





Whose DNA Was Left Behind?

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All experiment components are intended for educational research only. They are not to be used for diagnostic or drug purposes, nor administered to or consumed by humans or animals.

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Experiment Components

Storage:

Store this experiment at room temperature.

This experiment is designed for 10 groups.

Contents

- A Crime Scene Simulated DNA Sample 1
- B Crime Scene Simulated DNA Sample 2
- C Suspect One Simulated DNA Sample 1
- D Suspect One Simulated DNA Sample 2
- E Suspect Two Simulated DNA Sample 1
- F Suspect Two Simulated DNA Sample 2

Transfer pipets
UltraSpec-Agarose™
50x Electrophoresis Buffer
Practice Gel Loading Solution

None of the experiment components have been prepared from human sources. Simulated DNA samples are non-toxic, water-based dyes.

Experiment Requirements

Experiment Requirements

- Electrophoresis Apparatus, M-12 or equivalent
- D.C. Power Supply
- Heat Source
- 500 ml Beaker or Flask
- Hot Gloves
- Distilled Water (used to make buffer solutions)
- Balance
- Automatic Micropipet and tips (optional)

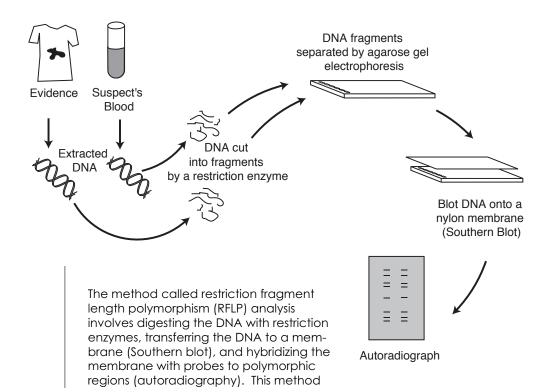


Whose DNA Was Left Behind?

Background and Introduction

DNA fingerprinting allows for the identification of the source of a DNA sample, which is very important in many forensic cases. DNA fingerprinting can provide positive identification with great accuracy by matching DNA obtained from a crime scene to individual suspects.

Several steps are involved in DNA fingerprinting. First, a suitable sample must be obtained. Forensic scientists use great care obtaining evidence from crime scenes so that the DNA will not be damaged. DNA is then isolated from the evidence, such as blood or hair samples. Once the DNA isolated, it is either digested with special enzymes called restriction endonucleases, or submitted to the Polymerase Chain Reaction (PCR).



requires relatively large amounts of DNA and takes several weeks. It is, how-

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ever, statistically very accurate.

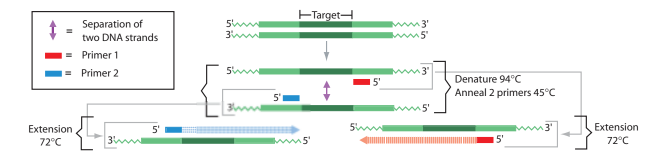
Whose DNA Was Left Behind?



Background and Introduction

More recently, the Polymerase Chain Reaction (PCR) has been used in forensics to analyze DNA. This technique requires much less (500-fold) DNA than RFLP analysis and is much less time-consuming. PCR amplification uses an enzyme known as Taq polymerase, which was originally purified from a bacterium that inhabits hot springs. It is stable at very high (near boiling) temperatures. The PCR reaction mixture also includes two (15-30 nucleotide) synthetic oligonucleotides, known as "primers". These items are mixed with the extracted DNA, known as the "template".

The region of DNA to be amplified is known as the "target". In the first step of the PCR reaction, the template's complimentary DNA strands are separated (denatured) from each other at 94°C, while the Taq polymerase remains stable. In the second step, known as annealing, the sample is cooled to an intermediate temperature, usually 40°-65°C, to allow hybridization of the two primers, one to each of the two strands of the template DNA. In the third step, known as extension, the temperature is raised to 72°C and the Taq polymerase adds nucleotides to the primers to complete the synthesis of the new complementary strands.



These three steps - denaturation, annealing, and extension - constitute one PCR "cycle". This process is typically repeated for 20-40 cycles, amplifying the target sequence exponentially. PCR is performed in a thermal cycler, an instrument that is programmed to rapidly heat, cool and maintain samples at designated temperatures for varying amounts of time.

In this experiment, you will analyze DNAs (represented by colored dyes) using aspects of RFLP analysis. In this hypothetical case, the dyes represent DNAs obtained from a crime scene and two suspects which have been cut by restriction enzymes and the fragmentation patterns serve as the individual fingerprint. The DNA (dye) fragmentation patterns are simple enough to analyze directly in the agarose gel. The objective is to analyze and match the DNA fragmentation patterns after agarose gel electrophoresis and determine if Suspect 1 or Suspect 2 was at the crime scene.



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Experiment Overview

BEFORE YOU START THE EXPERIMENT

- 1. Read all instructions before starting the experiment.
- 2. Write a hypothesis that reflects the experiment and predict experimental outcomes.

EXPERIMENT CONTENT OBJECTIVE

- Students will learn how restriction enzymes cut DNA molecules at specific base sequences producing DNA fragments of varying lengths.
- Students will learn how agarose gel electrophoresis separates different sizes of DNA fragments.
- Students will learn how these fragments form unique patterns for each person's DNA, which is the basis for DNA fingerprinting analysis.

WORKING HYPOTHESIS

If a DNA sample collected at the crime scene is cut with two different restriction enzymes and compared with DNA samples obtained from two suspects' DNA cut with the same two restriction enzymes, then one should be able to identify the real killer by the DNA fingerprint method.

MATERIALS FOR THE EXPERIMENT

Each Lab Group should have the following materials:

Activity One

- Electrophoresis Buffer
- Practice gel loading sample
- Sample delivery instrument

Automatic micropipet and tips, or Transfer pipet and beaker of distilled water

Activity Two

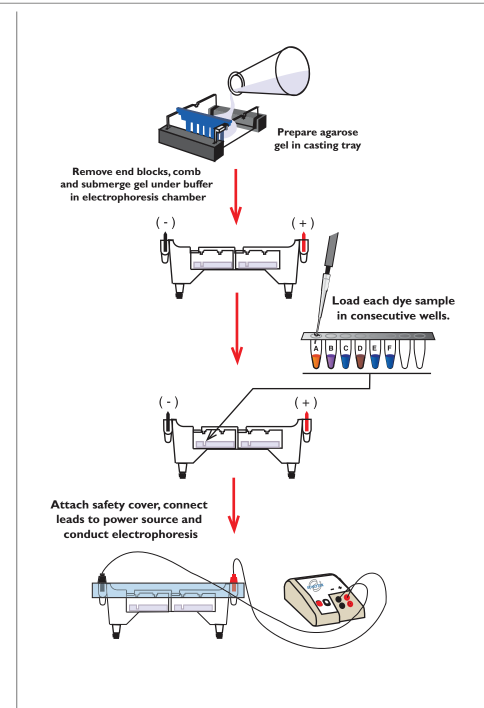
- Agarose gel
- Electrophoresis apparatus
- DC power source
- Dye Samples (A F) representing DNA
- Sample delivery instrument

Automatic micropipet and tips, or Transfer pipet and beaker of distilled water

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Experiment Overview





Whose DNA Was Left Behind?

Laboratory Safety

- 1. Gloves and goggles should be worn routinely as good laboratory practice.
- Exercise extreme caution when working with equipment that is used in conjunction with the heating and/or melting of reagents.
- DO NOT MOUTH PIPET REAGENTS USE PIPET PUMPS.



