Laboratory Safety

General Safety Rules

• Work in a laboratory only during regular, assigned period when an instructor is present, unless specific authorization has been given by the instructor to work in the laboratory at other times.

• Read carefully and observe fully all laboratory instructions. In case there is any doubt about any procedure, check with your instructor.

• Learn the location and proper use of emergency showers, fire extinguishers, and eye wash stations.

• Avoid inhaling chemical vapors or gases. Use fume hoods for hazardous materials.

• Immediately wash off and chemicals spilled on the skin with lots of water. In case of a serious spill, remove contaminated clothing immediately and flush affected area with lots of water.

• Do not eat, smoke, or drink in the laboratory. Do not bring any food items into the laboratory.

• Do not leave experiments in progress unattended without authorization.

• Keep working areas neat and clean at all times.

• Report all accidents to the instructor immediately.

Personal Protective Equipment

• All persons working with hazardous chemicals should wear gloves.

• All persons working with chemicals that could be splashed in the eyes are required to wear safety goggles or glasses.

• Contact lenses should not be worn in lab when hazardous chemicals or vapors are being used.

• Because of the danger of broken glassware or spilled chemicals, covered shoes should be worn in the laboratory. (No types of open toe shoes are permitted in labs.)

Chemical Safety

• Almost every chemical, whether solid, liquid or gaseous, is poisonous to the human body to some degree. Always use proper caution when handling chemicals.

• Consult a physician if you are pregnant or have any other medical condition that might render you susceptible to exposure to the chemicals used in this laboratory.

• When handling chemicals, keep your hands away from your face, eyes and body until your hands have been washed thoroughly.

• Do not taste any chemical. Label every container so items can be identified.

• When diluting acids, ALWAYS POUR ACID INTO WATER SLOWLY.

• Do not pipet anything into the mouth.

Waste Disposal

• Always treat laboratory glassware as if it were fragile. If glassware breaks, do not pick it up with your hands. Use a broom and dustpan, then place pieces in the cardboard box labeled “Glass Disposal Box.”

• Do not pour any chemicals down the drain. The instructor will advise you how to use the proper waste containers.

• Discarded animals parts must be placed in a red cardboard “Biohazardous Waste” box.

• Discarded sharp items including: scalpels, dissecting pins, probes, and needles must be placed in a red, plastic “Sharps Box.”
Safety Agreement
Biology III

I have read the Lab Safety Rules and procedures for the prevention of injuries in the laboratory, and I will observe them in my lab work.

INSTRUCTOR'S NAME:

______________________________

STUDENT'S NAME (PRINT)

______________________________

STUDENT'S SIGNATURE

______________________________

DATE

______________________________

NOTE - There is a COPY of this safety agreement form in the last section of the appendix for this Study Guide/Lab Manual for you to tear out and sign. Bring your signed copy to the first lab session. (Keep this copy here in your lab manual for reference.)
How to Use a Micropipettor

Micropipettors (a.k.a. pipette men) are used to measure and transfer small amounts of liquids (≤ 1 mL). You will find them in almost every biology laboratory in the world. They are expensive instruments (~$250/each) that must be shared by many scientists. Thus it is imperative that you treat our micropipettors as delicate and calibrated instruments. The scales on micropipettors are in microliters (1000 µL = 1 mL). In this course you will use four different types of micropipettors. Their properties are summarized in the table below.

<table>
<thead>
<tr>
<th>Name</th>
<th>P20</th>
<th>P200</th>
<th>P1000</th>
<th>Multichannel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tip color</td>
<td>yellow</td>
<td>yellow</td>
<td>blue</td>
<td>yellow</td>
</tr>
<tr>
<td>Minimum Volume</td>
<td>1 µL</td>
<td>20 µL</td>
<td>200 µL</td>
<td>5 µL</td>
</tr>
<tr>
<td>Maximum Volume</td>
<td>20 µL</td>
<td>200 µL</td>
<td>1000 µL</td>
<td>50 µL</td>
</tr>
</tbody>
</table>

A few important directions for the operation of any micropipettor:

1. Know the limits of your micropipettor (and don’t exceed those limits) of these pipettors. If you go above or below the minimum or maximum volume for a given pipettor, you will jeopardize the instrument’s calibration. (Note: just because you can dial 210 on a P200, doesn’t mean that you should!)

2. Set the desired volume by turning the centrally located rings clockwise to increase volume or counterclockwise to decrease volume.

3. Place a disposable plastic tip on the discharge end of the pipettor. NOTE: If sterile conditions are necessary, do not allow the yellow or blue plastic pipet tip to touch any object (including your hands, the bench, the side of a test tube, etc.).
4. **The plunger will stop at two different positions when it is depressed.** The first of these stopping points is the point of initial resistance and is the level of depression that will result in the desired volume of solution being transferred. Because this first stopping point is dependent on the volume that is being transferred, the distance you have to push the plunger to reach the point of initial resistance will change depending on the volume being pipetted. The second stopping point can be found when the plunger is depressed beyond the initial resistance until it is in contact with the body of the pipettor. At this point, the plunger cannot be depressed further. This second stopping point is used for the complete discharging of solutions from the plastic tip. You should not reach this second stop when drawing liquid into the pipettor, only when expelling the last drop. Before continuing, practice depressing the plunger to each of these stopping points until you can easily distinguish between these points.

5. **Depress the plunger until you feel the initial resistance (first stop) and insert tip into your solution, just barely below the surface of the liquid and not as deep as possible.** The wide (top) portion of the disposable pipette tip should never be underwater. Only the disposable pipet tip should touch the liquid; the pipettor should never touch any of the liquids. You should never rest the pipette tip on the bottom of the container (even if you have a shallow volume of liquid) because this could lead to inaccurate measurements.

6. **Carefully and slowly release plunger.** If you release the plunger too quickly, it will suck liquid up into the pipettor and damage it. **NOTE:** If the solution you are pipetting is viscous, allow the pipet tip to fill to final volume before removing it from solution to avoid the presence of bubbles in the plastic tip, which will result in an inaccurate volume.

7. **Discharge the solution into the appropriate container by depressing plunger.** This time, depress the plunger to the point of initial resistance, wait one second, and then continue pressing the plunger as far as it will go in order to discharge the entire volume of solution.

8. **Discard the tip by pressing down on the tip discarder over a waste container.**

9. **Always change tips between solutions.** You do not want to contaminate or mix your solutions with a dirty pipet tip.
Introduction to Spectrophotometry

Focused Reading
• p 192-194 “How does photosynthesis…” to “Several pigments…”

✓ Special Note
• Bring a calculator to lab today

Introduction
The purposes of this laboratory are to introduce you to:
1. Conventions used in making solutions: molarity, and percent.
2. Spectrophotometry and the use of the microplate reader.
3. Procedures for obtaining, recording, and analyzing data.
4. Conventions used in presenting data in graphs.
5. Procedures for planning and working through a series of related experiments.

Note: Record all your data, answers to questions and problems, and notes directly in this lab manual. Keep this lab manual beside you and write in it as you work. Good note taking skills are absolutely critical to successful scientific experiments.

Concentrations of solutions
It is important that you understand the units of the metric system (i.e., milli and micro). If you do not, please review these units of measurement as well as the Celsius (centigrade) temperature scale.

In the instructions below, the solute is the substance dissolved, the solvent is the liquid in which the solute is dissolved, and the resulting mixture is the solution.

Molar Concentrations
In technical terms, a mole of a compound is 6.02 x 10^{23} molecules of that compound. Practically speaking, a mole is the compound’s molecular weight in grams. A one molar (1.0 M) solution has one mole of a compound (the solute) dissolved in solvent so that the final volume is 1000 mL (one liter). The molecular weight of NaCl is 58.54. Therefore:
• A 1.0 M solution of NaCl has 58.54 g NaCl dissolved in dH_2O with a final volume of 1000 mL (or 5.85 g in 100 mL).
• A 0.1 M solution of NaCl has 5.85 g in 1 liter - or 0.585 g in 100 mL - or 0.058 g in 10 mL.
• A 0.2 M solution of NaCl has 11.71 g in 1 liter - or 1.17 g in 100 mL - or 0.117 g in 10 mL.

Concentration in Percent (Weight/Volume = w/v)
By definition, percent means “in a hundred” and by convention, a 10% w/v solution contains 10 grams of a solute in a total volume of 100 mL of the solution. A w/v solution is not made by adding 10 grams of a solute to 100 mL of solvent, but instead by dissolving 10 grams solute in enough solvent to dissolve the solid and then more solvent is added to reach a total volume of 100 mL of the solution.

Concentrations in Percent (Volume/Volume = v/v)
Aqueous solutes may be specified as percent solutions volume/volume (v/v). Volume/volume solutions differ from weight/volume solutions – and it is important to understand the difference. One hundred mL of a 5% v/v solution will have 5 mL solute diluted to 100 mL with solvent. Thus, a 5% v/v aqueous solution of ethanol is made by diluting 5 mL 100% ethanol with enough dH_2O (95 mL) to make a total of 100 mL (or by diluting 5 liters 100% ethanol with 95 liters dH_2O).
A handy formula for calculating v/v solutions:

\[ C_s V_s = C_w V_w \]

- \( C_s \) = concentration of the stock (original) solution
- \( V_s \) = volume of the stock solution (usually what you will solve for)
- \( C_w \) = concentration of the working solution desired (more dilute than the stock)
- \( V_w \) = total volume of the working solution desired (the solution you are making)
- \( V_w - V_s \) = volume of the solvent (often water) to be mixed with \( V_s \)

Suppose you have a stock bottle of 70% ethanol and you need to make 1 L of 50% ethanol.

\[ C_s V_s = C_w V_w \]
\[
70\% \times V_s = 50\% \times 1 \text{ L} \\
V_s = (50\% \times 1 \text{ L})/70\% \\
V_s = 0.714 \text{ L} \\
\]

\[ V_w - V_s = 1 \text{ L} - 0.714 \text{ L} = 0.286 \text{ L} \]

Thus you would measure out 714 mL of 70% ethanol and mix it with 286 mL of water to get your 1 L of 50% ethanol.

