

Quick Protocol

DNJia Plus Blood & Cell Pro Kit

DNA isolation based on silica technology

MiniPrep

For DNA Isolation from

Blood

Buffy coat

Cultured cell



Kit Content

Component	50 preps	100 preps
GLB	10ml	20ml
BWB1 (concentrate)	17ml	35ml
BWB2 (concentrate)	24ml	48ml
RRB	10ml	20ml
RJ-Protease	2 x 1ml	4 x 1ml
HiPure DR Column	50	100
Collection Tube	100	200

Recommended Starting Material

The size of recommended starting material to use with determined lysis volume are listed here.

Table 1: Recommended starting material and Lysis Buffer amount

Sample	Size of Starting Material	Lysis Buffer Amount
Blood*	Up to 200µl	200μΙ
Buffy coat	Up to 200µl	200μΙ
Cultured cell	≤5 x 10 ⁶ cells	200μΙ

^{*}White Blood Cell counts are extremely variable from a person to person and therefore, for optimum results its recommended to determine cell counts for choosing the best protocol for purification. For healthy individuals, white blood cell counts normally from 4-10 million per 1 ml blood; however, depending on disease state, cell counts may be higher or lower than the normal range.

Before Start

- If GLB forms precipitate, please warm it to 56°C until the precipitate has fully dissolved. This is due to storage condition and will not influence the efficiency of buffer.
- Not forget to add the appropriate amount of ethanol (96–100%) to BWB1 and BWB2 as indicated on the bottle, before using for the first time, refer to washing buffer preparation.



Washing Buffer Preparation

Before the first use, add appropriate amount of ethanol (96-100%) to each washing buffer tube, then mix thoroughly to prepare washing buffer, refer to Table 2. 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 2: Washing buffer preparation

Buffer Name	Concentrated Volume	Amount of Ethanol	Final Volume
BWB1	16ml	24ml	40ml
BWB2	16ml	24ml	40ml



Procedure of silica-based DNA isolation in quick look





Protocols

Protocol: Isolation of Genomic DNA (Animal blood and Buffy coat) Sample type:

- Animal blood
- Buffy coat

Some tips to know

- All centrifugation steps are carried out at room temperature (15–25°C).
- If RNase treatment is desired, Prime-RNase can be ordered separately from ROJETechnologies, Cat NO EB983013.
- Do not forget to add the appropriate amount of ethanol (96–100%) to BWB1 and BWB2 as indicated on the bottle, before using for the first time, refer to washing buffer preparation part.
- If GLB forms precipitate, please warm it to 56°C until the precipitate has fully dissolved.

Process

• Add 40µl RJ-Protease to a 1.5ml clean microcentrifuge tube. Add 200µl buffy coat (blood) to the tube. Then add 200µl GLB. Invert several times, Pulse vortex for 15s and incubate at 70°C for 10min.

Optional: This protocol is specialized for DNA isolation. For usual and routine applications RNase treatment isn't needed. If RNA-free genomic DNA is required, add $10 \, \mu l$ Prime-RNase A (Cat NO. EB983013), mix by vortexing, and incubate for 5 min at room temperature before going to step 2.

- Add 200 µl ethanol (96-100%) to the lysate, mix by pulse vortexing for 15 s, then centrifuge briefly.
- Gently, pipette the mixture to a HiPure DR column placed in a 2 ml collection tube (supplied in the kit box). Centrifuge at 13000 rpm for 1 min. Discard flow-through and place the HiPure DR column in new collection tube (provided in the kit box).

Note: If the lysate did not pass the column, centrifuge at full speed until it passes through the column.



- Add 600 µl BWB1 and centrifuge for 1 min at 13000 rpm, discard the flow-through and place the HiPure DR column in the previous collection tube.
- Add 500 µl BWB2 and centrifuge for 1 min at 13000 rpm, discard the flow-through and place the HiPure DR column in the previous collection tube.
- Add 500 µl BWB2 and centrifuge for 3 min at 14000 rpm. Discard both the flow-through and the collection tube. Place the HiPure DR column in a new clean 1.5 ml microcentrifuge tube (not provided).

Note: To avoid ethanol carry over, be careful that the column does not come into contact with the flow-through, if it happens discard the flow-through, place the column back in a collection tube and centrifuge for another 1 min at 14000 rpm.

• Pipette 100-200 µl **pre-warmed (warmed at 70 °C)** RRB directly onto HiPure DR column. Incubate at 56 °C for 5 min. Centrifuge it at 13000 rpm for 1 min.