- 4. Exercise caution when using any electrical equipment in the laboratory.
 - Although electrical current from the power source is automatically disrupted when the cover is removed from the apparatus, first turn off the power, then unplug the power source before disconnecting the leads and removing the cover.
 - Turn off power and unplug the equipment when not in use.
- 5. EDVOTEK injection-molded electrophoresis units do not have glued junctions that can develop potential leaks. However, in the unlikely event that a leak develops in any electrophoresis apparatus you are using, IMMEDIATELY SHUT OFF POWER. Do not use the apparatus.
- 6. Always wash hands thoroughly with soap and water after handling reagents or biological materials in the laboratory.



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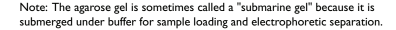
Activity One - Practice Gel Loading

Accurate sample delivery technique ensures the best possible gel results. Pipeting mistakes can cause the sample to become diluted with buffer, or cause damage to the wells with the pipet tip while loading the gel.

If you are unfamiliar with loading samples in agarose gels, it is recommended that you practice sample delivery techniques before conducting the actual experiment. EDVOTEK electrophoresis experiments contain a tube of practice gel loading solution for this purpose. Casting of a separate practice gel is highly recommended. One suggested activity is outlined below:

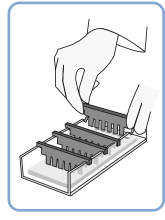
- 1. Cast a gel with the maximum number of wells possible.
- 2. After the gel solidifies, place it under buffer in an electrophoresis apparatus chamber.

Alternatively, your teacher may have cut the gel in sections between the rows of wells. Place a gel section with wells into a small, shallow tray and submerge it under buffer or water.



- 3. Practice delivering the practice gel loading solution to the sample wells. Take care not to damage or puncture the wells with the pipet tip.
 - For electrophoresis of dyes, load the sample well with 35-38 microliters of sample.
 - If using transfer pipets for sample delivery, load each sample well until it is full.
- 4. If you need more practice, remove the practice gel loading solution by squirting buffer into the wells with a transfer pipet.
- Replace the practice gel with a fresh gel for the actual experiment.

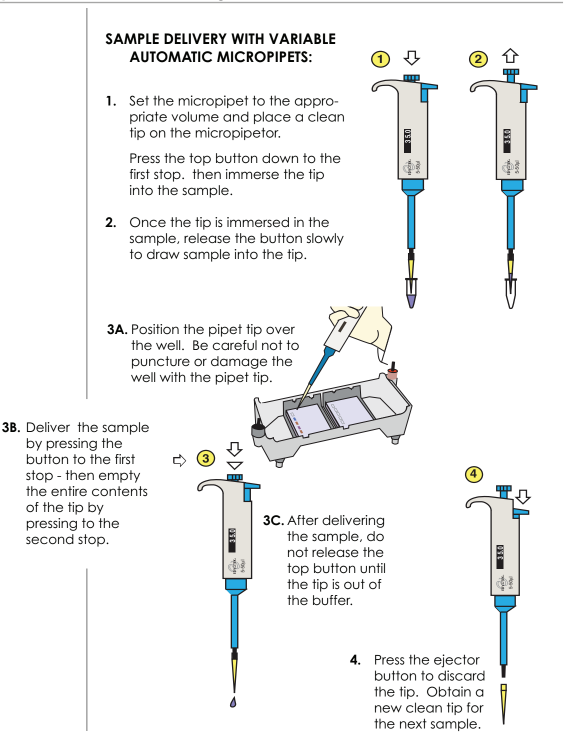
Note: If practicing gel loading in the electrophoresis chamber, the practice gel loading solution will become diluted in the buffer in the apparatus. It will not interfere with the experiment, so it is not necessary to prepare fresh buffer.





Whose DNA Was Left Behind?

Activity One - Practice Gel Loading



Whose DNA Was Left Behind?



Activity One - Practice Gel Loading

SAMPLE DELIVERY WITH PLASTIC TRANSFER PIPETS:

1. Gently squeeze the pipet stem to slowly draw the sample up into the pipet. The sample should remain in the lower portion of the pipet.

If the sample is overdrawn and becomes lodged in the bulb or on the walls, tap until the sample moves down into the lower stem of the pipet. Eject it back into the tube. Try step 1 again.

To control the delivery of small sample volumes with transfer pipets, gently squeeze the pipet stem, instead of the bulb.



- 2. While holding the pipet tip above the sample tube, slowly squeeze until the sample is nearly at the opening of the pipet tip.
- 3. Place the pipet tip in the electrophoresis buffer so it is directly above barely inside the sample well.
 - Avoid placing the pipet tip all the way inside the well this will minimize the chances of inadvertently piercing the bottom of the well.
- 4. MAINTAIN STEADY PRESSURE on the pipet stem to prevent buffer from being drawn in and diluting the sample.
- 5. Slowly squeeze to eject the sample. Stop squeezing when the well is completely full. Put any remaining sample in the pipet back into the sample tube.
- 6. Rinse the pipet with distilled water before obtaining the next sample for gel loading.



Whose DNA Was Left Behind?

Activity Two - Conducting Agarose Gel Electrophoresis

ELECTROPHORESIS SAMPLES

Samples in EDVOTEK Series 100 and S-series electrophoresis experiments are packaged in one of two different formats:

Pre-aliquoted Quickstrip™ connected tubes (new format)

To remove samples from the Quickstrip™ tubes, simply pierce the foil top with the micropipet tip and withdraw the sample.



Quickstrips patent pending



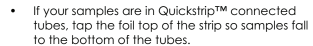
2. Individual 1.5 ml or 0.5 ml microtest tubes

Your instructor may have aliquoted these into a set of sample tubes for each lab group. Alternatively, you may be required to withdraw the appropriate amount from the experiment stock tubes.

LOADING THE SAMPLES

1. Check the Sample Volumes

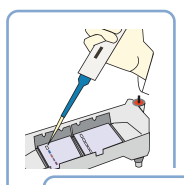
Sometimes a small amount of sample will cling to the walls of the tubes. Make sure the entire volume of sample is at the the bottom of the tubes before starting to load the gel.



• If your samples are in individual 1.5 ml or 0.5 ml microtest tubes, briefly centrifuge the sample tubes, or tap each tube on the tabletop to get all the sample to the bottom of the tube.



Load each of the dye samples in tubes A - F into the wells in consecutive order. The amount of sample that should be loaded is $35\text{-}38~\mu\text{l}$.



Lane Label Sample ı Α Crime scene DNA I 2 В Crime scene DNA 2 3 С Suspect I DNA I 4 D Suspect I DNA 2 5 Ε Suspect 2 DNA I 6 Suspect 2 DNA 2

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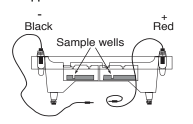
Activity Two - Conducting Agarose Gel Electrophoresis

RUNNING THE GEL

3. After the samples are loaded, carefully snap the cover down onto the electrode terminals.

Reminders:

During electrophoresis, the samples will migrate through the agarose gel towards the positive electrode. Before loading the samples, make sure the gel is properly oriented in the apparatus chamber.



Make sure that the negative and positive color-coded indicators on the cover and apparatus chamber are properly oriented.