This formula is simply determining proportions, so it can also be used to calculate how to dilute any solution. To make \( V_w \) units (mL, L, etc.) of a solution at \( C_w \) (M, mM, g/L, % etc.) from water and a stock solution of \( C_s \), you need to be certain you are using equivalent measures of volume for both \( V_s \) and \( V_w \) (i.e., make sure both are measured in mL or both are measure in L, etc.) and you need to be certain that you are using equivalent measures of concentration (i.e., make sure both \( C_s \) and \( C_w \) are both in mg/L or both in mM, etc.). Then you simply solve for \( V_s \) to determine how much stock solution you will need. Then you simply subtract \( V_s \) from \( V_w \) to determine how much solvent should be added. Here's an example. Suppose you have a 9.5 mM solution of KCl and want to make 927 mL of 4 mM KCl. You need to know how many mL of your 9.5 mM KCl solution you will mix with how many mL of H\(_2\)O in order to make 927 mL of 4 mM KCl. Using the \( C_s V_s = C_w V_w \) formula:

\[ C_s V_s = C_w V_w \]
\[
9.5 \text{ mM} \times V_s = 4 \text{ mM} \times 927 \text{ mL} \\
V_s = (4 \text{ mM} \times 927 \text{ mL})/9.5 \text{ mM} \\
V_s = 390.3 \text{ mL} \\
\]

\[ V_w - V_s = 927 \text{ mL} - 390.3 \text{ mL} = 536.7 \text{ mL} \]

Thus you would measure out 390.3 mL of 9.5 mM KCl and mix it with 536.7 mL of water to get your 927 mL of 4 mM KCl.

**Exercise 1: Evaluating a w/v solution**

Exercise 1 is a demonstration experiment for you to observe and think about.

1. Locate the graduated cylinder with a NaCl solution in it. Before lab your instructor mixed 5 grams (g) of sodium chloride (NaCl) with exactly 50 mL of distilled water (dH\(_2\)O) and the solution was stirred vigorously to dissolve the salt.

2. Determine the excess volume by reading the volume of the graduated cylinder. This solution represents a failed attempt to make a 10% w/v solution. The number of mL in excess of 50 represents the volume of liquid displaced by the 5 g of dissolved NaCl.

3. What is the actual percentage of NaCl (w/v) in this solution?
Exercise 2: Making a v/v solution of neutral red

On the side or front bench is a stock solution (100%) of a dye, neutral red. Use this stock solution and water to make 6 mL of a 4% v/v solution. Label your tube and save this solution; you will use it later today. Make sure to mix the solution thoroughly and use parafilm to cover the top of the test tube during mixing.

\[
\text{Cs} \quad V_s \quad C_w \quad V_w \quad V_w - V_s
\]

\[_______ \text{mL 100% neutral red} + \quad _______ \text{mL H}_2\text{O} = \quad 6 \text{ mL 4% neutral red}\]

Exercise 3: Diluting solutions

Using the 4% (v/v) neutral red solution you prepared in exercise 2, set up the following solutions in eight plastic microfuge tubes using distilled water and neutral red. Label the tubes with the concentrations in the left column of the table below. (Note: you can use a Sharpie marker to write directly on the tubes.)

You must calculate the volumes and fill in the table for the two unshaded columns before you make your solutions. Each tube should contain a final volume of 1000 µL.

<table>
<thead>
<tr>
<th>neutral red concentration</th>
<th>4% neutral red solution to add</th>
<th>distilled water to add</th>
<th>concentration of neutral red solution (mg/mL)</th>
<th>concentration of neutral red solution (M)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 %</td>
<td>0 µl</td>
<td>1000 µl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.36 %</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.84 %</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.56 %</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.96 %</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2.52 %</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4 %</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 %</td>
<td>1000 µl</td>
<td>0 µl</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complete these two columns later (when answering study question #6)

Spectrophotometry

A solution, such as neutral red, appears colored because it absorbs certain wavelengths of light in the visible spectrum and transmits or reflects others. Each solution with a different solute has its own characteristic absorption properties or "spectrum." A spectrophotometer is an optical machine that measures and lets you see (sense) how much light energy is transmitted by a substance in solution at different wavelengths of radiant energy. Biologists use the spectrophotometer for two different purposes (we will do both in the laboratory today):

1. to determine the absorption spectrum of a pure substance in solution
2. to determine the concentration of a solution.
A spectrophotometer consists of a white light source (light of all visible wavelengths), a prism or diffraction grating that separates the light into different wavelengths, a slit through which a narrow beam of the desired wavelength ($\lambda$) passes (the incident light, $I_0$), a sample solution holder, a photosensitive detector that measures the energy of light transmitted through the solution ($I$), and a recording device that displays the amount of transmitted light energy digitally or on a dial. See Fig. 1 below.

![Schematic diagram of a spectrophotometer](image)

**Figure 1. A schematic diagram of the components of a spectrophotometer.** The arrows indicate the pathway of light.

Transmittance is the ratio of the transmitted light energy ($I$) to the incident light energy ($I_0$); percent transmittance is 100X that ratio. Transmittance, however, is not proportional to solute concentration, so it is usually converted into absorbance that is proportional to solute concentration. Digital spectrophotometers have readouts for both percent transmittance and absorbance, but we will always measure the absorbance.

$$\%T = \left(1 - \frac{I}{I_0}\right) \times 100$$

$$\text{Abs.} = \log_{10} \left(\frac{100}{\%T}\right)$$

**Microplate Reader**

Figure 1 is a simplified diagram of a spectrophotometer that can measure one sample at a time. In our experiments, we will be using a microplate reader that is capable of measuring the absorbance of 96 samples in just a few seconds. The basic design is exactly the same; a selected wavelength of light passes through the samples and a phototube measures the amount of light transmitted through the sample, which the plate reader converts to absorbance. However, the samples are located in microwells that are arranged in an 8 x 12 matrix in one plastic plate (see figure 2). You can put your samples in any or all of the microwells. The plate is moved over an array of eight fiber optic light sources and eight phototubes. Each row of eight is scanned and then the plate advances by one row and the process continues until all 12 rows are scanned. The absorbance data are then displayed on a screen in an 8 x 12 array. You also can program the plate reader to measure the absorbance of all 96 samples at time intervals of your choice (e.g., every 30 seconds). You should take advantage of these capabilities when you design your experiments for next week.

![Schematic diagram of a microplate reader](image)

**Figure 2. Schematic diagram of a microplate reader.** Samples are placed in the 96 microwells, analyzed by the eight channel spectrophotometer.
The Absorption Spectrum

Because solutions of pure substances do not absorb the energy of all wavelengths of light equally, a substance may be identified by the unique pattern of wavelengths absorbed. The chlorophylls in plants absorb strongly in the blue wavelengths (about 450 nm) and red wavelengths (about 650 nm), but reflect the green wavelengths (about 525 nm). A plot of absorbance versus visible wave lengths (400 to 700 nm) for a solution of chlorophyll a shows two major peaks, one at 450 and one at 650 nm, and a valley from 500 to 625 nm (See Figure 3). This spectrum is characteristic for chlorophyll a and may be used as an aid in its identification.

By measuring the absorbance of an uncharacterized solution over a range of wavelengths and plotting the absorbance value on the Y-axis and the wavelength on the X-axis, one can determine the absorption spectrum of a sample. The absorption maximum of any pure substance in solution is the wavelength where absorption is the greatest.

Figure 3. The absorption spectrum of chlorophyll a. The absorbance of visible light by chlorophyll a is measured spectrophotometrically as a function of wave length. The absorption maximum is about 460 nm.

Standard Concentration Curve & Determining the Concentration of a Substance

One can construct a standard concentration curve of a solution by measuring the absorbance of several different known concentrations of the solution and graphing the results by plotting absorbance on the Y-axis and concentration on the X-axis. Spectrophotometry can be used to measure the absolute or relative concentration of a characterized substance in solution. To determine the absolute concentration of a pure substance, one first constructs a standard concentration curve from known concentrations and then takes the absorbance reading of the unknown concentration. The unknown concentration can be determined from the standard curve by drawing a horizontal line on the graph parallel to the X-axis and through the point on the Y-axis that corresponds to the absorbance. This line will intersect the standard curve; at this intersection, a vertical line is dropped to the X-axis and the concentration read from the X-axis.

Three factors are important in determining unknown or relative concentrations. The absorption maximum should be used, and absorbance rather than percent transmittance should be plotted because absorbance is directly proportional to concentration and transmittance is not. You cannot measure outside the standard curve.

Figure 4. An example of a standard curve. A best fit line has been generated and the resulting equation and $r^2$ value are shown below the X-axis label.
**Exercise 4: Determination neutral red’s absorption maximum**

For each of your eight different dilutions of neutral red (made in exercise 3) pipette 200 ul into triplicate wells of your 96-well plate. Keep track of which concentration corresponds with each particular well. You will collect the data for experiments 4 and 5 at the same time (see below).

**Exercise 5: Generating a neutral red standard curve**

You will generate these data at the same time as those from Exercise 4. Use only the data from the wavelength that is the absorption maximum for neutral red. With the appropriate data, you will generate a curve to measure the unknown concentration of a solution. Follow the directions below to collect the data.

**Using the Plate Reader** (a type of spectrophotometer)

1. Your instructor will have turned on the plate reader and let it warm up. Open the KCjunior program on the PC and click OK – the software will start up.

2. Click on the “Open Protocol” button and select the “340 nm endpoint” option then click on the “open” button, then on the “read plate” button.

3. You will see a read plate dialog box. Click on the “read plate” button. (There’s no need to fill in the results ID or plate description boxes).

4. You will get another read plate dialog box. Place your 96-well plate in the carrier and make sure the lid is shut, then press the OK button.

5. You should hear the machine moving and in a few seconds you should see an 8x12 report of the absorbance values in all 96 wells.

6. Go up to “Results” in the upper left menu portion of the monitor and select the “print results” option from the drop-down menu. (Make sure “brother HL-514 series” is selected in the printer options). Click the “OK” box to print your data.

7. Close your results using the “close results” command under the “Results” menu in the upper left. (There is no need to save your results on the computer after you have your printout).

8. Now you will repeat this procedure for five more wavelengths of light, using protocols named “405 endpoint”, “450 endpoint”, “490 endpoint”, 595 endpoint”, and “650 endpoint”.

9. When you have collected data for all six wavelengths, remove your plate from the plate reader and take your six printouts back to your bench to analyze your data.

<table>
<thead>
<tr>
<th>Wavelength (nm)</th>
<th>Absorbance values for 3.4% neutral red</th>
</tr>
</thead>
<tbody>
<tr>
<td>340</td>
<td>Value 1</td>
</tr>
<tr>
<td>405</td>
<td></td>
</tr>
<tr>
<td>450</td>
<td></td>
</tr>
<tr>
<td>490</td>
<td></td>
</tr>
<tr>
<td>595</td>
<td></td>
</tr>
<tr>
<td>650</td>
<td></td>
</tr>
</tbody>
</table>

To generate an absorption spectrum that will reveal the absorption maximum, plot the average absorbance of the 3.4% neutral red solution on the graph paper on the next page. Use absorbance (which is a ratio and therefore has no units) on the y-axis and the wavelength of light (in nm) on the x-axis.
Exercise 4 Data analysis:
Use your graph on the previous page to identify the wavelength that produced the highest absorbance values. This wavelength is the absorption maximum for neutral red.

