for  $\bigcirc 00:01:00$ , then spin at 316000 x g for  $\bigcirc 00:01:00$  to elute DNA.

Nuclease-free water (p+7-p+8.5) can also be used to elute the DNA. Delivery of the Monarch DNA Elution Buffer should be made directly to the center of the column to ensure the matrix is completely covered for maximal efficiency of elution. Additionally, yield may slightly increase if a larger volume of DNA Elution Buffer is used, but the DNA will be less concentrated as a result of dilution. For larger plasmids ( $\geq$  10 kb), heating the DNA Elution Buffer to § 50 °C prior to eluting and extending the incubation time after buffer addition to 5 minutes can improve yield.