- Insert the plug of the black wire into the black input of the power source (negative input). Insert the plug of the red wire into the red input of the power source (positive input).
- 5. Set the power source at the required voltage and conduct electrophoresis for the length of time determined by your instructor. General guidelines are presented in Table C.
- Check to see that current is flowing properly - you should see bubbles forming on the two platinum electrodes.

Table	C Time and Voltage
Electro	ophoresis of Dyes
Volts	Recommended Time
125	20 min
70	45 min
50	I hr 30 min

Staining is not required for Experiment # S-51, but results must be analyzed upon completion of the electrophoretic separation. Because dye molecules are extremely small they will diffuse out of the gel. Therefore, the gel cannot be saved.

- 7. After approximately 10 minutes, you will begin to see separation of the colored dyes.
- 8. After the electrophoresis is completed, turn off the power, unplug the power source, disconnect the leads and remove the cover.
- 9. Document the gel results.

A variety of documentation methods can be used, including drawing a picture of the gel, taking a photograph, or scanning an image of the gel on a flatbed scanner.



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Critical Thinking and Hypothesis Development



Record the following in your Laboratory Notebook or on a separate sheet of paper:

- 1. Based on the evidence obtained from analysis of the gel, which suspect committed the crime? Explain.
- 2. What is the variable in this experiment?
- 3. What would you change in the experiment if you had to do it over again?
- 4. Write a hypothesis that would reflect these changes.

Study Questions



Record the answers to the following Study Questions in your Laboratory Notebook or on a separate sheet of paper, as instructed by your teacher:

- 1. Why is it important to position the sample wells near the negative electrode?
- 2. What kind of evidence would you look for at a crime scene to obtain DNA?
- 3. Why is it important to wash the pipet between uses?
- 4. How will you be able to tell who committed the crime?
- 5. Who is the suspect that committed the crime?
- 6. What determines that each person has a unique pattern within their DNA?
- 7. Can you think of a case when two people will have identical DNA patterns?

Whose DNA Was Left Behind?



Notes to the Instructor

Class size, length of laboratory sessions, and availability of equipment are factors which must be considered in the planning and the implementation of this experiment with your students. These guidelines include Suggestions for Lesson Plan Content which can be adapted to fit your specific set of circumstances.

APPROXIMATE TIME REQUIREMENTS

- 1. UltraSpec-Agarose[™] gel preparation: Your schedule will determine when to prepare the gel(s) for an experiment. Whether you choose to prepare the gel(s) or have the students do it, allow approximately 30-40 minutes for this procedure. Generally, 20 minutes of this time is required for gel solidification.
- 2. The approximate time for electrophoresis will vary from 20 minutes to 1.5 hours.

ELECTROPHORESIS HINTS AND HELP

EDVOTEK Ready-to-Load Electrophoresis Experiments are easy to perform and are designed for maximum success in the classroom setting. However, even the most experienced students and teachers

> occasionally encounter experimental problems or difficulties.

The EDVOTEK web site provides a variety of resources which are continuously being updated and added. Several suggestions and reminders for conducting electrophoresis are available, as well as answers to frequently asked electrophoresis questions.

If you do not find the answers to your questions in this section or at the EDVOTEK web site, Technical Service is available from 9:00 am to 6:00 pm, Eastern time zone. Call for help from our knowledgeable technical staff at 1-800-EDVOTEK (1-800-338-6835).

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• The experiment number and title

Please have the following information:

- · Kit Lot number on box or tube
- · The literature version number (in lower right corner)
- · Approximate purchase date



Whose DNA Was Left Behind?

Instructor's Notes

SUGGESTIONS FOR LESSON PLAN CONTENT

This lesson plan outline, written by a teacher, can be used as a guideline to fit your specific classroom experience. The connections to the **National Content and Skills Standards** appear on pages that follow.

1. This experiment can be presented as a hypothetical burglary case. The detectives found some hair at the crime scene. You have isolated and prepared DNA from a strand of the hair. The detectives have four suspects, all of whom have submitted DNA samples. Using your DNA Fingerprinting skill, you must determine who the criminal is. When the DNA samples are run through an agarose gel, each sample will leave a specific pattern. The pattern matching the crime scene DNA will belong to the guilty person.

Alternatively, have students write a creative scenario that is based on the analysis of the gel or divide students into small groups and have each group write a short play based on the evidence obtained from the gel. Share scenarios with the class or give class time for a "short" play performance.

- 2. Using electrophoresis to separate DNA fragments is used in areas other than forensics. Have students conduct an "on-line" search to see how this procedure is used in the following cases:
 - Identifying individuals that are carriers of genetic diseases...
 - Identifying men and women killed in service.
 - Determining paternity
- 3. Discuss the importance of being meticulous in the collection and analysis of DNA that will be used as evidence in court cases.
- 4. From the vocabulary list of words below, have students write application sentences about each:

DNA Restriction Enzyme RFLP

Electrophoresis DNA Fingerprinting Polymerase Chain Reaction

- 5. List and discuss with students the essential parts of an experiment.
 - Writing a logical hypothesis
 - Making careful observations
 - Differentiating between an experiment and a control
 - Identifying variables
 - Predicting experimental outcomes
 - Recording results in a concise and accurate manner
 - Drawing valid interpretations of results
 - Formulating alternative explanations

Whose DNA Was Left Behind?

Instructor's Notes

ELECTROPHORESIS ANALYSIS OF SIMULATED DNA SAMPLES

DAY ONE

- 1. Have gels prepared in advance and hand out lab instructions.
- 2. The electrophoresis samples in this experiment are packaged in new prealiquoted QuickstripTM tubes. Several Quickstrips are connected in a block sealed with foil top cover. The sample block can easily be separated by scoring and tearing, or cut with scissors into strips of connected tubes. Each strip of connected tubes comprise a complete set of samples for each gel.
- Have students practice loading sample wells before doing the experiment.
 - If using micropipets, review proper use.
 - Thoroughly rinse wells before proceeding with experiment.
 - Remind students to keep track of what samples were loaded into which wells.
- 5. Clean up and answer student questions.

DAY TWO

- 1. Conduct the actual electrophoresis experiment.
 - Have students load the experiment samples.
 - Remind students to keep track of what samples were loaded into which wells.
- 2. At the end of the electrophoresis run, have students view and sketch results.
- 3. Have students answer a list of Study Questions and go over answers to the Study Questions.

OPTIONAL ACTIVITY FOR THE STUDENTS:

- Provide students with a list of topics for further research (library or newspaper research, written report, etc.).
- Set a date for students to report their research results.



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Connections to National Content Standards

- 1. Students will develop abilities necessary to do scientific inquiry.
 - Student questions will be answered through conduction of a scientific investigation.
- 2. Students will develop an understanding through inquiry.
 - Students will develop a logical hypothesis
 - Students will make careful observations.
 - Students will interpret results correctly.
 - Students will understand the difference between the experiment and the control.
 - Students will identify and control variable.
 - Students will predict experimental outcomes.
 - Students will formulate explanations from evidence.
 - Students will recognize and analyze alternative explanations.
- 3. Students will use equipment, materials, and techniques for experimentation and direct investigation of phenomena.
 - Students will understand the principles of agarose electrophoresis.
 - Students will understand how different sizes of DNA fragments are separated by agarose gel electrophoresis resulting in unique DNA Finger-prints for each individual.
- 4. Students will develop an understanding of the function of restriction enzymes.
 - Students will understand that restriction enzymes are endonucleases which catalyze the cleavage of bonds within both strands of DNA.
 - Students will understand that points of cleavage occur in or near very specific sequences of bases called recognition sites.
 - Students will understand that the number of bases in a recognition site
 and the distance between the recognition sites determines the size of
 the DNA fragment produced.
 - Students will be introduced to the concept of Polymerase Chain Reaction (PCR). Specific sequences or genes in DNA can selectively be purified by the enzyme DNA polymerase under specific conditions to yield sufficient DNA for analysis, as in the case of DNA fingerprinting.
- 5. Students will understand the principle behind DNA Fingerprinting
 - Students will understand that differences in restriction enzyme cleavage patterns among individuals will result in unique Restriction Fragment Length Polymorphisms (RFLPs) for each individual.
 - RFLPs are the basis for DNA fingerprinting which provides positive identification with great accuracy.