Neutral red absorption maximum: ________ nm

All subsequent experiments should use only the data collected at the absorbance maximum wavelength.

*What is the advantage of using this absorbance max for subsequent experiments with neutral red?

*What can you predict about the absorbance maximum wavelength for a blue dye?
Exercise 5:

Look back to your printouts and use only the absorbance data that were generated at neutral red’s absorption maximum wavelength to model the relationship between the concentration of neutral red and the absorbance. This is done by plotting the absorbance values for the set of known (standard) concentrations and then performing a linear regression to get the equation of the line that best describes that curve. This equation is your mathematical model. The process is referred to as determining the standard curve for neutral red. In the images below, neutral red is abbreviated “NRD,” and absorbance is abbreviated “ABS”.

To plot the standard curve, open a spreadsheet in Excel, and set up the following rows and columns.

2) Enter your data into the columns labeled 1st, 2nd, and 3rd well.
3) Calculate the average absorbance value for each concentration of neutral red.

In the ‘Avg ABS’ column, enter the formula that tells Excel to take the average of the first 3 columns and propagate that formula down the column.

The absorbance values determined for each well also include some contribution due to the plastic and other parts of the solution that are not neutral red. These values are removed from the calculations by correcting for the background absorbance as follows:

5) In cell F4 type the formula =E4-$E$4 and propagate the function down the column. This tells Excel to subtract the value in E4 (the ‘0% NRD’ average) from each of the other averages so that the data are corrected for background absorbance. (The use of $E$4 makes the program always go back to the same cell in the table.)

6) Plot the points as follows:

   a) Select the NRD concentrations by clicking and dragging A3 to A11. Hold the ‘command’ key (⌘) while selecting the Background Corrected Avg abs values (F3-F11).
b) In the menu bar above the spreadsheet select ‘Chart’, then ‘Scatter’, and ‘Marker scatter’. The graph should appear.

7) Determine the linear regression as follows:
   a) Single click on a data point on the graph—this should cause all of the points to become selected.
   b) Right click on that data point and choose ‘add trend line’.
   c) Make sure that the ‘type’ of trend line chosen is ‘linear’ but don’t choose OK yet.
   d) In the left part of the box choose ‘option’ then click on ‘show equation’ and ‘show R-squared value’
   e) Now click OK. The equation that appears on the graph is the equation modeling the relationship between the Concentration of NRD and the Absorbance value (at NRD’s ABS<sub>max</sub> wavelength)

   **Print your neutral red standard curve and attach it here:**
**Exercise 6:**
**Using your standard curve to determine concentrations**

Obtain four neutral red solutions of unknown concentration from your instructor. Determine the concentration of your four unknowns by putting each unknown (A-D) into three individual wells (200 µL/well). Record well names (e.g., H3 - H6) corresponding with each unknown on the chart below. Use the plate reader at the wavelength of light that is absorbed maximally by neutral red. Subtract the absorbance of water (the blank) from these values and use these corrected values to determine the concentrations of the four unknowns. Using the equation determined in the standard curve, determine the concentration of your unknown dilutions of neutral red.

For each unknown, the unknown concentration can be determined from the standard curve by drawing a horizontal line on the graph parallel to the X-axis and through the point on the Y-axis corresponding to the corrected absorbance for the unknown. This line will intersect the standard curve; at this intersection, a vertical line is dropped to the X-axis and the concentration read from the X-axis. The unknown concentration can also be determined by using the experimentally determined ‘y’ value (the solution’s blank corrected average Abs) in the equation provided by the linear regression.

1) On the same spreadsheet used in Exercise 5, set up a table containing the triplicate absorbance readings for unknowns A-D, the average absorbance, and the blank-corrected absorbance.

2) Determine the estimated concentration for solutions A-D first by eye from your graph, and then using the equation from the linear regression of the standard curve. Place those estimates in the table below.

3) Ask you instructor to reveal the actual concentrations of the four unknowns. How well did your standard curve predict the concentration on the four unknown solutions?

<table>
<thead>
<tr>
<th>Neutral Red Unknown</th>
<th>‘By Eye’ Concentration</th>
<th>Calc. Estimated concentration (from your std. curve eq.)</th>
<th>Actual concentration (provided by instructor)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Exercise 7:**
**Determining the absorption maximum for NADPH and NADP⁺**

Over the subsequent two weeks, we will use NADPH and NADP⁺ as indicators of enzyme activity. Consequently, we will need to know which wavelengths of light will give the best information on NADPH concentration. You will want to perform a series of experiments similar to experiment four. Put 200 µL NADPH into each of three empty wells and 200 µL of NADP+ into another three empty wells. Record your averaged results in the table below.

<table>
<thead>
<tr>
<th></th>
<th>340 nm</th>
<th>405 nm</th>
<th>450 nm</th>
<th>490 nm</th>
<th>595 nm</th>
<th>650 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>NADPH</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NADP+</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Look at your data above. Which wavelength is absorbed the best and the amount of light absorbed by each chemical. Record your responses in the table below. Verify your results with the instructor before you leave today.

<table>
<thead>
<tr>
<th>Absorbance maximum (nm)</th>
<th>Amount of absorbance at maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>NADPH</td>
<td></td>
</tr>
<tr>
<td>NADP+</td>
<td></td>
</tr>
</tbody>
</table>

**Cleaning Up**

Leave your workstation cleaner than you found it. Although this step may seem obvious, it is quite important. It is basic laboratory courtesy to leave the workspace as clean as you found it with equipment back in its proper place(s).

BEFORE LEAVING make sure that all members of the lab group have access to the Excel file Although one of you may have created it—all of you are responsible for understanding how to determine and use a standard curve.

❖ **Study Questions** *(to answer on your own time):*

1. How do you account for the excess volume you observed in Exercise 1?

2. The solution made in experiment 1 does not constitute a 10% w/v solution of sodium chloride but something less than that. Calculate the actual percentage from your data.

3. Why do you have to dissolve the solute in a volume of solvent less than the final volume you eventually want? Does it matter if the salt is added first or second to the graduated cylinder?

4. Describe precisely how you made up the 4% v/v solution of neutral red.

5. In Experiment 1, does it matter whether the solute is added to the graduated cylinder before or after the water?
6. The aqueous stock solution of neutral red was actually a 1.0 M solution. Determine the molar concentration of the six solutions you made. If neutral red has a molecular weight of 87, how many µg/µL are contained in your 4% (v/v) solution? Calculate the concentration of neutral red in each tube (µg/µL) and add these data to the table from exercise 3.

7. Describe how to prepare 50 mL of 70% ethanol when your only source is a stock container of 95% ethanol.

8. What is the molarity of your 10% w/v NaCl solution?

9. What is the percent concentration of a 2 M NaCl solution?

10. The molecular weight of Na₂C₂O₄ is 106. Describe how you would make up 100 mL of a 0.15 M solution.

11. What is the absorption maximum for NADPH?

Lab Notes
I. Introduction

Enzymes are biological catalysts with remarkable power, increasing reaction rates by at least a million-fold. They increase reaction rates by lowering activation energies, allowing chemical reactions to proceed under physiological conditions. Enzymes are highly specific as to substrates and reactions catalyzed. They are usually proteins, although some enzymes are other types of biological molecules. Enzymes function best in dilute aqueous solutions under limited conditions of temperature, pH, and salt concentration. Some enzymes require one or more non-protein components, called “coenzymes” if they are organic, and “cofactors” if inorganic. Isocitrate dehydrogenase [IDH] requires Mg$^{2+}$ or Mn$^{2+}$ as a divalent metal cofactor.

IDH is a ubiquitous enzyme found in all living organisms and has two catalytic activities (Figure 1). As its name implies, IDH removes hydrogens from its primary substrate, isocitrate. It also converts NADP$^+$ to NADPH. In addition, it is a decarboxylase, removing a CO$_2$ from the six-carbon substrate to generate a five-carbon product, $\alpha$-ketoglutarate.

Figure 1. IDH catalyzes the sequential dehydrogenation and decarboxylation of isocitrate to $\alpha$-ketoglutarate. Note that IDH catalyzes two sequential reactions to convert isocitrate into $\alpha$-ketoglutarate.
Two distinct forms of IDH are found in higher organisms. They differ in their distribution within the cell and in whether they use NADP⁺ or NAD⁺ as a substrate. The soluble form of IDH requires NADP⁺ (Figure 2). This NADP⁺-dependent form of IDH is thought to be the only IDH in most bacteria. In higher organisms, this form of IDH is found in the cytoplasm of cells within all organs and tissues, and it is used in lipid synthesis. The other form of IDH is NAD⁺-dependent; it is localized in the mitochondria of eukaryotic organisms. You may be familiar with this second form of IDH from previous study of the Krebs cycle (and if you’re not, you will meet it again in the bioenergetics section of this course). Both forms of IDH require a divalent metal ion.

![Figure 2. The molecular structure of NADP⁺. The active site is where the hydrogen atom will be added to convert NADP⁺ to NADPH. This diagram illustrates what the letters N-A-D-P represent.](image)

NEWS ITEMS: A team of researchers found a species of voles that was resistant to mutations caused by radiation. When they analyzed their cells, they found that the voles had elevated levels of IDH, which they believe is protecting them from radiation-induced mutations. (Science 273)

NADP⁺-dependent IDH activity is especially high in cardiac tissue and is often monitored in the blood of heart attack patients. Detectable IDH activity in the arterial blood suggests severe tissue damage with leakage of the soluble (cytosolic) IDH into the blood system.

How IDH activity is measured

IDH activity routinely is measured using a spectrophotometer to monitor the reduction of NADP⁺ to NADPH. While performing assays, the spectrophotometer is set at 340 nm, the absorption maximum of NADPH (and results from last week’s lab). Assays are performed at a standard temperature, usually 25°C to 30°C.

Designing a protocol

Before a scientist begins an experiment, he or she must first define a problem and suggest possible explanations based upon previous knowledge or observations. In other words, develop an hypothesis, which might be considered an “educated guess” or a tentative explanation as to the cause and effects relating to that problem. A good hypothesis is one that is testable and fosters predictions that consider one variable at a time. The hypothesis may turn out to be incorrect, but it is a good hypothesis if it can be tested. In fact, an hypothesis that cannot be tested is useless to science - it may be good philosophy, but not good science. Hypotheses can not be proven to be correct - they may be tested extensively and rigorously and they may be proven to be incorrect, but an hypothesis can never be proven to be true.
A scientist must first define a problem and then develop an hypothesis. Next, one must devise predictions that will hold, or will not hold, if the hypothesis were true. These predictions lead to experiments. Many experiments may be possible, and all may be tried eventually; however, it is important to perform one discrete experiment at a time. After designing an experiment, our scientist must outline a series of logical procedures to be completed in the laboratory or in the field. This written sequence of steps is called a protocol. A well-planned protocol will include the following elements:

1. An outline of the sequence of detailed procedures.
2. Calculations of volumes, concentrations, etc., of all reagents to be used.
3. Tables constructed for recording data.
4. Procedures for testing and organizing data for presentation.

