Gel Electrophoresis - TBE 0.5X
Forked from Gel Electrophoresis

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ABSTRACT
Separates molecules based on size.

MATERIALS

- Ethidium P212121
- 1 kb DNA Ladder - 1,000 gel lanes New England Biolabs Catalog #N3232L
- Gel Loading Dye, Purple (6X), no SDS - 4.0 ml New England Biolabs Catalog #B7025S
- TAE Buffer (Tris-acetate-EDTA) Contributed by users Catalog #B49
- Agarose Contributed by users Catalog #A5304
- 1 Kbp Plus DNA Ladder Invitrogen - Thermo Fisher Catalog #10787018
- DNA Gel Loading Dye (6X) Thermo Fisher Scientific Catalog #R0611

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Protocol status: Working
We use this protocol and it's working

Created: Aug 29, 2018
Last Modified: Nov 16, 2018
PROTOCOL MATERIALS

- **DNA Gel Loading Dye (6X)** Thermo Fisher
  - Catalog #R0611
  - Materials, Step 7

- **Ethidium Phosphate** Materials
  - Catalog #P212121

- **1 kb DNA Ladder - 1,000 gel lanes** New England Biolabs Catalog #N3232L
  - Materials

- **Gel Loading Dye, Purple (6X), no SDS - 4.0 ml** New England Biolabs Catalog #B7025S
  - Materials

- **TAE Buffer (Tris-acetate-EDTA)** Contributed by users Catalog #B49
  - Materials

- **Agarose** Contributed by users Catalog #A5304
  - In Materials, Materials, Step 2

- **1 Kb Plus DNA Ladder** Invitrogen - Thermo Fisher Catalog #10787018
  - Materials, Step 7

SAFETY WARNINGS

- Ethidium Bromide potentially acts as a mutagen or carcinogen.

BEFORE START INSTRUCTIONS

Have a DNA Sample ready, typically either from PCR or a recently performed Restriction Digest. Dilute down the 50X TAE Buffer to 1X.

**Prep Work**

1. Pour 150 mL of 0.5X TBE Buffer into a 250 ml Duran Bottle.
   - 150 mL 0.5X TBE buffer

2. Weigh out 1.5 g Agarose and add it to the Duran bottle.
   - 1.5 g agarose
   - Agarose Contributed by users Catalog #A5304
3 Place Duran bottle in a microwave on medium power for two minutes or until solution is clear and agarose is completely dissolved, occasionally stirring.

4 Remove the Duran bottle from the microwave with a glove/pot holder and let it cool under tap water until you can comfortably pick it up without protection.

5 Place gel tray on clamp.
Add well combs and use a level to ensure it is balanced.
Pour 40 ml (small gel) or 70 ml (big gel) of melted agarose into the gel tray and let it sit for 15 minutes, or until solid.

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### Loading the Gel

6 Remove the well comb/s carefully as to not tear the gel and remove the tray from the clamp, but ensure the gel remains in the tray.

7 Place the gel tray into the gel electrophoresis apparatus with the wells closer to the negative/black end.
Pour additional 0.5X TBE Buffer to fill each side of the apparatus and to create a thin layer of buffer covering the top of the gel.
Pipette 5 μL of the 1kb DNA Ladder with Loading Dye into a well.
Prepare your DNA samples by adding an adequate amount of Gel Loading Dye (6X) and load them into the wells.

- **1 Kb Plus DNA Ladder Invitrogen - Thermo Fisher** Catalog #10787018
- **DNA Gel Loading Dye (6X) Thermo Fisher Scientific** Catalog #R0611

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### Running the Gel

8 Place lid on apparatus and plug cables into the power supply. Set the power supply to stay at a constant voltage of 90 V (small gel) - 120 V (big gel).
Let the gel run until the bromophenol blue reaches the end of the gel.
**Staining**

9. Remove the gel from the gel tray after draining excess of TBE Buffer. Place gel into a tray with 1% Ethidium Bromide ensuring the solution does not come into contact with your skin, then place the tray over a shaker with gentle agitation. Let the gel stain for 15 minutes.

**Destaining**

10. Carefully remove the gel from the Ethidium Bromide solution and place it in a tray with distilled water, then place the tray over a shaker with gentle agitation. Let the gel destain for 10 min.

**UV exposure**

11. Place the gel on top of a UV light source with a camera. Turn on the UV light and capture a picture of the gel.