

GeneFlow™ Gel Extraction Kit Quick Protocol

For research use only

Catalogue Number

DFG004, DFG100, DFG300

Instruction Manual Download

When using this product for the first time, or if you are unfamiliar with the procedure, please scan the QR code and download the complete instruction manual.

Geneaid



Instruction Manual Download

Gel Extraction Protocol

1. Gel Dissociation

Excise the agarose gel slice containing relevant DNA fragments and remove any extra agarose to minimize the size of the gel slice. Transfer up to 300 mg of the gel slice to a 1.5 ml microcentrifuge tube. Add **500 µl of QG Buffer** and mix by vortex. Incubate at 55-60°C for 10-15 minutes to ensure the gel slice has been completely dissolved. During incubation, invert the tube every 2-3 minutes. If the color of the mixture has turned to purple, add 10 µl of 3M Sodium Acetate (pH5.0) and mix thoroughly. Cool the dissolved sample to room temperature.

2. DNA Binding

Place a **DFH Column** in a **2 ml Collection Tube**. Transfer **800 µl of the sample mixture** to the **DFH Column**. Centrifuge at 14-16,000 x g for 30 seconds. Discard the flow-through and place the **DFH Column** back in the **2 ml Collection Tube**. If the sample mixture is more than 800 µl, repeat the DNA Binding step.

3. Wash

Add **400 µl of W1 Buffer** into the **DFH Column**. Centrifuge at 14-16,000 x g for 30 seconds then discard the flow-through. Place the **DFH Column** back in the **2 ml Collection Tube**. Add **600 µl of Wash Buffer** (make sure ethanol was added) into the **DFH Column**. Let stand for 1 minute at room temperature. Centrifuge at 14-16,000 x g for 30 seconds then discard the flow-through. Place the **DFH Column** back in the **2 ml Collection Tube**. Centrifuge at 14-16,000 x g for 3 minutes to dry the column matrix.

4. DNA Elution

Transfer the dried **DFH Column** to a new 1.5 ml microcentrifuge tube. Add **20-50 µl of (60-70°C) pre-heated Elution Buffer** or TE into the center of the column matrix. Let stand for at least 2 minutes to ensure the **Elution Buffer** is completely absorbed. Centrifuge for 2 minutes at 14-16,000 x g to elute the purified DNA.

Gel Extraction For Sequencing Protocol

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Components

Component	DFG004	DFG100	DFG300
QG Buffer	3 ml	80 ml	240 ml
3M Sodium Acetate (pH5.0)	N/A	200 µl	200 µl
W1 Buffer	2 ml	45 ml	130 ml
Wash Buffer (Add Ethanol)	1 ml (4 ml)	25 ml (100 ml)	50 ml + 25 ml (200 ml) (100 ml)
Elution Buffer	1 ml	6 ml	30 ml
DFH Columns	4	100	300
2 ml Collection Tubes	4	100	300

GenepHlow™ Gel Extraction Kit Functional Test Data

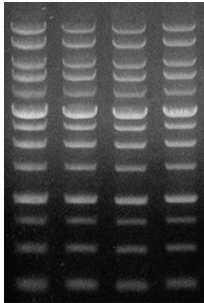


Figure 1. Gel slice DNA fragments ranging from 250 bp-10 kb were extracted using the GenepHlow™ Gel Extraction Kit (lane 1, 2, 3). The purified DNA from a 50 µl eluate was analyzed by electrophoresis on a 1% agarose gel.

M = Geneaid 1 Kb DNA Ladder (control, total DNA = 1100 ng)

Test	DNA Conc.	260/280	Yield	Recovery
1	18.2 ng/µl	1.84	910 ng	82%
2	18.7 ng/µl	1.82	935 ng	85%
3	19.2 ng/µl	1.82	960 ng	87.3%

M 1 2 3

Storage

Dry at room temperature (15-25°C)