**Experiment 1: Performing an IDH assay**

**Hypothesis 1:** A successful assay for IDH activity simultaneously requires the enzyme (IDH), as well as the substrate isocitrate and NADP+.

**Hypothesis 2:** Under ideal conditions, IDH activity will be linear for at least three minutes.

To test your hypotheses, you will need to set up assays as in the table below. You should ask yourself, "What is the purpose of each assay? And why are assays done in triplicate?"

**Important notes for all experiments:**

1) Bubbles and/or drops of liquid clinging to the sides of your wells (rather than mixed in the bottom) will yield unusual absorbance readings. Get in the good habit of ejecting your liquids into the bottom of the wells and then checking your plate for bubbles and drops on the sides before you place it in the spectrophotometer.

2) You will usually initiate the reactions by adding 10 µL of isocitrate solution as the last step. (You will use a multi-tip pipet, at the plate reader, to add isocitrate to all wells that get isocitrate.) HOWEVER, in later experiments, isocitrate will not always be the last reagent added. Look closely at the instructions!

<table>
<thead>
<tr>
<th>Well</th>
<th>Buffer</th>
<th>Substrate 1</th>
<th>Enzyme</th>
<th>Substrate 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>A 1</td>
<td>200 µL</td>
<td>0 µL</td>
<td>0 µL</td>
<td>0 µL</td>
</tr>
<tr>
<td>A 2</td>
<td>200 µL</td>
<td>0 µL</td>
<td>0 µL</td>
<td>0 µL</td>
</tr>
<tr>
<td>A 3</td>
<td>200 µL</td>
<td>0 µL</td>
<td>0 µL</td>
<td>0 µL</td>
</tr>
<tr>
<td>B 1</td>
<td>170 µL</td>
<td>10 µL</td>
<td>20 µL</td>
<td>0 µL</td>
</tr>
<tr>
<td>B 2</td>
<td>170 µL</td>
<td>10 µL</td>
<td>20 µL</td>
<td>0 µL</td>
</tr>
<tr>
<td>B 3</td>
<td>170 µL</td>
<td>10 µL</td>
<td>20 µL</td>
<td>0 µL</td>
</tr>
<tr>
<td>C 1</td>
<td>180 µL</td>
<td>10 µL</td>
<td>0 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>C 2</td>
<td>180 µL</td>
<td>10 µL</td>
<td>0 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>C 3</td>
<td>180 µL</td>
<td>10 µL</td>
<td>0 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>D 1</td>
<td>170 µL</td>
<td>0 µL</td>
<td>20 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>D 2</td>
<td>170 µL</td>
<td>0 µL</td>
<td>20 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>D 3</td>
<td>170 µL</td>
<td>0 µL</td>
<td>20 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>E 1</td>
<td>160 µL</td>
<td>10 µL</td>
<td>20 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>E 2</td>
<td>160 µL</td>
<td>10 µL</td>
<td>20 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>E 3</td>
<td>160 µL</td>
<td>10 µL</td>
<td>20 µL</td>
<td>10 µL</td>
</tr>
</tbody>
</table>

**FYI:** Standard assay conditions in these experiments include 0.2 M Tris buffer at pH of 8.5 with 1 mM MgCl₂, 0.144 mM NADP+, and 0.23 mM isocitrate.
Procedure

1. Use the appropriate pipettors to add the appropriate amounts of buffer, NADP+, and IDH to the appropriate wells. (You will add the isocitrate at the last second before you start the plate reader.)

2. Use KCJunior software to select the “IDH (340 nm every 30 sec)” option and click on the “open” button, then on the “read plate” button.

3. Click on the “read plate” button. (There’s no need to fill in the results ID or plate description boxes.)

4. Place your 96-well plate in the carrier (with well A1 at the top left position). Add the last ingredient (isocitrate) to the appropriate wells as rapidly and simultaneously as you can. Then quickly shut the lid and press the OK button to start recording the absorbance.

5. You should hear the machine shaking the plate (to mix the solutions) and moving the plate. The plate reader will wait 30 seconds and repeat the procedure six more times for a total of three minutes (and seven total reads).

6. When the plate reader has finished all seven readings, go up to “Results” in the upper left menu portion of the monitor and select the “print results” option from the drop-down menu. (Make sure “Brother HL-514 series” is selected in the printer options). Click the “OK” box to print your data. You should get a two-page printout with seven sets of data.

7. Set up an Excel file as below, and type your raw triplicate absorbance data into the appropriate cells. This spreadsheet is very similar to the one you used last week—it just has more parts. The “corrected” reading for each assay is found by subtracting the average reading of a “control” experiment, in this case wells A1-3. (Would wells B1-3, C1-3, or D1-3 provide better “control” data?)

<table>
<thead>
<tr>
<th>Time</th>
<th>A1 abs</th>
<th>A2 abs</th>
<th>A3 abs</th>
<th>A avg abs</th>
<th>Corrected A average absorbance</th>
<th>B1 abs</th>
<th>B2 abs</th>
<th>B3 abs</th>
<th>B avg abs</th>
<th>Corrected B average absorbance</th>
<th>C1 abs</th>
<th>C2 abs</th>
<th>etc…</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>1</td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>1.5</td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>2</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>2.5</td>
<td></td>
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<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

8. In Excel, prepare a graph of your data with time (independent variable) on the x-axis and corrected average absorbance (dependent variable) on the y-axis. Include data for all experimental conditions on one graph, so that the graph will have a separate line representing each experimental condition. See the subsequent pages for instructions.
A Beginner's Guide to Graphing with Excel

Click on the Excel icon in the tool bar. When Excel opens, you will see a spreadsheet. In the first cell, enter a name for your X axis values (for instance, "Time"). Enter the appropriate time values in the first column. Enter headings for dependent variables in subsequent columns. For example, at the top of the second column, enter "A1 abs." representing the absorbance values for well A1 of the 96 well plate. Enter the raw absorbance data for well A1 at each time point. And repeat for all other wells with sample in them.

Leave two blank columns to the right of each set of triplicate data, and type headings for these columns as in the image below. At this point, you should have a spreadsheet whose first portion looks similar to the following image:

![Spreadsheet Screenshot](Image)

Remember that, using Excel, we can determine the average values for the experimental replicates. You can type in the formula as you did in last week, or else click on the empty cell for the average values of well A at time 0 and click on the Insert menu at the top. Select the Function option and double click Average in the Formula Builder box that pops up. You might notice that Excel "wants" to calculate the average of the values in cells A2 to D2 but our experimental values are in cells B2 to D2. Using the mouse, make any necessary changes to these cell designations by highlighting the cells that you want to average (in this case B2, C2, & D2) or typing B2:D2 into the box at the bottom of the Formula Builder window. Hit the return key, and the average value of your triplicates will be inserted into cell E2. To determine the averages for the other triplicate readings, click in cell E2 and propagate the formula down the column as you have done before. When you let go you should have averages in every cell. Repeat this process for the raw data in the B wells, the C wells, etc. until you have computed the averages for all your data.

The main window should now look something like this:

![Spreadsheet Screenshot](Image)
Now “background correct” the data by subtracting the absorbance that is due to the buffer and the plastic. The remaining values will represent the true absorbances of just the substance we are interested in. Wells A1-3 contained just buffer. We will subtract the A averages from all averages to achieve the corrected values. Note that this will result in values of 0 for our corrected A averages. Click in the first open cell of the “corrected A average” column (cell F2). Type “=E2-$E2” and hit enter. This tells Excel to subtract the value in cell E2 from itself and put the result in this cell. The dollar sign is important for later. Propagate the function down the column. Notice that the formula is copied such that each time it refers to and subtracts the cell that has the average for the cell to its left.

To set up the functions in the other background corrected average columns, copy and paste cells F2:F8 into cells K2:K8 and every other column where corrected values are required. The formula, not the absolute values, is transferred. The dollar sign tells the pasted formula always to subtract the 0 minute values. You should now have corrected values in every column. Spot check the values by eye to make sure things look right. If not, go back and correct your formulas. If you want a different condition to be used as your background correction, this formula is where you set it up. For example using ‘E2-$E2’ tells Excel to subtract the values from the E columns but match the time point (i.e., 15 seconds or 30 seconds), so in column K, cell K4 would be ‘K4-E4’. However ‘E2-$E$2’ will have Excel always subtract the value found in cell E2, so K4 would have K4-E2.

To graph the data, highlight column A (time values), hold the apple/command key down to highlight a second column while keeping the first column highlighted. **NOTE: Be sure to highlight columns in one dragging motion; it won’t work if you click each individual cell separately.** It’s OK to include the cells that contain the headings, as Excel usually recognizes those as labels. Go to the green menu about the spreadsheet and select Chart, and then the Scatter option at the far right. Choose one of these five graph types (which do you think is most appropriate?). At any time later you can switch to a different one of these five types.

You will then see a graph of your data with all of the data plotted as separate colors on the same graph. Note that Excel automatically used the first highlighted column as the X axis values, with the remaining columns as Y axis values in separate lines on the graph. If you ever want to change which data are on which axis, just right-click on the white area in the graph above the key and pick Select Data. Position the cursor in the “X values” or “Y values” box, and then highlight the cells whose data are meant for that axis.

You can change the appearance of your graph by double-clicking and/or right-clicking various features on the graph and selecting the formatting options, or by using the Formatting Palette.
• **Remove gridlines:** Right-click the lines themselves and select Delete.

• **Remove or modify the key (confusingly called a legend by Excel):** If you have only one data series represented as a line on the graph, just highlight the "series 1" key and hit delete. However, if you have a graph of several lines (as you will need to eventually), you should keep and properly label the keys to indicate what each line represents, if Excel hasn’t done this automatically. Go back to the Select Data option as above, and type a new title in the “name” box for each data series that you can highlight at the left. (Note that you can also add data lines to the graph with the Add button; in the box that comes up, just highlight the proper data cells for the X and Y axis).

• **Label the axes and title the graph:** Click on the graph so that it has blue circles at the corners and blue squares along the sides, and make sure the Formatting Palette is visible (if not, find it in the View menu). Under Chart Options, the Titles section lets you easily add the appropriate information.

• **Modify colors, fonts, and sizes of graph features:** You can adjust many parameters simply by double-clicking on specific parts of the graph. For example, to change the color, size, or shape of the data points, double click on the data series within the graph and you will get a window that lets you change those parameters. To change the scale of either axis, double click on the line of the x- or y-axis and you can change the minimum, maximum, major/minor units, etc. To change the font, size, or orientation of the labels double click on the text itself.