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Connections to National Skill Standards

In this experiment students will learn to load and run agarose gel electrophoresis. Analysis of th experiment will provide students the means to transform an abstract concept into a concrete explanation.

Students will be able to:

- 1. Use scientific equipment such as calibrated pipets for metric measurements and run electrophoresis units.
- 2. Accurately load and run an agarose gel.
- 3. Make careful observations and record results.
- 4. Perform a DNA fingerprint procedure.
- 5. Compare and evaluate DNA fingerprint patterns.
- 6. Discuss the limitations of DNA profiling

Instructor's Guidelines

Whose DNA Was Left Behind?

Preparations for the Experiment

Electrophoresis samples and reagents in EDVOTEK experiments are packaged in various formats. Samples in Series 100 and Sci-On electrophoresis experiments will be packaged in one of the following ways:

- Pre-aliquoted Quickstrip™ connected tubes (new format)
 OR
- 2) Individual 1.5 ml or 0.5 ml microtest tubes

IMPROVED FEATURES

В DO NOT BEND С С С С D D D D D D 5 5 CC Ę Ε Ε 5 Ε Ε Ε Ε **EDVOTEK®** G G G G G G G G G нΙ н Н нΙ нΙ Н Carefully cut between each set of tubes

FORMAT: PRE-ALIQUOTED QUICKSTRIP™ CONNECTED TUBES

If the Quickstrip™ samples are not already cut into individual strips:

 Use sharp scissors to separate each set of tubes A-H in the block of samples.

Note: In this experiment, tubes G and H are empty.

- Cut carefully through the foil between the rows of samples. Do not cut or puncture the foil covering the top of the sample tubes.
- 3. Each group will require one strip of samples.
- Remind students to tap the foil or tubes before gel loading to ensure that all of the sample is at the bottom of the tube.

Instructor's Guidelines

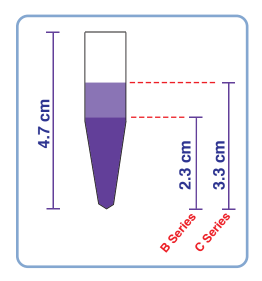
Whose DNA Was Left Behind?

Preparations for the Experiment



FORMAT: INDIVIDUAL 1.5 ML MICROTEST TUBES

It is recommended that samples packaged in 1.5 ml individual microtest tubes be aliquoted for each gel. Samples packaged in this format include bulk samples for certain EDVOTEK Series 100 or Sci-On electrophoresis experiments and are available in two standard quantities: the B-Series (480 μ l) and the C Series (960 μ l). Custom bulk quantities are also available by request.



Before aliquoting, check all sample volumes for possible evaporation. The samples will become more concentrated if evaporation has occurred.

If needed, tap or centrifuge the sample tubes. Then add distilled water to slightly above the following level:

- 2.3 cm level for the B-Series
- 3.3 cm level for the C-Series

Mix well by inverting and tapping the tubes several times.

After checking sample volumes and determining that the samples are at their proper total volumes:

 Aliquot the dye samples into appropriately labeled 0.5 ml or 1.5 ml microtest tubes:

 $38-40 \mu l$ of each sample

- 2. Students might have difficulty retrieving the entire aliquoted volume of sample because some of it may cling to the side walls of the tubes. Some suggestions are:
 - Remind students to make sure all of the sample is at the bottom of the tube before gel loading. They should centrifuge the samples tubes, or tap the tubes on the tabletop.
 - Instruct students to set their automatic micropipets to a volume that is 2 microliters less than the volume you have aliquoted.



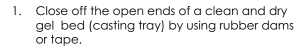
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Preparations for the Experiment

This experiment requires an 0.8% agarose gel. Agarose gels can be prepared before the laboratory period and stored under buffer. The simulated DNA samples (dyes) are ready-to-load for electrophoresis. Agarose gels can be prepared individually, or a batch preparation of agarose gel solution can be prepared to cast several gels at the same time. See page 26 for batch gel preparation instructions.

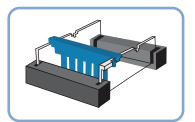


PREPARING THE GEL BED





- Place a rubber dam on each end of the bed. Make sure the rubber dam fits firmly in contact with the sides and bottom of the bed.
- B. Taping with labeling or masking tape:
 - With 3/4 inch wide tape, extend the tape over the sides and bottom edge of the bed.
 - Fold the extended edges of the tape back onto the sides and bottom. Press contact points firmly to form a good seal.
- Place a well-former template (comb) ii end of the gel bed. Make sure the cor bed.



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Preparations for the Experiment

CASTING AGAROSE GELS

- 3. Use a 250 ml flask to prepare the gel solution. Add the following components to the flask as specified for your experiment (refer to Table A).
 - Buffer concentrate
 - Distilled water
 - Agarose powder

Table A Individual 0.8% UltraSpec-Agarose™ Gel Electrophoresis of Dyes									
Size of EDVOTEK Casting Tray (cm)	Amt of Agarose - (g)	Concentrated Buffer (50x) (ml)	Distilled + Water = (ml)	Total Volume (ml)					
7 × 7	0.24	0.6	29.4	30					
7 × 15	0.48	1.2	58.8	60					

- 4. Swirl the mixture to disperse clumps of agarose powder.
- 5. With a marking pen, indicate the level of the solution volume on the outside of the flask.
- 6. Heat the mixture to dissolve the agarose powder. The final solution should appear clear (like water) without any undissolved particles.
 - A. Microwave method:
 - Cover the flask with plastic wrap to minimize evaporation.
 - Heat the mixture on High for 1 minute.
 - Swirl the mixture and heat on High in bursts of 25 seconds until all the agarose is completely dissolved.
 - B. Hot plate method:
 - Cover the flask with aluminum foil to prevent excess evaporation.
 - Heat the mixture to boiling over a burner with occasional swirling.
 Boil until all the agarose is completely dissolved.

At high altitudes, it is recommended to use a microwave oven to reach boiling temperatures.

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Preparations for the Experiment

7. Cool the agarose solution to 55°C with careful swirling to promote even dissipation of heat. If detectable evaporation has occurred, add distilled water to bring the solution up to the original volume as marked on the flask in step 5.

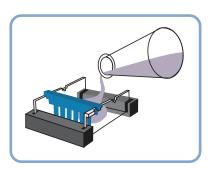
After the gel is cooled to 55°C:

If you are using rubber dams, go to step 9. If you are using tape, continue with step 8.

Cool the agarose to 55°C

DO NOT POUR BOILING HOT AGAROSE INTO THE GEL BED.

Hot agarose solution may irreversibly warp the bed.



- 8. Seal the interface of the gel bed and tape to prevent the agarose solution from leaking.
 - Use a transfer pipet to deposit a small amount of cooled agarose to both inside ends of the bed.
 - Wait approximately 1 minute for the agarose to solidify.
- 9. Pour the cooled agarose solution into the bed. Make sure the bed is on a level surface.
- 10. Allow the gel to completely solidify. It will become firm and cool to the touch after approximately 20 minutes.

