

Gene to Protein Pvt. Ltd.

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G2P Cutting Edge Gel Extraction kit

Cat# ☐ PUR12-50, ☐ PUR12-50-S Pack size: ☐ 50 Preps, ☐ 5 Preps Storage: Room Temperature

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Kit Content

S.No	Components	PUR12-50	PUR12-50-S
1	Buffer GE	30mL	3mL
2	Buffer D (Wash Buffer)	10mL	2mL
3	Buffer E (Elution Buffer)	5mL	500 μl
4	Spin Columns and Collection Tubes	50	5

Introduction:

The G2P CuttingEdge Gel Extraction kit from Gene to Protein Pvt Ltd is a powerful tool for DNA fragment purification. Utilizing proprietary silica-based membrane technology, the kit delivers high yields of pure DNA fragments in a simple and rapid protocol that takes less time to complete. The kit is compatible with a wide range of downstream applications and is backed by a team of experienced technical support specialists.

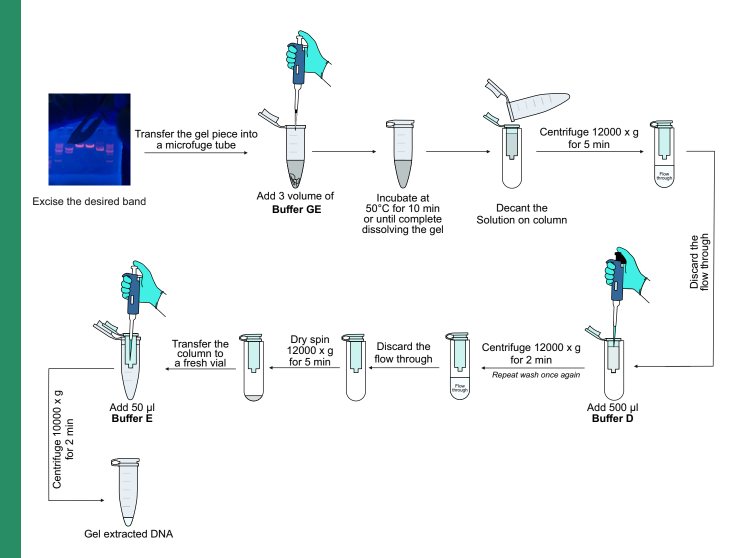
Procedure

Add 40ml ethanol to 10mL of buffer D before starting the protocol

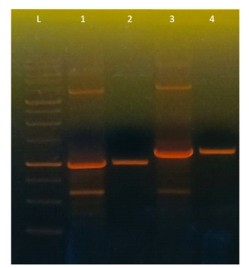
- 1. Excise the gel band containing the DNA fragment using a clean, sharp scalpel. Remove any excess agarose from the sides of the gel band.
- 2. Weigh the gel slice in a microfuge tube and add 3 volumes of Buffer GE to 1 volume of gel (approximately 300 μl of Buffer GE to 100 mg of gel).
- 3. Incubate the mixture at 50°C for 10 minutes or until the gel slice is completely dissolved. You may shake the tube or vortex it lightly, or pipette mix. If using higher percentage gels (>2%), increase the incubation time.
- 4. Load up to 600 μl of the mixture onto a spin column and spin at high speed (10,000xg). Load the leftover solution and repeat this step if necessary.
- 5. Discard the flow-through.
- 6. Add 600 µl of Buffer D and spin for 3 minutes.
- 7. Discard the flow-through.
- 8. Spin again for 3 minutes to dry.
- 9. Place the column on a fresh microfuge tube (not included in the kit) and add approximately 30 µl of Buffer E. Water may be used in place of Buffer E for elution. When using water, ensure that the pH of the water is 8.0. When using lower volumes of elution buffer, add the buffer dropwise to the center of the membrane. Wait for 1 minute after adding the Buffer E.
- 10. Spin at high speed and collect the elution (gel-extracted product).

Note: It is highly recommended to use low melting agarose or use a low percentage gel (~0.8%) for better recovery.

Graphical Protocol



Result



- L. 1Kb G2P ladder (Catalogue No. L12)
- 1. Before gel extraction of amplified 1kb template
- 2. After gel extraction of amplified 1kb template
- 3. Before gel extraction of amplified 1.2kb template
- 4. After gel extraction of amplified 1.2kb template

