| Step | Well | Name    | Waiting<br>Time<br>(min :<br>ss) | Mixing<br>Time<br>(min<br>: ss) | Magnet<br>Time<br>(min<br>: ss) | Adsorption | Speed | Volume<br>(µL) |
|------|------|---------|----------------------------------|---------------------------------|---------------------------------|------------|-------|----------------|
| 1    | 1    | Lysis   | 00:00                            | 20:00                           | 00:00                           | Normal     | F     | 700            |
| 2    | 6    | Beads   | 00:00                            | 00:15                           | 00:30                           | Normal     | М     | 200            |
| 3    | 1    | Binding | 00:00                            | 10:00                           | 03:00                           | Strong     | F     | 700            |
| 4    | 2    | Wash 1  | 00:00                            | 03:00                           | 01:00                           | Strong     | F     | 500            |
| 5    | 3    | Wash 2  | 00:00                            | 02:00                           | 01:00                           | Strong     | F     | 800            |
| 6    | 4    | Wash 3  | 00:00                            | 02:00                           | 01:00                           | Strong     | F     | 800            |
| 7    | 5    | Elution | 01:00                            | 05:00                           | 01:00                           | Strong     | F     | 80             |
| 8    | 6    | Discard | 00:00                            | 00:30                           | 00:00                           | Normal     | S     | 200            |

Lysis temperature : 65°C, lysis heating end step 2;

Elution temperature : 75°C, elution start heating step 7.

## (Analysis of Nucleic Acid)

Get some DNA, diluted in a advisable factor with elution buffer.

Survey the OD260, OD280 and OD320.

Concentration  $(ng/\mu L) = 50 \times OD260 \times dilution$  fact

1.7 ≤ OD260-320/ OD280-320 ≤ 2.1

Notice:  $0.1 \le OD260 \le 1.0$ , the result of ratio is much reliable.

# **Company Information**

Manufacturer: Hangzhou Bioer Technology Co.,Ltd

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# MagaBio plus Whole Blood Genomic

# **DNA Purification Kit**

**Kit Components** 

| Cat#            | BSC08S1E                        | BSC08S1B   | BSC08M1B   |  |
|-----------------|---------------------------------|--|--|--|
| Components      | 32 Tests                        | 50 Tests   | 100 Tests  |  |
| PK Solution     | 0.32 mL                         | 0.5 mL   | 1 mL   |  |
| Lysis Buffer    |                                 | 25 mL  | 50 mL  |  |
| WB1 Buffer      | 96 Well                         | 12 mL<br>(add 18 mL absolute<br>ethanol before use)    | 24 mL<br>(add 36 mL absolute<br>ethanol before use)    |  |
| Wash Buffer     | pre-packed<br>plate<br>2 Pieces | 15 mLx 2<br>(add 35 mL absolute<br>ethanol before use) | 15 mLx 4<br>(add 35 mL absolute<br>ethanol before use) |  |
| Elution Buffer  |                                 | 10 mL  | 20 mL  |  |
| MagaBio Reagent |                                 | 0.75 mL  | 1.5 mL   |  |
| Handbook        | 1                               | 1  | 1  |  |

# Storage and expiry date

The kit can be transported at room temperature. 1.

2. The kit should be stored at 2~8°C.

3. All reagents are valid for 12 months if stored properly.

## [Introduction]

The kit provides a very simple, fast and cost effective technique to isolate high quality DNA. Using one simple protocol, high yield of purified DNA can be isolated from whole blood. MagaBio sample processing is based on proprietary magnetizable particles--MagaBio Reagent. The pure DNA can be applied extensively in PCR, Real-time PCR, sequencing, Southern hybridization, mutant analysis, SNP and the others.

According to the special interaction, use MagaBio nucleic acid separation system with a general protocol---sample processing, MagaBio adsorption, washing and elution, and can go high-throughput.

#### [Principle and Advantage]

DNA in the sample is released by PK Solution and Lysis Buffer. Released DNA is bound 1

exclusively and specifically to the MagaBio Reagent. DNA bound to Magnetic particles are captured by a magnetic tool and contaminants are removed by Wash Buffer once or more. The DNA is then eluted from particles by Elution Buffer or molecular grade water.

