

Quick Protocol

PCR-Pure Kit

PCR Purification based on silica technology

MiniPrep



Kit Content

Component	100 preps	
RPB	12ml	
GWB	10ml	
RSB	5ml	
HiPure DR column	50	
Collection tube	100	

Buffer Preparation

Before the first use, add appropriate amount of ethanol (96-100%) to each washing buffer tube and RPB tube, then mix thoroughly to prepare buffers, refer to Table 1. Do not forget to tick the check box on the bottle label to indicate that ethanol has been added. Before each use mix reconstituted buffer by shaking. Storing at room temperature.

Table 1: buffer preparation

Buffer Name	Concentrated Volume	Amount of Ethanol	Final Volume
GWB	10ml	40ml	50ml
RPB	12ml	28ml	40ml



Procedure of silica-based DNA isolation in quick look





Protocol: Purification of PCR product (based on silica technology)

Sample type: PCR products

Some tips to know:

- All centrifugation steps are carried out at room temperature (15–25°C).
- Not forget to add the appropriate amount of molecular biology grade ethanol to RPB as indicated on the bottle, before using for the first time.
- Not forget to add the appropriate amount of molecular biology grade ethanol (%96–100) to GWB as indicated on the bottle, before using for the first time (refer to buffer preparation section).
- RPB should be pink, if it turns yellow or orange, contact the technical support group.
- If RPB forms precipitate during storage, please warm it to 56°C until the precipitate has fully dissolved.
- RPB should be prepared. Not forget to add the appropriate amount of molecular biology grade ethanol (%96–100) to RPB as indicated on the bottle, before using for the first time (refer to buffer preparation section).

Process

- Add 350 μ l RPB to clean 1.5ml tube and 50-150 μ l PCR Product then invert 10 times and 3s pulse vortex.
- Transfer the solution to a Micro Column placed in a 2ml collection tube (supplied in the kit box). Centrifuge at 13000 rpm for 15s at room temperature.
- Discard the flow-through. Add 750µl GWB to the spin column. Centrifuge for 15s at 13000 rpm at room temperature. Discard the flow-through, and then centrifuge the Micro Column for an additional 1min at 14000 rpm.

Note: Discarding the flow-through before the second centrifuge is necessary to remove ethanol.

• Place the Micro Column in a new 1.5ml microcentrifuge tube. Add 30µl RSB directly to center of the spin column. Incubate at room temperature (15-25℃) for 3 min. Centrifuge for 2 min at 14000 rpm to elute the nucleic acid.