Excel will also determine a best-fit line of your data and the line’s slope. After you have completed your graph, right-click on any data point, and select “Add Trendline.” To format the trendline, under “Type,” choose “Linear”. Under “Options,” click the boxes for “Display equation on chart”, and “Display R-squared value on chart”. Click the "OK" button. The equation for your trendline (in \( y = mx + b \) format where \( m \) = slope and \( b \) = the value of the y-intercept) and an \( R^2 \) values will then appear on your graph. Briefly, the closer your \( R^2 \) value is to 1.0, the better your experimental data fit the trendline, your best fit line.
Experiment 1 • Considerations

Using Microsoft Excel as according to the instructions in the previous pages, enter data from all the wells into a spreadsheet, and have Excel calculate averages and corrected averages. Construct a graph that visually portrays your data by plotting corrected average absorbance (y-axis) as a function of time (x-axis). Your graph should include five lines on one graph, one line for the averaged data from wells A1-3 on the 96 well plate, one with averaged data from B1-3, etc. Was there activity in wells A1 - D3? Was there enzyme activity in wells E1 - E3? Was activity the same in wells E1, E2, and E3?

Was activity linear for three minutes? If so, the slope of the best fit line tells you the rate of reaction. If not, explain your observations. The initial rate of a reaction may be determined from the slope of the line joining each successive point.

Do your data support your hypotheses? If not, how would you change the protocol?

Save your completed graph and data worksheet in an appropriate folder (such as Tuesday AM lab) with a descriptive name. Email the file to yourself, or place it in a shared folder on the public server) so that the raw data are preserved. Include your notes below on this page, and attach a copy of your graph (Graph 1) to the next page. Make sure all aspects of the graph are properly labeled within Excel, and make sure to include an appropriately descriptive title that distinguishes this graph from others by its content (not experiment number)
Attach Graph 1 to this page:
Experiment 2: Effects of varying enzyme concentration on IDH activity

**Problem:** What is the relationship between the rate of a reaction and the amount of enzyme in the assay solution when substrates are abundant (non-limiting)? This question might become “In subsequent experiments, how much enzyme solution should I use in each assay?”

**Hypothesis:** IDH activity will vary directly with the amount of enzyme in each assay.

To test this hypothesis, you will need to follow a protocol that holds all conditions constant except the amount of enzyme added to each assay. All tests should be run more than once; routinely, enzyme assays are run “in triplicate.” For example, wells B4, B5, and B6 in the table below are triplicate assays containing 5 µL of IDH. Set up reactions as shown the table below.

Your instructor will tell you (or write on the board) that today’s IDH stock concentration today is: __________________________. You will need to know this stock concentration for later calculations.

<table>
<thead>
<tr>
<th>Wells</th>
<th>Buffer</th>
<th>NADP⁺</th>
<th>IDH</th>
<th>Isocitrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>A 4-6</td>
<td>180 µL</td>
<td>10 µL</td>
<td>0 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>B 4-6</td>
<td>175 µL</td>
<td>10 µL</td>
<td>5 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>C 4-6</td>
<td>170 µL</td>
<td>10 µL</td>
<td>10 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>D 4-6</td>
<td>160 µL</td>
<td>10 µL</td>
<td>20 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>E 4-6</td>
<td>130 µL</td>
<td>10 µL</td>
<td>50 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>F 4-6</td>
<td>100 µL</td>
<td>10 µL</td>
<td>80 µL</td>
<td>10 µL</td>
</tr>
<tr>
<td>G 4-6</td>
<td>200 µL</td>
<td>--</td>
<td>--</td>
<td>--</td>
</tr>
</tbody>
</table>

**Procedure:**

1. Use the appropriate pipettors to add appropriate amount of buffer, NADP⁺, and IDH to the appropriate wells.

2. Again use the “IDH (340 nm every 30 sec)” protocol on KCjunior to collect your data. Make sure to add the isocitrate at the plate reader, just before you start collecting your data.

3. When the plate reader has finished all seven readings, print your results.

4. Return to your station and organize your data in Excel in a similar manner as you did in Experiment 1. Type in the raw triplicate data. Use Excel to calculate the averages absorbance values and corrected average absorbance values (with blanks subtracted). Create a single graph in Excel that includes a separate line for each experimental condition. Make sure all aspects of the graph are properly labeled within Excel, and make sure to include an appropriately descriptive title. Attach your graph (**Graph 2A**) to the next page of your lab notebook. Save the raw data file in your email or public shared folder.
Attach Graph 2A to this page:
**Experiment 2 • Considerations**

Look at the graph you just created to answer the following questions:

Was there activity in all wells?

Did activity vary with the amount of enzyme in each assay?

How consistent was the activity in the three wells with the same amount of enzyme?

Was activity linear for the first three minutes for each volume of enzyme? If not, explain.

Do your data support your hypothesis?

Return to Excel and use its ability to generate a formula for the best fit line for each of the five lines on the graph you just made. For each best fit line you will get an equation in the $y = mx + b$ format where $b$ is the $y$ intercept value and $m$ is the slope or change in absorbance over time which is the definition of activity. Put the slopes in the table below.

<table>
<thead>
<tr>
<th>IDH Volume/Well</th>
<th>IDH Concentration*</th>
<th>Slope (Abs/min) (a.k.a. activity)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 µL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5 µL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10 µL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20 µL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>50 µL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>80 µL</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*You will need to calculate the IDH concentration by multiplying the stock concentration by the proportion of IDH in the wells. (Hint: $C_sV_s = C_wV_w$)

Use the table above to construct another graph (Graph 2B) in Excel that compares the concentration of enzyme on the x-axis to the slope of the appropriate lines (slope indicates enzyme activity) from your previous graph. What conclusions can you reach from your results? Attach Graph 2B to the next page. Make sure all aspects of the graph are properly labeled within Excel, and make sure to include an appropriately descriptive title.

Why is it inappropriate to graph enzyme activity versus IDH volume/well for these data? Why is it far more appropriate to graph enzyme activity versus IDH concentration?
Attach a copy of Graph 2B to this page:
Experiment 3. Effects of varying isocitrate concentration

Problem: What is the relationship between the rate of a reaction and the amount of isocitrate in the assay solution when the amounts of IDH and NADP+ in the assay are held constant? Before you start this experiment, develop an hypothesis and sketch a graph predicting the relationship of activity vs. isocitrate concentration.

Write your hypothesis here:

Sketch a simple graph predicting the relationship between activity (y-axis) and isocitrate concentration (x-axis) here:

Procedure: To test your hypothesis, you will need to follow a protocol that holds all conditions constant except the amount of isocitrate added to each assay. The table below outlines such a protocol using five concentrations of isocitrate. Each concentration is tested in triplicate. Add reagents to your wells as listed from left to right.

*Important Note: in this experiment you will add the enzyme (IDH) last.

<table>
<thead>
<tr>
<th>Wells</th>
<th>Buffer</th>
<th>NADP+</th>
<th>Isocitrate</th>
<th>*IDH</th>
</tr>
</thead>
<tbody>
<tr>
<td>A 7-9</td>
<td>170 μL</td>
<td>10 μL</td>
<td>0</td>
<td>20 μL</td>
</tr>
<tr>
<td>B 7-9</td>
<td>167.5 μL</td>
<td>10 μL</td>
<td>2.5 μL</td>
<td>20 μL</td>
</tr>
<tr>
<td>C 7-9</td>
<td>165 μL</td>
<td>10 μL</td>
<td>5 μL</td>
<td>20 μL</td>
</tr>
<tr>
<td>D 7-9</td>
<td>160 μL</td>
<td>10 μL</td>
<td>10 μL</td>
<td>20 μL</td>
</tr>
<tr>
<td>E 7-9</td>
<td>150 μL</td>
<td>10 μL</td>
<td>20 μL</td>
<td>20 μL</td>
</tr>
<tr>
<td>F 7-9</td>
<td>130 μL</td>
<td>10 μL</td>
<td>40 μL</td>
<td>20 μL</td>
</tr>
<tr>
<td>G 7-9</td>
<td>90 μL</td>
<td>10 μL</td>
<td>80 μL</td>
<td>20 μL</td>
</tr>
<tr>
<td>H 7-9</td>
<td>200 μL</td>
<td>--</td>
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</tbody>
</table>

1. Use the appropriate pipettors to add appropriate amount of buffer, NADP+, and isocitrate to the appropriate wells (NOTE – this experiment adds IDH at the plate reader instead of isocitrate.)

2. Again use the “IDH (340 nm every 30 sec)” program to collect your data. Make sure to add the IDH at the plate reader as simultaneously as possible and just before you collect your data.

3. When the plate reader has finished all seven readings, print your results.
4. Return to your station and organize your data in Excel in a similar manner as you did in Experiments 1 and 2. Type in the raw triplicate data. Use Excel to calculate the average absorbance values and corrected average absorbance values (with blanks subtracted). Create a single graph in Excel that includes a separate line for each experimental condition. Make sure all aspects of the graph are properly labeled within Excel. Attach a copy of your graph (Graph 3A) to the next page of your lab notebook. Save the raw data file in your email or public shared folder.

5. Determine the activity for each concentration of isocitrate by determining the slopes of the lines in the graph you just made above. Then, use the table below to construct another graph that compares activity (y-axis) as a function of isocitrate concentration (x-axis). Attach that graph (Graph 3B) to the next available page of your lab notebook.

<table>
<thead>
<tr>
<th>Isocitrate Used</th>
<th>Isocitrate Concentration*</th>
<th>Slope (Abs/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>none</td>
<td>0 mM</td>
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<tr>
<td>2.5 µL</td>
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<td>20 µL</td>
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<td>40 µL</td>
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<tr>
<td>80 µL</td>
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</tbody>
</table>

*You will need to calculate the isocitrate concentration by multiplying the stock concentration by the proportion of isocitrate in the wells. (Hints: The isocitrate stock concentration is 4.6 mM; and \( C_{V_s} = C_w V_w \))

Do your data support your hypothesis?

Is the relationship between activity and concentration of substrate linear?
Attach a copy of Graph 3A to this page:
Attach a copy of Graph 3B to this page:
Preparation for Next Week’s Lab:

During next week’s lab, we will study the effects of environmental conditions on enzyme activity. Each group of students will design an experimental protocol to address one of the following questions. Hint: look at the following few pages of the lab manual to get a preview of these experiments. Your team will need to sign up for one of the following options by the end of lab today.

Option A: Does pH influence IDH activity?
Option B: Does IDH have a metal ion cofactor requirement?
Option C: Does temperature influence IDH activity?
Option D: Does NADP+ concentration influence IDH activity?
Option E: Does NaCl concentration influence IDH activity?
Option F: Does IDH activity vary among different organisms and/or tissues?

