

## High Pure Plasmid Purification Resin

Product Number: PLR16

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### Shipping and Storage

Sealed storage at 15~25°C for 3 years.

### Product content

Index	Parameter
Carrying capacity	500µg plasmid/g filler
Particle size	50µm
PH tolerance range	6~9
Form of Preservation	White solid powder

### Description

Plasmid purification packing is based on anion exchange column technology, utilizing gravity flow to efficiently purify plasmid DNA. The yield can reach 10mg (Giga), 2.5mg (Mega), 500µg (Maxi), or 100µg (Mini) of high copy plasmid DNA. And ensure that the purified plasmid DNA has low endotoxin levels. The entire purification process avoids toxic substances such as phenol, chloroform, ethidium bromide, and CsCl, reducing the harm to users and the environment. The plasmid DNA purified by plasmid purification packing and reagent kit is suitable for transfection, sequencing, labeling, cloning, and any other experimental operations.

### Protocol

#### 1. Preparation of buffer solution

- 1.1. P1, P2, P3, ER buffer solution, equilibration solution, washing solution, and eluent need to be customized;
- 1.2. All water and buffer solutions are prepared using endotoxin free water, and it is recommended to filter them through a 0.22µm membrane before use;
- 1.3. Consumables without heat source;

#### 2. Filling of fillers

- 2.1. Add an appropriate amount of plasmid purification packing into the empty column with a lower sieve plate, compact it, and press it into the upper sieve plate;

Note: Add approximately 1.4g of powder filler to a 15mL empty column.

#### 3. sample preparation

##### 3.1. Bacterial preparation

- 2.5.1. Take out the bacterial strain or select monoclonal bacterial spots from the -80°C refrigerator, inoculate them into a 4 mL culture medium test tube, shake at 37°C for 8 hours at a speed of 300 rpm, and after 8 hours, transfer 4 mL of culture medium into 100 mL of rich culture medium. Shake overnight at 37°C at a speed of 300 rpm.
- 2.5.2. The bacterial solution cultured overnight was centrifuged at 4°C and 5000rpm for 15 minutes, the culture medium was discarded, and the precipitate was retained.

##### 3.2. Bacterial lysis

- 2.5.3. Add 10mL of P1 Buffer to the bacterial precipitate and suspend the bacterial cells thoroughly. Note: Please add RNase A to P1 Buffer before use, with a final concentration of 100µg/mL.
- 2.5.4. Add 10mL of P2 Buffer to the suspended bacterial solution, slowly mix evenly (without vigorous shaking), and perform lysis for no more than 5 minutes (to prevent genome contamination).
- 2.5.5. Add 10mL of P3 Buffer to the cracking system, mix slowly and evenly (without vigorous shaking), and a sheet-like precipitate appears. 7000rpm, Centrifuge for 10 minutes, then filter the supernatant using filter paper. Note: If endotoxin removal is required, please add one tenth of the filtered supernatant volume of ER Buffer to the

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supernatant, usually 3mL, and incubate at 4°C for 30 minutes.

**4. Sample purification**

- 4.1. Place the pre installed column on a plastic rack, add 2 \* 10mL of balancing solution, and wait for the liquid to flow out by gravity to complete the packing balance. Place the filler in the same buffer system as the target protein to provide protection.
- 4.2. Add the prepared bacterial solution and wait for it to flow out by gravity.
- 4.3. Add 2 \* 10mL washing buffer, add in batches according to the column capacity, and wait for gravity flow to complete.
- 4.4. Add 10mL elution buffer and collect the effluent using a 50mL centrifuge tube.

**5. Isopropanol precipitation**

Add 7mL of isopropanol to the centrifuge tube containing the eluent, mix it upside down, and centrifuge at 4°C and 11000RPM for 15 minutes to precipitate the plasmid. Slowly pour out the supernatant.

Note: Pre storing isopropanol at -20°C is more conducive to precipitation formation and improves recovery rate.

**6. Ethanol cleaning**

Add 10mL of 70% ethanol to the precipitate at 4°C and 11000RPM for 15 minutes. Slowly pour out the supernatant and let it stand at room temperature until ethanol evaporates.

**7. Plasmid preservation**

Dissolve the precipitate in an appropriate amount of endotoxin free water (or TE) to obtain the target plasmid solution. Using a spectrophotometer to measure plasmid concentration, the ratio of A260/A280 should be between 1.8 and 2.0. Use agarose electrophoresis to detect plasmid purity.

**Product ordering information**

Product name	Item number	Specifications
High Pure Plasmid Purification Resin	PLR16-50	50g
	PLR16-100	100g
	PLR16-500	500g
	PLR16-1kg	1kg