

Restriction Enzyme Digest & Gel Electrophoresis Field Trip Background MS

Background Information

Restriction enzymes are proteins that cut double-stranded DNA at specific recognition sites. Think of them as a pair of molecular scissors making cuts along the genetic code. Restriction enzymes are isolated from bacteria. Bacteria use them as protection against the invasion of foreign DNA. They were proposed in the 1960's as an explanation to the question, "Why can bacteriophages (viruses that infect bacteria) infect some strains of bacteria but not others?" In 1968, similar enzymes were purified that were able to cut DNA. These early enzymes cut the DNA at random positions. In 1970, a more useful enzyme was purified and characterized. This enzyme, called *HindII*, cut DNA at a specific recognition site every time. *HindII* was used to create the first **restriction map**. The ability to cut DNA at specific sequences became the first step toward molecular cloning.



Think of a restriction enzyme cutting DNA like a pair of scissors

Restriction of DNA Using Restriction Enzymes

In the **Restriction Enzyme Digest & Gel Electrophoresis** field trip, students cut lambda bacteriophage DNA with 3 different restriction enzymes. Then, they will use gel electrophoresis to visualize the DNA. Molecular biology laboratory skills, equipment, and laboratory safety will be discussed and used in this lab.

Lambda bacteriophage DNA is used in this experiment. The DNA is linear and 48,502 base pairs long. It is useful because it has recognition sites for many different restriction enzymes. It is commonly used for molecular weight size markers in gel analysis of DNA.

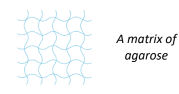
This process of cutting DNA is known as a **restriction digest**. Different restriction enzymes (and there are hundreds) recognize and cut different DNA sequences. After a restriction digest, the sizes of the resultant DNA fragments correspond to the distances (in base pairs) between restriction sites. The reaction is incubated at the enzyme's optimum temperature to digest the DNA. This usually takes about 45 minutes. During this time students will learn about **REs**, **restriction digest** and **gel electrophoresis**.



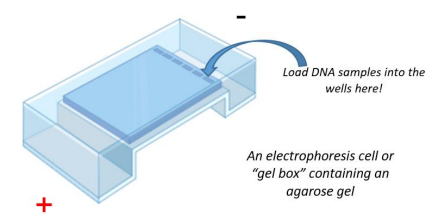
Agarose Gel Electrophoresis of DNA

Electrophoresis is a laboratory technique that is used to separate charged molecules from one another based on size and charge. DNA is negatively charged and will move under the force of an electric current. It will move through a gel made from a sugar called **agarose**. DNA molecules will separate by size. The small pieces will move more rapidly through the gel than the large ones.

Agarose is a powder that is insoluble in water (or buffer) at room temperature but melts in boiling water (or buffer). After it's heated, agarose undergoes **polymerization** as it cools, where the liquid agarose sugar molecules solidify into a gel (matrix of agarose). Higher concentrations of agarose produce firmer gels. 0.8% agarose gels in a sodium borate running buffer are used for this field trip. Molten agarose is cooled to 55°C prior to pouring into casting trays around a six-well comb. Then, it is left to solidify (instructors prepare the agarose gel). The comb is removed from the solidified gel to form small pockets called wells.



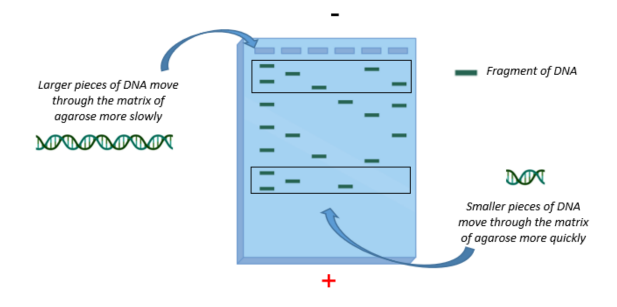
To perform **gel electrophoresis**, an agarose gel is placed into an electrophoresis cell that contains both positive and negative electrodes. A buffered solution (sodium borate) is added to act as an electrically conductive medium. Restriction enzyme-digested DNA samples are mixed with a loading dye containing tracking dyes, a thickening agent, and a chemical that serves as a DNA stain. The prepared samples are then loaded into the wells of the agarose gel. When the electrophoresis cell is connected to a power supply, DNA fragments move through the agarose gel.





Visualization of DNA in a Gel

After running the gel, DNA is visualized by exposing the gel to a medium-wavelength ultraviolet (UV) light source^{*}. The DNA stain component of the loading dye binds itself to the DNA molecule. When the gel is exposed to UV light, the DNA fragments will fluoresce. A photograph of the gel is taken so that it is easier to study the banding pattern. *Students should ALWAYS wear safety glasses while in the lab; ESPECIALLY when using a UV light source.



If you have any questions or would like more information before you bring your students to the BTCI for this field trip, please contact us. Alternatively, bring your questions along and we can discuss them during the lab. We look forward to seeing you and your group on your scheduled field trip day. Thank you for your interest in the BTC Institute's Biotechnology Field Trips program!

Images for this background obtained from Biorender.com



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