PrimePrep™ PCR Purification Kit

[지금 받으신 제품은 Sample size로써, 10회 prep.을 하실 수 있습니다. Buffer PW에 첨가할 Ethanol의 양은 bottle label에 표기되어있습니다.]

Introduction

PrimePrepTM PCR Purification Kit offer simple, rapid and cost-effective method for purification from PCR/enzyme reaction mixtures.

The purified DNA can be directly used in ligation, sequencing and other downstream application.

Kit Components

Cat. No. Reagents	K-7000 (50 prep.)
Spin column	50 ea
Buffer PCR-B	30 ml
Buffer PW	10 ml
Buffer PE	10 ml

Tel. 1599-4003, 0502-523-3484 Fax. 0502-523-3485

Before you begin

- Add ethanol to Buffer PW before use.
- → bottle의 label에 첨가량이 표시되어 있습니다.

Experimental Protocol

1. Add 5 volumes of Buffer PCR-B to 1 volume of the sample and mix well by vortexing.

If the PCR reaction product is 50 ul, add 250 ul of Buffer PCR-B.

- 2. Centrifuge the tube briefly at room temperature.
- 3. Transfer the mixture to a Spin column.
- 4. Centrifuge for 30 sec ~ 1 min at 13,000 rpm. Discard the flow-through and re-inserting the spin column to the collection tube.
- 5, Add 700 ul Buffer PW and centrifuge for 30 sec, at 13,000 rpm. Discard the flow-through and re-inserting the spin column to the collection tube.
- 6. Centrifuge for an additional 1 ~ 2 min at 13,000 rpm to remove residual wash buffer.

Residual ethanol of washing buffer may inhibitor subsequent enzymatic reaction.

7. Transfer the spin column to new 1.5 ml microcentrifuge tube.

The 1.5 ml microcentrifuge tube is not provided.

8 Add 50 ul of Buffer PE or deionized distilled water to the center of the membrane in the column, let stand for 1 min and centrifuge for 1 min at 13,000 rpm.

For larger fragment(>5kb), use pre-warmed (70°C) Buffer PE for best efficiency.