Whose DNA Was Left Behind?

Preparations for the Experiment

PREPARING THE GEL FOR ELECTROPHORESIS

11. After the gel is completely solidified, carefully and slowly remove the rubber dams or tape from the gel bed.

Be especially careful not to damage or tear the gel wells when removing the rubber dams. A thin plastic knife, spatula or pipet tip can be inserted between the gel and the dams to break possible surface tension.

- 12. Remove the comb by slowly pulling straight up. Do this carefully and evenly to prevent tearing the sample wells.
- 13. Place the gel (on its bed) into the electrophoresis chamber, properly oriented, centered and level on the platform.



14. Fill the electrophoresis apparatus chamber with the required volume of diluted buffer for the specific unit you are using (see guidelines in Table B).

For DNA analysis, the same EDVOTEK 50x Electrophoresis Buffer is used for preparing both the agarose gel buffer and the chamber buffer. The formula for diluting EDVOTEK (50x) concentrated buffer is I volume of buffer concentrate to every 49 volumes of distilled or deionized water.

The electrophoresis (chamber) buffer recommended is Trisacetate-EDTA (20 mM tris, 6 mM sodium acetate, I mM disodium ethylenediamine tetraacetic acid) pH 7.8. Prepare the buffer as required for your electrophoresis apparatus.

Table B	Dilution of Electrophoresis (Chamber) Buffer								
EDVOTEK Model #	Concentrated Buffer (50x) + (ml)	Distilled Water (ml)	Total = Volume (ml)						
M6+	6	294	300						
MI2	8	392	400						
M36 (blue)	10	490	500						
M36 (clear)	20	980	1000						



Note: The UltraSpec-

Agarose[™] kit component is

often labeled with the amount it contains. In many cases, the

entire contents of the bottle is

3.0 grams. Please read the label

carefully. If the amount of

the bottle's plastic seal has been broken, weigh the agarose

to ensure you are using the

correct amount.

agarose is not specified or if

Sci-On® Biology

Whose DNA Was Left Behind?

Preparations for the Experiment

BATCH AGAROSE GEL PREPARATION

To save time, the agarose gel solution can be prepared in a batch for sharing by the class. Any unused prepared agarose can be saved and remelted for gel casting at a later time. For a batch (375 ml) preparation of 0.8% agarose gel:

- 1. Use a 500 ml flask to prepare the diluted gel buffer.
 - Add 7.5 ml of buffer concentrate
 - Add 367.5 ml of distilled water.
- Pour 3.0 grams of UltraSpec-Agarose[™] into the prepared buffer. Swirl to disperse clumps.
- 3. With a marking pen, indicate the level of solution volume on the outside of the flask.
- Heat the agarose solution as previously described for individual gel preparation. The heating time will require adjustment due to the larger total volume of gel buffer solution.
- 5. Cool the agarose solution to 55°C with swirling to promote even dissipation of heat. If evaporation has occurred, add distilled water to bring the solution up to the original volume (375 ml) as marked on the flask in step 3.



Table D Batch Preparation of 0.8% UltraSpec-Agarose™

Amt of Agarose (g)	+	Concentrated Buffer (50x) (ml)	+	Distilled Water (ml)	=	Total Volume (ml)
3.0		7.5		367.5		375

- Dispense the required volume of cooled agarose solution for casting the gels. The volume required is dependent upon the size of the gel bed (refer to Table A for individual gel casting guidelines).
- Allow the gel to completely solidify. It will become firm and cool to the touch after approximately 20 minutes. Then proceed with preparing the gel for electrophoresis.

Preparations for the Experiment

ELECTROPHORESIS TIME AND VOLTAGE

Your schedule will dictate the length of time samples will be separated by electrophoresis. In general, longer electrophoretic runs will increase the

separation between fragments of similar size. Because this experiment involves the electrophoresis of dyes, it can be easily monitored visually. Follow general guidelines as presented in Table C, but monitor the electrophoresis to make sure that the dyes do not migrate off the end of the gel.

Table	C Time and Voltage		
Electrophoresis of Dyes			
Volts	Recommended Time		
125	20 min		
70	45 min		
50	I hr 30 min		

If you don't find answers to your questions in this section, call our

Technical Service Department

EN O. TECH SERVICE **-800-EDVOTE**K (1-800-338-6835) Fri 9am Spm

Mon

Mon - Fri 9:00 am to 5:00 pm EST

24-hour FAX: (301) 340-0582

web: www.edvotek.com email: edvotek@aol.com

Please have the following information:

- · The kit number and title
- · Kit lot number on box or tube
- The literature version number (in lower right corner)
- Approximate purchase date



Visit our web site for information about EDVOTEK's complete line of experiments for biotechnology and biology education.



Whose DNA Was Left Behind?

Notes regarding electrophoresis

- 1. Do not move the apparatus immediately after the samples have been loaded.
 - Moving the apparatus will dislodge the samples from the wells into the buffer and will compromise results.
 - If it is necessary to move the apparatus during electrophoresis, you may safely do so after the tracking dye has migrated at least 1 cm from the wells into the gel.
- For optimal separation, do not use voltages higher than 125 volts for agarose gel electrophoresis. Higher voltages can overheat and melt the gel.
- Electrophoresis should be terminated when the dyes have moved 3 to 4 centimeters from the wells and before it moves off the gel.

AVOIDING COMMON PITFALLS

Potential pitfalls and/or problems can be avoided by following the suggestions and reminders listed below.

- To ensure that dyes are well resolved, make sure the gel formulation is correct (see Table A) and that electrophoresis is conducted for the optimal recommended amount of time.
- Correctly dilute the concentrated buffer for preparation of both the gel and electrophoresis (chamber) buffer. Remember that without buffer in the gel, there will be no sample mobility. Use only distilled water to prepare buffers. Do not use tap water.
- For optimal results, use fresh electrophoresis buffer prepared according to instructions.
- Before performing the actual experiment, practice sample delivery techniques to avoid diluting the sample with buffer during gel loading.
- To avoid loss of samples into the buffer, make sure the gel is properly oriented in the electrophoresis unit so the samples are not electrophoresed in the wrong direction off the gel.

Whose DNA Was Left Behind?



Experiment Results and Analysis

(+)

Idealized results are shown in the figure at left. Actual results will yield bands of varying intensity. The idealized schematic shows the relative positions of the bands, but are not depicted to scale.