Now that you have selected a variable to test next week, you will need to think about the following things. (Reading forward to next week’s lab notes will be helpful.)

1. Develop a hypothesis about the effects of the environmental condition you chose to vary upon enzyme activity.

2. Design an experiment to test that hypothesis.

3. Prepare a protocol to carry out that experiment. Write your protocol in the space below.
IDH Physical Models

✓ **Special Note**
When handling the IDH models please be very careful – they are fragile, one-of-a-kind objects that cannot be easily (or inexpensively) replaced.

**In lab today your group must:**

**A.** Find on the space-filling and the wire models of IDH:
- the active site
- isocitrate
- Mg$^{2+}$
- NADP$^+$ and/or NAD$^+$
- an alpha helix
- a beta sheet

**B.** Have one person randomly and gently tumble the two models and have another person move both models so they are sitting in the same orientation.

**C.** Answer the question: Why does IDH prefer NADP$^+$ over NAD$^+$ given they have the same structure on the reactive end of the molecule?
Parameters that Affect Enzyme Activity

Focused Reading

- p 158-161 “Molecular Structure...” to “How are Enzyme...”
- p 161-162 “Enzymes can be...” to “Allosteric enzymes...”
- p 164-165 “Enzymes are affected...” to “Chapter summary...”

✓ Special Note
- Bring a calculator to lab

The more you have prepared before lab the smoother it will proceed. This includes setting up spreadsheets and performing calculations of concentrations of test substances.

Goals for This Lab:

This week, we will determine the effects of environmental perturbations of our standard IDH assay conditions. We will use the techniques and information we learned last week and apply that to this week’s experiments.

Methods and Materials

You will use the same general methods that we used in the previous lab. All equipment, solutions and supplies required to carry out the experiments have been prepared for you. You should review your protocol and assign specific tasks to specific team members before you start your experiments. You will find helpful tables and suggestions for the different experimental options on the following pages. Find the specific page that relates to the experiment that your group chose to perform today.

The stock solutions provided for all groups are:

2.88 mM NADP+

4.6 mM isocitrate

_____ IDH (look on the board for the IDH concentration)

If you are using other reagent(s) for your experiment the concentration(s) will be provided on the reagent itself and/or in the notes.

Before You Leave Lab (All Groups):

1. Be certain that you have collected all of the data you need to make your experiment complete.

2. Be certain that each member of the group fully understands what was done and has a copy of all of your data recorded in her/his lab manual.

3. Be certain that you have analyzed your results and determine if your results support your hypothesis.
Option A: Does pH influence IDH activity?

Write out your hypothesis in this space:

To determine if pH influences IDH activity, we will need to follow a protocol that holds all conditions constant except the pH of the assay buffer. Remember to include blank(s).

<table>
<thead>
<tr>
<th>Wells</th>
<th>pH</th>
<th>Buffer</th>
<th>NADP⁺</th>
<th>IDH</th>
<th>Isocitrate</th>
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Considerations:
Does IDH activity vary when the pH of the assay mixture varies, or do levels of activity remain constant regardless of pH? Explain how the pH of the assay mixture might affect activity of an enzyme.

Make an Excel file of your data, formatted in the same way as last week. Attach a graph of your data below.
Option B: Does IDH Have a Metal Ion Requirement?

Write out your hypothesis in this space:

You will need to follow a protocol that holds all conditions constant except for the presence or absence of specific divalent metal ions. Your stock ion solutions were all prepared at 20 mM and your EDTA solution is 0.1 M. (EDTA is a chelator that removes ions from solutions.)

<table>
<thead>
<tr>
<th>Wells</th>
<th>Buffer **</th>
<th>Metal</th>
<th>NADP⁺</th>
<th>IDH</th>
<th>Isocitrate</th>
</tr>
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<tbody>
<tr>
<td>None</td>
<td>EDTA</td>
<td>Mg²⁺</td>
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<td>10 µL</td>
<td>EDTA</td>
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<tr>
<td>10 µL</td>
<td>Mg²⁺</td>
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<td>10 µL</td>
<td>Mn²⁺</td>
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<td>10 µL</td>
<td>Ca²⁺</td>
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<tr>
<td>10 µL</td>
<td>Zn²⁺</td>
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<td>Blank</td>
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**You will use a special buffer that does not contain Mg²⁺. Do not share solutions with other lab groups.

Considerations:
Does IDH require a divalent metal ion for activity? Does additional Mg²⁺ added to the standard assay buffer increase activity? What does this observation mean? Does the addition of Mn²⁺ added to the standard assay buffer increase activity? What does this observation mean? How can the effects of Ca²⁺ and Zn²⁺ on IDH activity be explained?

Make an Excel file of your data, formatted in the same way as last week. Attach a graph of your data below.
Option C: How does temperature influence IDH?

Write out your hypothesis in this space:

You will need to follow a protocol that holds all conditions constant except temperature. The temperature on the plate reader can be set for a variety of temperatures above ambient (room) temperature. Remember to make a blank.

<table>
<thead>
<tr>
<th>Wells</th>
<th>Temperature</th>
<th>Buffer</th>
<th>NADP⁺</th>
<th>IDH</th>
<th>Isocitrate</th>
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Make an Excel file of your data, formatted in the same way as last week. Attach a graph of your data below.
**Option D: Does enzyme activity vary with NADP⁺ concentration?**

**Write out your hypothesis in this space:**

What is the relationship between the rate of a reaction and the amount of NADP⁺ in the assay solution when the amount of enzyme is held constant? Before starting this experiment, develop a hypothesis and sketch a graph predicting the relationship of activity vs. NADP⁺ concentration.

<table>
<thead>
<tr>
<th>Wells</th>
<th>Buffer</th>
<th>NADP⁺</th>
<th>IDH</th>
<th>Isocitrate</th>
<th>*NADP⁺ concentration</th>
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<tbody>
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<td></td>
<td></td>
<td>0</td>
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*Note: 2.88 mM NADP⁺ stock provided.

**Considerations:**
Organize the data from the experiment. Determine the mean activity for each concentration of NADP⁺. Construct a graph that compares activity as a function of NADP⁺ concentration. Do the data support the hypothesis? Is the relationship between activity and concentration of NADP⁺ linear? Explain this relationship.

**Make an Excel file of your data, formatted in the same way as last week. Attach a graph of your data below.**
Option E: Does NaCl concentration influence IDH activity?

Write out your hypothesis in this space:

To test this hypothesis, you will need to follow a protocol that holds all conditions constant except concentration of NaCl in the assay solution. You will be given a 5 M NaCl solution to dilute to a wide variety of different concentrations. FYI – blood is ~0.9% salt and the FW of NaCl is 58.44. Remember to set up appropriate blanks.

<table>
<thead>
<tr>
<th>Wells</th>
<th>Buffer</th>
<th>5M NaCl</th>
<th>NaCl Concentration</th>
<th>NADP+</th>
<th>IDH</th>
<th>Isocitrate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0</td>
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Considerations:

Were differences in activity between the treatments observed? What was the relationship between the concentration of NaCl and activity? Explain how salt might affect enzyme activity.

Make an Excel file of your data, formatted in the same way as last week. Attach a graph of your data below.
Option F: Is IDH present in specific organisms and/or tissues?

Write out your hypothesis in this space:

This experiment is “open ended”; we may design a single, additional experiment, comparable to the ones listed above, or expand these topics into a research project of wider magnitude.

1. You may choose to survey IDH activity:
   i. in a wide variety of related species.
   ii. in different tissues of a single species.
2. Homogenize the samples in cold TEM Buffer, using a kitchen blender.
3. Filter the homogenate through two layers of cheesecloth into a small beaker on ice.
4. Transfer 1 mL samples to 1.5 mL microfuge tubes, spin for five minutes.
5. Transfer the supernatant to clean 1.5 mL microfuge tubes, on ice.
6. Use standard conditions to assay IDH activity.

<table>
<thead>
<tr>
<th>Wells</th>
<th>Buffer</th>
<th>Tissue</th>
<th>NADP⁺</th>
<th>IDH</th>
<th>Isocitrate</th>
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</thead>
<tbody>
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Make an Excel file of your data, formatted in the same way as last week. Attach a graph of your data below.
Lab Notes
Graphing Exercise: Introduction to Data Presentation

The enzyme wildcatase catalyzes the conversion of wildcatin to catin. Previously, researchers determined that catin absorbs maximally at 605 nm, while wildcatin has a minimal absorbance at this wavelength. To characterize the properties of wildcatase, you have conducted a series of experiments. The scientific community is extremely excited to hear about your results. In fact, the organizers of the 16th Annual International Wildcatase Convention ask you to present your findings at their meeting in Cancun. Normally, you would decline this invitation, preferring to stay in Davidson. Because your friend is one of the meeting organizers, though, you agree to attend and present your findings. Of course, now you must organize your data and make graphs and/or tables of your results for your presentation.

Brief descriptions of these experiments and the resulting data are provided on subsequent pages. For each experiment, determine the most effective graphical means of presenting your data and generate a graph in Excel that you can include in your lab notebook on the following pages.
Experiment 1: After a series of experiments, you have concluded that two different enzymes can catalyze the conversion of wildcatin to catin. You have named these enzymes wildcatase A and wildcatase B. A study of 100 mammal species has revealed that 50 of these species contain wildcatase A and wildcatase B, 40 contain only wildcatase A, and the remaining 10 contain only wildcatase B.

Attach a copy of your graph on this page:
Experiment 2: Your studies indicate that both deer mice and field mice contain only wildcatase A. To investigate the location of wildcatase A within these organisms, you looked for the presence of this enzyme in various tissues. You discovered that, in the deer mouse, wildcatase A is present in lung, brain, spleen, and liver. It is not found in muscle, kidney, or heart tissues. In the field mouse, wildcatase is present in lung, brain, and spleen tissues. It is not found in liver, muscle, kidney, or heart tissues.

Attach a copy of your graph on this page:
Experiment 3: In examining the tissues of the deer mouse, you discovered that the different tissues containing wildcatase A contain different amounts of wildcatase A. Lung contains 350 units of wildcatase A per gram of tissue. Spleen contains 250 units of wildcatase per gram of tissue. Liver contains 100 units of wildcatase A per gram of tissue. Brain contains 500 units of wildcatase A per gram of tissue.

Attach a copy of your graph on this page:
**Experiment 4:** To investigate the kinetics of wildcatase A and wildcatase B activity, you combined 100 mM wildcatase A and 100 mM wildcatin. You then measured the absorbance at 605 nm at 30 second intervals for two minutes, resulting in the following data: 0 min: 0.010; 0.5 min: 0.100; 1.0 min: 0.20; 1.5 min: 0.300; 2.0 min: 0.400. You then did a similar experiment, but used wildcatase B instead, resulting in the following data: 0 min: 0.010; 0.5 min: 0.150; 1.0 min: 0.300; 1.5 min: 0.450; 2.0 min: 0.600.