#### [Apparatus and materials to be prepared by the user]

- 1. Magnetic Rack or Bioer NPA-32P purification instrument;
- 2. Water bath or Dry bath;
- 3. Vortex mixer;
- $\label{eq:solute} 4. \ \ Absolute \ alcohol \ (For \ BSC08S1B \ and \ BSC08M1B).$

# [Important Notes]

- > Add the ethanol (as the volume marked on bottle label) to the WB1 Buffer and mix them well.
- > Add the ethanol (as the volume marked on bottle label) to the Wash Buffer and mix them well.
- The following procedures are applicable to the Bioer NPA-32P purification instrument. If use other purification instrument, the operating procedures shall be adjusted according to the performance of different instruments.

# [Protocol]

#### The manual purification

Please add absolute ethanol to WB1 Buffer and Wash Buffer and mix thoroughly before the first use.

#### 1. Sample processing

- 1) Equilibrate all reagents and samples to room temperature.
- 2) Pipet 10µL of PK Solution into the bottom of a 1.5 mL microcentrifuge tube.
- 3) Add  $200\mu$ L of sample to the microcentrifuge tube from the above.
- Add 500μL of the Lysis Buffer to the sample from the above and mix by pulse-vortexing intensively for 15-20 seconds.

*Note:* Mix the Lysis Buffer thoroughly before use, make sure that no crystal in the Lysis Buffer.

- 5) Incubate at 56°C for 20 minutes. Mixing every 10 minutes.
- 6) Remove the tube from  $56^{\circ}$ C.

## 2. MagaBio adsorption

- 1) Add  $15\mu$ L of the well-mixed (particles should be suspended) MagaBio Reagent.
- Mix the tube gently and incubate for 10 minutes at room temperature while mixing. *Note:* Using an end-over-end rotator or manual mixing every 2-3 minutes.
- 3) Aggregate MagaBio particles bound with DNA by using a magnetic rack. Discard the supernate, remove the tube from the magnetic rack and wash particles as described below.

- Add 500µL of WB1 Buffer to the tube from the above. Mix well by inverting the tube several times to ensure the particles are completely dispersed. Sediment the particles on the magnetic rack and discard the supernate.
- Add 800µL of Wash Buffer to the tube from the above. Mix well by inverting the tube several times to ensure the particles are completely dispersed. Aggregate the particles on the magnetic rack and discard the supernate.
- 3) Remove the tube from the magnetic rack and repeat washing step 2) one more time follow the above step.
- 4) Open the cap, dry at room temperature for 5 minutes.

### 4. Elution

- Add 80µL of Elution Buffer and mix, incubate at 60°C for 10 minutes.
  *Note:* Vortex gently every 2-3 minutes.
- 2) Aggregate the particles on the magnetic rack and transfer the supernate contained the isolated DNA carefully into a clean tube. The product is ready for further analysis. If the isolated DNA sample is not going to be tested on the same day, freeze at -20°C.

## 【The automatic purification】

With automatic machine, the kit is highly suitable for various samples, which provide a convenient platform to achieve high-throughput and fast and effective purification.

## 1. Reagent preparation

## 1) For BSC08S1B and BSC08M1B

Add 500µL Lysis Buffer to the 2.2mL 96 Deep Well column 1 and 7; 500µL WB1 Buffer to column 2 and 8; 800µL Wash Buffer to column 3,4 and 9, 10; 80µL Elution Buffer to column 5and 11; 185µL Pure Water and 15µL MagaBio Reagent to column 6 and 12.

2) For BSC08S1E

Turn the 96-well plate upside down three times after placed at room temperature, then rip off plastic film, centrifuge in 96-well centrifuge for seconds (or swing by hand) to avoid adhered liquid. Rip off aluminum foil film of 96-well plate; make sure the direction of the plate (magnetic beads in column 6th&12th).

2. Add 200µL sample and 10µL PK Solution to the 96-Deep Well column 1 and 7.

3. Put 96-Deep Well plate into the instrument, then plugs in 8-strip Tip and start the program.

3. Washing

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