Lane	Tube	
1	Α	Crime Scene DNA 1
2	В	Crime Scene DNA 2
3	С	Suspect 1 DNA 1
4	D	Suspect 1 DNA 2
5	Е	Suspect 2 DNA 1
6	F	Suspect 2 DNA 2

Instructor's Guidelines

Please refer to the kit insert for the Answers to Study Questions



Material Safety Data Sheet
May be used to comply with OSHA's Hazard Communication
Standard. 29 CFR 1910.1200 Standard must be consulted for

LO VOICE	Standard, 29		pecific requirements.	it be consulted it	л	
IDENTITY (As Used on Label and List)			Note: Blank spaces are not permitted. If any item is not applicable, or no information is available, the space must			
Agarose Section I		ı .	be marked to indicate th	at.		
Manufacturer's Name		Emer	gency Telephone Nur	mher		
		Elliel	gency relephone Nui	(301) 2	51-5990	
EDVOTEK, Inc.		Telepi	none Number for inform	ation		
Address (Number, Street, City, State,	Zip Code)			(301) 2	51-5990	
14676 Rothgeb Drive		Date I	Prepared 07/01/0	3		
Rockville, MD 20850		Signat	ture of Preparer (option	al)		
Section II - Hazardous Ingred	ients/Iden	tify Ir	formation			
Hazardous Components [Specific Chemical Identity; Common Name(s)]	OSHA	PEL		ther Limits commended	% (Optional)	
This product contains no hazardous r	naterials as de	fined b	v the OSHA Hazard	Communication	1	
Standard.			,			
CAS #9012-36-6						
Section III - Physical/Chemic	al Charact	eristi	cs			
Boiling Point For 1% solution	194° F	Spe	cific Gravity (H ₂ 0 = 1)		No data	
Vapor Pressure (mm Hg.)	No data	mening i emi			No data	
Vapor Density (AIR = 1)	No data	Evaporation Rate (Butyl Acetate = 1)			No data	
Solubility in Water Insoluble - cold						
Appearance and Odor White p	owder, no odo	or				
Section IV - Physical/Chemic	al Charact	eristi	cs N.D. = No da	ta		
Flash Point (Method Used) No data		Flam	imable Limits	LEL N.D.	UEL N.D.	
Extinguishing Media Water spray, dry chemical, carbon dioxide, halon or standard foam						
Special Fire Fighting Procedures Possible fire hazard when exposed to heat or flame						
Unusual Fire and Explosion Hazards	None					

Stability	Unstable		Conditions to Avoid	
	Stable	X	None	
Incompatibility No	data available			
Hazardous Decomposition	or Byproducts			
Hazardous	May Occur		Conditions to Avoid	
Polymerization	Will Not Occu	r X	None	
Section VI - Health	n Hazard Data			
Route(s) of Entry:	Inhalat	tion? Ye	es Skin? Yes	Ingestion? Yes
Health Hazards (Acute a	and Chronic) ion: No data avail:	-1-1-	Iti I	
Carcinogenicity:	NTP		Ingestion: Large amou IARC Monographs?	
Signs and Symptoms of	Exposure No da	ata availa	ble	
Medical Conditions Gen	erally Aggravated	by Expos	Sure No data available	
Emergency First Aid Pro	ocedures			
	Treat	symptom	atically and supportively	
Section VII - Preca				
Steps to be Taken in cas			•	
	Sweep up an	d place ii	n suitable container for dis	posal
Waste Disposal Method				
	Normal solid	l waste di	isposal	
Precautions to be Taken		toring		
	None			
Other Precautions				
	None			
Section VIII - Cont	rol Measures			
Respiratory Protection (Specify Type)	hemical	cartridge respirator with fu	ıll facepiece.
Ventilation	Local Exhaus	t	Spec	cial
	Mechanical (C	General)G	en. dilution ventilationOth	er
Protective Gloves Y	es		Eye Protection	Splash proof goggles
Other Protective Clothing	g or Equipment	Impervi	ous clothing to prevent ski	n contact
Work/Hygienic Practices	8	None		



Special Fire Fighting Procedures

Unusual Fire and Explosion Hazards

Material Safety Data Sheet

May be used to comply with OSHA's Hazard Communication
Standard. 29 CFR 1910.1200 Standard must be consulted for
specific requirements.

IDENTITY (As Used on Label and List) 50x Electrophoresis Buffer		Note: Blank spaces are not permitted. If any item is not applicable, or no information is available, the space mube marked to indicate that.			
Section I		•			
Manufacturer's Name	Emer	rgency Telephone Number (301) 251-5990			
EDVOTEK, Inc.		` ,			
Address (Number, Street, City, State, Zip Code)	Telepi	phone Number for information (301) 251-5990			
14676 Rothgeb Drive Rockville, MD 20850	Date Prepared 07/01/03				
Hockville, MD 20850	Signa	ature of Preparer (optional)			
Section II - Hazardous Ingredients/Iden	tify Ir	nformation			
Hazardous Components [Specific Chemical Identity; Common Name(s)] OSHA	PEL	Other Limits ACGIH TLV Recommended % (Options			
This product contains no hazardous materials : Communication Standard.	as defin	ned by the OSHA Hazard			
Section III - Physical/Chemical Charact	eristi	ics			

Section ii - nazardous ingredients/identity information								
Hazardous Components [Specific Chemical Identity; Common Name(s)]	Hazardous Components [Specific Other Limits Chemical Identity; Common Name(s)] OSHA PEL ACGIH TLV Recommended							
This product contains no hazardous materials as defined by the OSHA Hazard Communication Standard.								
Section III - Physical/Chemica	al Characte	ristics						
Boiling Point	No data	Specific Gravity (H20	No data					
Vapor Pressure (mm Hg.)	No data	Melting Point	No data					
Vapor Density (AIR = 1) No data Evaporation Rate (Butyl Acetate = 1)				No data				
Solubility in Water Appreciable, (§	greater than 10	%)						
Appearance and Odor Clear, liquid, slight vinegar odor								
Section IV - Physical/Chemical Characteristics N.D. = No data								
Flash Point (Method Used) No da	ata	Flammable Limits	LEL N.D.	UEL N.D.				
Extinguishing Media	se extinguishin	ng media appropriate fo	or surrounding fire					

None identified

Wear protective equipment and SCBA with full facepiece operated in positive pressure mode.

Section V - Reactive	ty Data						
Stability	Unstable		Condit	ions to Avoid	d		
	Stable	X		None			
Incompatibility	Strong oxidi:	zing age	ents				
Hazardous Decomposition o	r Byproducts Ca	arbon mo	onoxide,	Carbon dic	xide		
Hazardous	May Occur		Condi	tions to Avo	id		
Polymerization	Will Not Occur	X		None			
Section VI - Health	Hazard Data						
Route(s) of Entry:	Inhalatio	on? Ye	es	Skin?	Yes	I	ngestion?
Health Hazards (Acute an	d Chronic) Non-	e					
Carcinogenicity: None ide	entified NTP?		IAF	RC Monogr	aphs?	OSH	A Regulation?
Signs and Symptoms of E	xposure Irritati	on to up	per respi	ratory tract	, skin, ey	es	
Medical Conditions Gene	rally Aggravated b	y Expos	ure	None			
Emergency First Aid Prod	edures Ingestic	on: If co	onscious	give large	amounts	of water	
Eyes: Flush with water	Inhalation: !	Move to	fresh air	Skin: Was	sh with so	ap and wat	ter
Section VII - Precau	itions for Safe	e Hand	dling a	nd Use			
Steps to be Taken in case					ole protec	tive clothir	ng. Mop up spill
and rinse	with water, or colle	ect in ab	sorptive	material an	d dispose	of the abso	orptive material.
Waste Disposal Method	Dispose in accor environmental re			pplicable fe	ederal, sta	te, and loca	al
Precautions to be Taken i	n Handling and St	oring					
	Avoid eye and s	kin cont	act.				
Other Precautions							
	None						
Section VIII - Contr	ol Measures						
Respiratory Protection (S	pecify Type)						
Ventilation	Local Exhaust	Y	es		Special	Non	e
	Mechanical (Ge	eneral)	Yes		Other	None	e
Protective Gloves Ye	s			Eye Prote	ection	Safety go	ggles
Other Protective Clothing	or Equipment	None		-			
Work/Hygienic Practices		None					



Material Safety Data Sheet
May be used to comply with OSHA's Hazard Communication
Standard. 29 CFR 1910.1200 Standard must be consulted for specific requirements.