**Attach a copy of your graph on this page:**
What is your genotype? A PCR Approach

Focused Reading
• p 279-283 “DNA polymerase…” to “Telomeres…”
• p 324-327 “Gel electrophoresis separates…” to “15.2 Recap”

Web Reading
• Movie of PCR method: http://www.bio.davidson.edu/misc/movies/pcr2.mov
• Polymerase Chain Reaction: http://www.dncal.org/resources/animations/pcr.html
• Gel electrophoresis: http://learn.genetics.utah.edu/content/labs/gel/

Introduction
Movies, TV crime series and the news are full of court cases that employ “DNA fingerprinting”. The technology available is so sensitive that unbelievable sources of DNA have been used to convict criminals. In Minnesota, for example, DNA was extracted from the back of a postage stamp because some epidermal cells from a person’s tongue had been deposited on the glue when the stamp was licked. Most likely, a pivotal point of contention in future court cases will be the collection and handling of the evidence. For example, what if some DNA from the crime scene is proposed to have come from the accused? The defense attorney could suggest that the police collected some epidermal cells from the sidewalk (from a visit the day before) at the same time as the blood drops. What do you think of this possibility as a defense?

There are three standard methods for “DNA fingerprinting”: 1) Southern blots (which are being used less and less), 2) PCR, and 3) a combination of PCR and DNA sequencing. Over the next two weeks, we will use PCR to determine the genotype of every student in class. Dr. Kary Mullis, the inventor of PCR, was awarded a Nobel Prize in 1993 for his revolutionary innovation. PCR allows you to amplify a single small region of DNA (out of a complex background mixture) into millions of copies. One requirement is that the ends of the DNA of interest must have been previously sequenced, because you have to supply primers that will specifically hybridize near the target gene. We are using hair roots as our source of genomic DNA and are looking at a locus called DIS80. DIS80 contains a Variable Number of Tandem Repeat sequence (VNTR). As the term implies, there is a section of DNA that is repeated to varying degrees in each person. As a simplified example, let’s say the repeat unit is the two nucleotides sequence CG. If we were to sequence this portion of the DIS80 locus from four different DNA sources, we might see the following:

1) ATGCCGTATTACGCCGCGCGCGCGCGCTATTAGGTATTAG
2) ATGCCGTATTACGCCGCGCGCGCGCGCGCGCGCGCGCTATTAGGTATTAG
3) ATGCCGTATTACGCCGCGCGCGCGCTATTAGGTATTAG
4) ATGCCGTATTACGCCGCGCGCGCGCGCGCGCGCGCGCTATTAGGTATTAG
In this example, there are four alleles of this VNTR with four different lengths. If we electrophoresed these four segments of DNA on a gel, we would observe bands of four different sizes (2>4>1>3). In a criminal case, we might have four suspects and one DNA sample from the crime scene. The resulting gel might look like this:

Questions:
1) So, “who done it?”
2) What is wrong with the above gel? Why is this example too easy?

Now it is time for us to determine our genotypes. You should be forewarned – this technique is a delicate procedure that does not always work for everyone. In order to process this kind of evidence for a criminal case, a technician usually has a master’s degree in Forensic Science, and a few years of on-the-job training. Nevertheless, even these experts sometimes make mistakes. So, do not be discouraged if your sample does not work, but try to avoid this situation by observing these guidelines:

• Follow the protocol as carefully as possible.

• Do not contaminate your hair or DNA with that of others (remember one cell contains enough DNA to be amplified).

• Immediately after the DNA extraction is finished, visually check to verify that you have extracted DNA by gently removing the tube from the thermocycler and flicking the tube holding it up to a light and looking very carefully. You might be able to see a denser region of the solution at the bottom of the tube as it mixes with the less dense water.

• The most common mistakes are pipetting errors. Be sure to check that you are transferring the correct volumes and always use clean pipette tips; when in doubt use a new tip.

• Be very careful loading the gel. We will have time to practice loading gels this week so that when you are loading your real sample next week, you will be a pro.

**DNA extraction**

1) Pluck a hair so that a large portion of root is removed from your head (yikes!) The root can be anywhere from dark to translucent depending on your individual heritage and biology. Regardless of the color, the root will be sticky so you can test it by touching it to the bench top to see if it adheres. Check to make sure you got some root and not all shaft.

2) Put the hair into a small microfuge tube with the root at the bottom of the tube. Cut off most of the hair but keep the root (~5 mm). Be careful, sometimes the root will jump away when you cut the hair.
3) Incubate the root in 100 µL digestion buffer (which contains 6 µg of proteinase K) for 1 hour at 55°C, then 10 minutes at 95°C (what is the purpose of this step?). Use thermocycler program HAIR 1 - lid disabled.

During this waiting period, we will practice loading gels so you will be ready for next week.

**PCR Reaction Mixtures**

4) When the DNA extraction cools, **vortex the tubes for 30 seconds** and then set up a new 500 µL microfuge tube by adding the following: 10.0 µL of your DNA and 15 µL of reaction mixture. The reaction mixture contains H₂O, PCR buffer, dimethylsulfoxide (DMSO; a solvent), dNTPs (dATP, dTTP, dCTP, & dGTP), the two primers, Taq DNA polymerase, and gel loading dye. (The Taq is added immediately before distribution by your instructor, 0.4 µL Taq per 10 µL mixture.)

**PCR**

The PCR temperature conditions are as follows:

- **Step 1:** 5 minutes at 95˚C
- **Step 2:** 1 minute at 95˚C
- **Step 3:** 1 minute at 65˚C
- **Step 4:** 1 minute at 72˚C
- **Step 5:** repeat steps 2 - 4 twenty-nine more times
- **Step 6:** hold at 4˚C

When the PCR is completed, the tubes are removed and stored at 4˚C.

**D1S80 factoids**

- >80% of all people tested are heterozygous
- 28 alleles have been published
- Repeat unit is 16 nucleotides long
- If there were zero repeat units, the PCR product would be 142 bp long
- PCR products range from 430 to 814 base pairs long
- 44 repeated units have been observed in the largest allele
- Primer sequences²:
  - #1 5' GAAACTGGCCTCACAACACTGCCCACC 3'
  - #2 5' GTCTTGTGAGAGATGCACGTGCCCCTTGC 3'

**The PCR Results**

Load 15 µl of your DNA/dye mixture into a lane on the gel. It is important to note which lane contains your DNA. We will also add DNA of known sizes to several lanes on the gel. This DNA often called a “ladder” will provide a ruler of sorts for us to determine the size of your DNA. Electrophorese the DNA on a 1.5% agarose gel using 0.5X TBE and 3.6 µl of Midori Green. We usually run these gels at 90-100 volts for 1 - 1.5 hours. The exact time and voltage will depend on the gel box configuration. Appropriate conditions can be refined accordingly.

While the gel is running, we will learn how to calculate the molecular weights of bands.
**How to calculate the molecular weight (MW) of a molecule that has been separated in a gel**

The log\(_{10}\) of a molecule’s molecular weight is proportional to the distance that molecule has migrated. Therefore, the first step is to generate a **standard curve** using molecules of known size (the molecular weight markers).

You will do this once, for practice, using the image at right, and then again using your actual data. You must generate a new standard curve for your actual data since your gel was run at a different time and under different conditions. The molecular weights on your actual gel are determined by comparing the pattern of the bands on your gel picture to the picture on the right (e.g. three close together, then one bright one then another bright one then two close together). We are using the same molecular weight standard samples as seen here, so you can use the banding pattern to figure out the molecular weight.

Set up an Excel spreadsheet with columns for distance migrated (in millimeters, mm) and molecular weight (in kilobases= kb).

Measure from the well to each band (bright line) on the picture. These are the distances migrated for each DNA fragment.

You will now plot molecular weight (kb) on the Y axis and the distance the molecule migrated (mm) on the X-axis. After you have entered your values, select all the values by clicking and dragging over them.
Click on the ‘charts’ tab and the scatter pull down menu. Choose ‘marked scatter’ from the that menu (on older versions it is not named but has a picture of non-connected dots on a graph).

A new scatter plot should appear. Double click on the Y axis and choose the ‘scale’ option on the left. Select the ‘logarithmic scale’ button on the right and click ok.

The plot will change in appearance as the Y axis reconfigures itself to a log scale.

What we want from these data is the relationship between MW and distance migrated—a standard curve. This is the second time in our labs we have generated a standard curve to aid in data analysis.

Click on one of the dots on your graph so that the whole set of dots becomes selected. From the Chart menu at the top of the screen select ‘Add Trendline’. In the dialog box, select ‘Types’ and ‘Exponential’. In the same dialog box select ‘options’ on the left. Then select ‘Display equation on chart’ and ‘Display R-squared on chart’. Then click on ‘OK’.

Now that you have the equation of the line that best describes the relationship between the MW of the standards (the ‘known’ MWs) and the distances those DNA fragments migrated, you can use it to determine the MW of your DNA band (the ‘unknown’). Use the distance that your band migrated as the X value and solve the equation to find out what the corresponding Y is. This value is the number of kilobases of DNA that your D1S80 DNA fragment contains. If you have two bands, repeat this calculation with the distance that the other band migrated.

Footnotes:
Lab Notes (Including Picture of YOUR DNA)
**VNTR Online Exploration**


2. Make sure "All Databases" is selected in the pull-down menu at the top.

3. Enter D1S80 in the search box and click Search. (Note that it is the numeral one, not a letter, in the second position.)

4. List the databases that have matches to D1S80. (In other words, which databases show a number instead of "none" to the left of the name?) What kind of information do you expect to find in each (e.g., nucleotide sequences, protein sequences, journal articles, etc.)?

5. Click on the number in the box next to the Nucleotide sequence database (45 as of July 2015).

6. Look at the description of each entry. (Make sure to view all 45 entries.) How many species are described to have this VNTR? How many alleles are there in the database? Give the accession number of the largest (highest number of repeat units) and smallest VNTR alleles.

7. Click on any of the human entries that specify the number of repeat units; scroll to the bottom of the page and copy the DNA sequence into a Word document.

8. Can you identify a typical repeat unit? Hint: look at the FEATURES part of the database entry you opened; it identifies the region of the sequence in which repeats are found. See “D1S80 Factoids” on Lab Manual page 51 to remind yourself of the repeat size. Keep in mind that the repeats may not be completely identical, since mutations have gradually accumulated over time. Bold the first repeat unit (the 5’-most one) and the last repeat unit (the 3’-most one) in your Word document. Are all repeat units identical? If they are not, will that impair your ability to estimate the number of repeat units you have?