Signature of Preparer (optional)

IDENTITY (As Used on Label and List) Practice Gel Loading Solution		Note: Blank spaces are not permitted. If any item is not applicable, or no information is available, the space mus be marked to indicate that.
Section I		
Manufacturer's Name	Emer	gency Telephone Number (301) 251-5990
EDVOTEK, Inc.		(301) 251-5990

Address (Number, Street, City, State, Zip Code) 14676 Rothgeb Drive

Rockville, MD 20850

Telephone Number for information

(301) 251-5990 07/01/03

Section II - Hazardous Ingredients/Identify Information

Hazardous Components [Specific Chemical Identity; Common Name(s)] Other Limits
ACGIH TLV Recommended % (Optional) OSHA PEL This product contains no hazardous materials as defined by the OSHA Hazard Communication

Section III - Physical/Chemical Characteristics

Boiling Point	No data	Specific Gravity (H ₂ 0 = 1)	No data
Vapor Pressure (mm Hg.)	No data	Melting Point	No data
Vapor Density (AIR = 1)	No data	Evaporation Rate (Butyl Acetate = 1)	No data

Solubility in Water Soluble

Appearance and Odor Blue liquid, no odor

Section IV - Physical/Chemical Characteristics

Flash Point (Method Used) Flammable Limits No data No data No data Extinguishing Media

Dry chemical, carbon dioxide, water spray or foam Special Fire Fighting Procedures Use agents suitable for type of surrounding fire. Keep upwind, avoid

breathing hazardous sulfur oxides and bromides. Wear SCBA.

Unusual Fire and Explosion Hazards

Unknown

	Unstable	1 1	Conditions to Ave	лu		
	Stable	X	None			
Incompatibility	None					
Hazardous Decomposition	or Byproducts Sul	fur oxides,	and bromides			
Hazardous	May Occur	T	Conditions to Av	oid		
Polymerization	Will Not Occur	X	None			
Section VI - Healt	h Hazard Data					
Route(s) of Entry:	Inhalati	Yes		Yes	Ingestion?	Yes
Health Hazards (Acute		ute eye con er routes.			No data available fo	r
Carcinogenicity: No data ava	NTP?	1	IARC Mono	graphs?	OSHA Regulation	on?
Signs and Symptoms o	f Exposure May	cause skin	or eye irritation			
Medical Conditions Ger	nerally Aggravated b	y Exposure	None repo	rted		
Emergency First Aid Pr	110		atically and sup mounts of wate		inse contacted area	
Section VII - Prec)		
Steps to be Taken in ca	ise Material is Relea	sed for Spil	lled			
Wear eye and skin	protection and mop	spill area. I	Rinse with water	r.		
Waste Disposal Method			Rinse with water	r.		
Waste Disposal Method Observe all federal	d , state, and local regu n in Handling and St	ılations.	Rinse with wate	r.		
Waste Disposal Method Observe all federal	d , state, and local regu n in Handling and St	ılations.	Rinse with wate	r.		
Waste Disposal Method Observe all federal Precautions to be Take Avoid eye and skin Other Precautions None	, state, and local regu n in Handling and St contact.	ılations.	Rinse with water	r.		
Waste Disposal Method Observe all federal Precautions to be Take Avoid eye and skin Other Precautions None	d , state, and local regun in Handling and St contact.	ılations.	Rinse with water	r.		
Waste Disposal Method Observe all federal Precautions to be Take Avoid eye and skin Other Precautions None Section VIII - Con Respiratory Protection	d , state, and local regun in Handling and St contact.	ulations.	Yes	r. Special	None	
Waste Disposal Method Observe all federal Precautions to be Take Avoid eye and skin Other Precautions None Section VIII - Con Respiratory Protection	trol Measures (Specify Type)	ulations.			None None	
Waste Disposal Method Observe all federal Precautions to be Take Avoid eye and skin Other Precautions None Section VIII - Con Respiratory Protection Ventilation	trol Measures (Specify Type) Local Exhaust	ulations.	Yes	Special Other		ggles
Waste Disposal Method Observe all federal. Precautions to be Take Avoid eye and skin Other Precautions None Section VIII - Con Respiratory Protection Ventilation Protective Gloves	trol Measures (Specify Type) Local Exhaust Mechanical (Green	ulations.	Yes Yes Eye Pro	Special Other	None	ggles
Waste Disposal Method Observe all federal Precautions to be Take Avoid eye and skin Other Precautions None Section VIII - Con	trol Measures (Specify Type) Local Exhaust Mechanical (Gr	oring eneral)	Yes Yes Eye Pro	Special Other tection	None	ggles

Section V - Reactivity Data



Material Safety Data Sheet

May be used to comply with OSHA's Hazard Communication Standard. 29 CFR 1910.1200 Standard must be consulted for specific requirements.

IDENTITY (As Used on Label and List) B-1: Food Dye		Note: Blank spaces are not permitted. If any item is not applicable, or no information is available, the space must be marked to indicate that.
Section I		
Manufacturer's Name	Emer	gency Telephone Number (301) 251-5990
EDVOTEK, Inc.		. ,
Address (Number, Street, City, State, Zip Code)	Telepi	none Number for information (301) 251-5990
14676 Rothgeb Drive Rockville, MD 20850		O7/01/03
		Signature of Preparer (optional)

Section II - Hazardous Ingredients/Identify Information

Other Limits
ACGIH TLV Recommended % (Optional) Hazardous Components [Specific Chemical Identity; Common Name(s)] OSHA PEL This product contains no hazardous materials as defined by the OSHA Hazard Communication Standard.

Section III - Physical/Chemical Characteristics

Boiling Point	No data	Specific Gravity (H ₂ 0 = 1)	No data
Vapor Pressure (mm Hg.)	No data	Melting Point	N/A
Vapor Density (AIR = 1)	No data	Evaporation Rate (Butyl Acetate = 1)	No data
Solubility in Water			

Solubility in Water Soluble

Appearance and Odor Blue color, liquid, no odor

Section IV - Physical/Chemical Characteristics

Flash Point (Method Used)	Flammable Limits	LEL	UEL
No data		No data	No dat

Extinguishing Media N/A

Special Fire Fighting Procedures

Unusual Fire and Explosion Hazards

occion v modelivity	Data			
Stability	Unstable		Conditions to Avoid	
	Stable	Χ	Unknown	
Incompatibility		None		
Hazardous Decomposition or E	Syproducts Sul	fur oxid	les and bromides	
Hazardous	May Occur		Conditions to Avoid	
Polymerization	Will Not Occur	Χ	None	
Section VI - Health H	azard Data			
Route(s) of Entry:	Inhalatio	n? No	Skin? Yes	Ingestion? Yes
Health Hazards (Acute and	Chronic) A	cute e	ye contact: may cause	irritation
Carcinogenicity:	NTP?		IARC Monographs?	OSHA Regulation?
None	No	data	No data	No
Signs and Symptoms of Exp	oosure		May cause ski	n or eye irritation
Medical Conditions Genera	lly Aggravated by	/ Exposu	None reported	t
Emergency First Aid Proce		ntacted	d areas with copious am	ounts of water

Section VII - Precautions for Safe Handling and Use

Steps to be Taken in case Material is Released for Spilled

Wear eve and skin protection and mop/wipe spill area. Rinse with water.

Waste Disposal Method

Can be disposed in the trash or down the sink

Precautions to be Taken in Handling and Storing Avoid eye and skin contact

Other Precautions

None

Section VIII - Control Measures

Respiratory Protection (Specify Type) NIOSH/MSHA - approved respirator Ventilation Local Exhaust Special None

No Other None Mechanical (General) Eye Protection Protective Gloves Yes Splash prof goggles

Other Protective Clothing or Equipment None required

Work/Hygienic Practices Avoid eye and skin contact



Material Safety Data Sheet

EDVØTEK.		CFR 19		zard Communication must be consulted s.	
IDENTITY (As Used on Label and List)			Note: Blank space	s are not permitted. I	f any item is not
R-40 Food dye			be marked to indica	formation is available ate that.	, tne space must
Section I					
Manufacturer's Name		Emer	gency Telephone	Number (301)	251-5990
EDVOTEK, Inc.		Teleni	none Number for in	. ,	
Address (Number, Street, City, State,	Zip Code)	Тоюр	iono riambor ioi in		251-5990
14676 Rothgeb Drive Rockville, MD 20850			O7/01	,	
		Sigria	luie di Freparer (di	nioriai)	
Section II - Hazardous Ingred	lients/Iden	tify Ir	formation		
Hazardous Components [Specific Chemical Identity; Common Name(s)]	OSHA	PEL	ACGIH TLV	Other Limits Recommended	% (Optional)
This product contains no hazard Communication Standard.	lous materic	als as c	defined by the	OSHA Hazard	
Section III - Physical/Chemic	al Charact	eristi	cs		
Boiling Point	No data	Spe	cific Gravity (H ₂ 0	= 1)	No data
Vapor Pressure (mm Hg.)	No data	Mel	ting Point		N/A
Vapor Density (AIR = 1)	No data		poration Rate tyl Acetate = 1)		No data
Solubility in Water Soluble	le				
Appearance and Odor Red Co	olor, liquid, r	no odc	or		
Section IV - Physical/Chemic	al Charact	eristi	cs		
Flash Point (Method Used) No dat	ta	Flam	nmable Limits	LEL No data	UEL No data
Extinguishing Media				•	

Section v - Reactivity	Data					
Stability	Unstable		Condition	ons to Avoid		
	Stable	Х		Unknown		
Incompatibility		None				
Hazardous Decomposition or E	Syproducts Sul	lfur oxid	es and	l bromides	;	
Hazardous	May Occur		Condit	ions to Avoid		
Polymerization	Will Not Occur	X	1	None		
Section VI - Health H	azard Data					
Route(s) of Entry:	Inhalatio	on? No		Skin? Y	es	Ingestion? Yes
Health Hazards (Acute and	Chronic)	Acute e	ye cor	itact: may	y caus	e irritation
Carcinogenicity:	NTP?		IAF	C Monogra	phs?	OSHA Regulation?
None		data		No data		No
Signs and Symptoms of Exp	osure			Мау сс	ause sk	kin or eye irritation
Medical Conditions Genera	lly Aggravated by	y Exposu	ire	None re	eporte	ed
Emergency First Aid Proce	dures					
	Rinse cor	ntacted	d areas	with copi	ous ar	nounts of water
Section VII - Precaut	ions for Safe	e Hand	ling a	nd Use		
Steps to be Taken in case N	Material is Releas	sed for Sp	pilled			
Wear eye and ski	n protection o	and mo	p/wipe	spill area	. Rinse	with water.
Waste Disposal Method						
Can be disposed	in the trash or	r down	the sinl	<		
Precautions to be Taken in	Handling and Sto	oring				
Avoid eye and sk	in contact					
Other Precautions						
Nor	ne					
Section VIII - Control	Measures					
Respiratory Protection (Spe		IOSH/M	SHA - a	pproved res	pirator	
Ventilation	Local Exhaust		No)	Special	None
	Mechanical (Ge	eneral)	No)	Other	None
Protective Gloves	Yes			Eye Protec	tion	Splash prof goggles
Other Protective Clothing or	Equipment	None re	equired	d		-
Work/Hygienic Practices	Avoid eye	e and sk	din con	tact		
Section V - Reactivity	y Data					



Special Fire Fighting Procedures N/A

Unusual Fire and Explosion Hazards

Material Safety Data Sheet
May be used to comply with OSHA's Hazard Communication
Standard. 29 CFR 1910.1200 Standard must be consulted for
specific requirements.

HOCKVIIIE, IND 20030	Signat	ure of Preparer (optional)		
14676 Rothgeb Drive Rockville, MD 20850		Prepared 07/01/03		
Address (Number, Street, City, State, Zip Code)		(301) 251-5990		
EDVOTEK, Inc.		Telephone Number for information		
		(301) 251-5990		
Manufacturer's Name	Emer	gency Telephone Number		
Section I				
Xylene Cyanol		applicable, or no information is available, the space must be marked to indicate that.		
IDENTITY (As Used on Label and List)		Note: Blank spaces are not permitted. If any item is not		

None

Section II - Hazardous Ingredients/Identify Information

Hazardous Components [Specific Chemical Identity; Common Name(s)] Other Limits
OSHA PEL ACGIH TLV Recommended % (Optional) This product contains no hazardous materials as defined by the OSHA Hazard Communication Standard. CAS # 2650-17-1

Section III - Ph	nysical/Chemica	I Characte	ristics

Boiling Point	No data	Specific Gravity (H ₂ 0 = 1)	No data
Vapor Pressure (mm Hg.)	No data	Melting Point	N/A
Vapor Density (AIR = 1)	No data	Evaporation Rate (Butyl Acetate = 1)	No data
Solubility in Water			

Soluble

Appearance and Odor color, liquid, no odor

Section IV - Physical/Chemical Characteristics

conon iv i nyolodi onomodi ondraotoriodoo						
ilash Point (Method Used) No data	Flammable Limits	LEL No data	UEL No data			
Extinguishing Media						

Special Fire Fighting Procedures N/A

Unusual Fire and Explosion Hazards

None

Stability	Unstable Conditions to Avoid			
	Stable	Χ	Unknown	
Incompatibility		None		
Hazardous Decomposition or B	yproducts Sul	fur oxid	es and bromides	
Hazardous Polymerization	May Occur		Conditions to Avoid	
	Will Not Occur	Χ	None	
Section VI - Health H	azard Data			
Route(s) of Entry:	Inhalation? No		Skin? Yes	Ingestion? Yes
Health Hazards (Acute and	Chronic) A	Acute e	ye contact: may cause	irritation
Carcinogenicity:	NTP?		IARC Monographs?	OSHA Regulation?
None	No data		No data	No
Signs and Symptoms of Exp	osure		May cause skin	or eye irritation
Medical Conditions Generally Aggravated by Exposure None reported				
Emergency First Aid Proced		ntacted	l areas with copious amo	ounts of water

Section VII - Precautions for Safe Handling and Use

Steps to be Taken in case Material is Released for Spilled

Wear eye and skin protection and mop/wipe spill area. Rinse with water.

Waste Disposal Method

Can be disposed in the trash or down the sink

Precautions to be Taken in Handling and Storing

Avoid eye and skin contact

Other Precautions

None

ection	VIII	- Control	Measures
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Respiratory Protection (Specify Type) NIOSH/MSHA - approved respirator

Local Exhaust None None Ventilation Other Mechanical (General) Eye Protection Protective Gloves Yes Splash prof goggles

Other Protective Clothing or Equipment None required

Work/Hygienic Practices Avoid eye and skin contact