9. On the sequence in your Word document, underline the sequences that represent the primers you are using for the PCR reaction. What size fragment do you expect to see on the gel, if this were the sequence of one of your two alleles?

10. Put square brackets around the nucleotides that are not primers and are not considered a part of the repeats (note, they may be at more than one place in the sequence). How many of them are there? Your text says that there are 142 extra base pairs (bp) that you need to subtract from the total size of the PCR product to get the size of the repeat region. What accounts for this total?
Using Microscopes

Focused Reading
• p 79 “Microscopes reveal…” to “The plasma membrane…”
• p 78 fig. 5.1 (The Scale of Life)
• p 80 fig. 5.3 (Looking at Cells)

Web Reading
• How to Use Your Compound Microscope Properly
  www.bio.davidson.edu/Courses/Bio111/Bio111LabMan/IL/scopes.html
• Molecular Expressions Optical Microscopy Primer
  (optional — a great place to learn about microscopes, try interactive animations, and see cool images)
  http://microscopy.fsu.edu/primer/index.html

✓ Special Note
• You will be working with iodine in this lab and there is a risk of staining your clothing, so you may want to wear old clothes.

Goals for This Exercise
During this session, you will learn how to use a compound microscope that has the ability to view specimens in brightfield, darkfield, and phase-contrast illumination. You also will learn about a model research organism, Chlamydomonas. Chlamydomonas is a unicellular green alga that has two flagella and can reproduce asexually by mitosis, or sexually after undergoing gametogenesis.

Care and Use of the Compound Microscopes
A compound microscope is illustrated in Figure 1 and can magnify from 40 to 2000 times (40 – 2000X). Microscope quality, however, depends on resolving power in addition to magnification. Resolving power is the ability to distinguish between two points in the field of view. Thus, if you can magnify 1000-fold yet cannot resolve detail, then your microscope would be of little value. Even more important may be the abilities of the microscopist to learn the capabilities of her/his microscope and to gain proficiency in the use of the instrument.

Important things to know when using a microscope:
1. Always carry a microscope with both hands, one grasping the handhold in the back and one grasping the bottom.
2. Do not swing the microscope and do not bang it onto the bench top.
3. Never place the microscope near the edge of the bench and keep electrical cords out of the way.
4. All of our compound microscopes are parfocal, which means that the objects remain in focus as you change from one objective lens to another. Examine your material first using the lower power objective (i.e., 10X); then use a higher power objective (i.e., 20X or 40X). Because the objectives are parfocal, you need to use only the fine focus knob to fine-tune your image. Never use the coarse adjustment to focus downward. Replace and remove a slide only after the lowest power objective has been rotated into viewing position.
5. Never attempt to repair a microscope or force an adjustment knob. You may severely damage the instrument.

Parts of a Microscope:
Ocular: The piece you look through. Sometimes called an ocular lens or eyepiece, this unit is really a series of lenses. Our microscopes are binocular, having two oculars. Learn to use both eyes; focus your
eyes as if you were looking at an object about five to ten meters in front of you. You should adjust the width of the oculars to match the width of your eyes, similar to how you might adjust binoculars when birdwatching.

**Objective lens:** Sometimes called the objective; a set of self-contained lenses. The objective gathers light and directs it through the tube to the oculars. These microscopes have three phase contrast objectives (10X, 20X, and 40X with red lines on them) and one brightfield objective (20X with no red line).

**Nosepiece:** The rotating turret to which objectives are mounted. There are preset positions for each objective, detected by slight pressure changes while turning the nosepiece and usually a clicking noise. You should not grab the objectives to turn the nosepiece – use the black ring instead.

**Stage:** The flat surface upon which slides are placed. On your microscopes, the stage moves up and down and the slide is manipulated by a geared device. A moveable stage is sometimes called a mechanical stage. The slide is moved left/right and front/back by two knobs projecting downward from the stage.

**Condenser:** A lens system under the stage that gathers light from the light source and focuses it on the
specimen. There is a diaphragm in one part of the condenser that can be adjusted to allow the viewer to see different parts of the cell when using bright field illumination. You should experiment with this control. These condensers also have phase rings but you should not have to make any adjustments to them.

Condenser Adjustment Control: Under the stage on the left side is a small knob that is used to adjust the height of the condenser. Usually, the condenser always will be all the way up.

Light Switch Control: The light switch and intensity controls are on the right side of the microscope base, about half way up the side. There is an on/off switch as well as a brightness control. Use only as much light as necessary to illuminate the specimen.

Light Source: On our microscope the light source is built into the base and is directly under the condenser.

Adjustment (Focus) Knobs: Both coarse (large) and fine (small, inner) adjustment knobs are found on both sides of our microscopes. Remember that the coarse adjustment is used only with the low-power objective. These knobs control a gear mechanism that raises and lowers the stage.

Types of Microscopy
There are three different ways that we can view specimens with these microscopes. The type of illumination with which people are most familiar is called Brightfield. Think of the light source as producing a solid tube of light that travels up to and through the condenser. When you view specimens with all of this light, you are using bright field illumination.

Darkfield: Dark field illumination seems like an oxymoron, but in this case it describes an unusual way of viewing specimens in some compound microscopes. The light that passes directly through the condenser does not enter the objective lens. Only light that has been scattered or reflected by the specimen enters the objective. As a result, you wind up seeing bright objects on a dark background.

Phase-Contrast: Phase-contrast microscopy allows us to see otherwise transparent organelles and structures. We will make extensive use of this for viewing flagella. In a phase-contrast scope, the light hits the specimen and some of the light continues in a direct path. Other portions of the light pass through membranes that redirect the light. This redirected light is slowed down by 1/4 a wavelength (a phase shift of 1/4) by passing through a special filter. This special filter is shaped like a doughnut and is called a phase ring. The redirected and out of phase light eventually reaches your eyes but not at the same time as the unaltered light that passed straight through. The end result is that you can see transparent structures because they altered the pathway of light as it went through the structures. This phase shift allows us to view subcellular structures within living cells.

Viewing A Specimen:
Everyone will follow the procedure for viewing a specimen as a group. Your instructor will demonstrate how to make a wet mount (see below) and show you the differences between brightfield, darkfield, and phase-contrast microscopy using a microscope that is equipped with a camera and projector. In this session, we will be looking primarily at wet mounts. A "wet mount" is a specimen mounted in an aqueous solution but you do not expect to keep the slide for very long. If your preparation begins to dry out while you are working with it, make a new one.

Every time you work with a microscope:
1) Position the scope so it is directly in front of you and your chair is adjusted so that you do not have to strain to view a specimen.
2) Make sure the light intensity control is turned all the way off before turning on the microscope.
3) Make sure the 10X objective is in place over the specimen.
4) If you are making a wet mount, clean the microscope slide by fogging it with your breath and then wiping it with a Kimwipe.

**Bright Field**

1. Switch on the light source and then dial the adjustment knob to about 4.0. Start with the oculars set so they are at equal heights.
2. Turn the condenser so that the “O” is facing you. This position is the bright field slot on the condenser.
3. Position the low-power (10X) objective over the specimen and, looking from the side, raise the stage as high as possible. Notice how close to the objective the stage is.
4. Use the coarse adjustment to lower the stage away from the glass slide while looking through the oculars until the specimen comes into focus. Adjust the focus to its sharpest with the fine adjustment knob.
5. Now it is time to make sure both oculars are focused. Use the fine focus while looking through the right ocular and close your left eye. Pick one object to focus on. Then close your right eye and focus the left ocular by turning it up and down with the focusing ring for the left eye but do **not** touch the fine focus control during this time.
6. Readjust the light intensity to reduce glare and center the specimen in the field of view by moving the stage.
7. Use the knob on the left side of the condenser to move the condenser up as high as possible. You may also want to adjust the condenser’s diaphragm to maximize the resolution but minimize the “graininess” of the image.
8. Place the 20X objective (no red line) over the specimen and sharpen the focus with the fine adjustment knob (only) as necessary. Readjust light.
9. Adjust the condenser’s diaphragm to maximize the resolution of the structure you are trying to see. The actual setting will depend on what you are trying to see. Small translucent objects will be seen more easily with the diaphragm closed substantially while large pigmented structures are easier to see with the diaphragm wide open.
10. Repeat steps 8 and 9 but use the 40X objective instead of the 20X.

**Darkfield**

1. Turn the condenser ring clockwise so that the “D” is facing you. This position will permit you to see objects in dark field illumination. You also must adjust the condenser so that it is as high as it can go - use the knob on the left side of the scope. You can use dark field illumination with any of the four objective lenses.
2. What structures can you see now that you could not see in brightfield?
3. What is difficult to see in dark field that was easy to see in brightfield?

**Phase-Contrast**

1. When you use phase-contrast, you must match the objective lens with the phase ring in the condenser. Therefore, you must follow this table:

<table>
<thead>
<tr>
<th>Objective Lens</th>
<th>Phase Ring</th>
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<tr>
<td>10X, 20X (red lines)</td>
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<tr>
<td>40X (red line)</td>
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</table>

**A Reminder:** only the objectives with red lines can be used for phase. The 20X objective that
does not have a red line on it is not equipped with phase rings.

2. Select the appropriate objective lens and phase ring pair. You might need to increase the amount of light because images do not appear as bright in phase. When you have done this, you should adjust the condenser vertically with the knob on the left side of the condenser. When these adjustments are made, using a phase-contrast microscope is similar to using a bright field scope.

3. What structures can you see now that you could not see in brightfield? Darkfield?

4. What is difficult to see in phase-contrast that was easier to see in brightfield?

5. Do you see the same colors in phase that you saw in brightfield? Darkfield?

Lab Notes
Overview:
Over the next several weeks we will become comfortable with a fundamental tool in biology - the compound microscope and the use of digital cameras and digital imaging software to measure very small structures. We will conduct a series of experiments on a unicellular green alga, *Chlamydomonas reinhardtii*. *Chlamydomonas* is a biflagellated green plant that reproduces asexually (by mitosis) and sexually (via meiosis, mating, and zygote formation).

Background Information on *Chlamydomonas*
There are several reasons why *Chlamydomonas* is such a useful model organism. It is a haploid organism, which means there is only one copy of each chromosome. Therefore, the genotype is always expressed in the phenotype (unlike diploids that may have a recessive mutation that is not revealed in the phenotype). It has a generation time of two weeks (from mating of one generation to when the next generation can mate). Finally, there are hundreds of mutant strains (stored at Duke University) that have been generated over the years and can be used for research. For example: *ac-17* cannot fix carbon during photosynthesis, *arg-7* requires the amino acid arginine to be added to the medium because it cannot synthesize its own; *act-1* is resistant to the translational inhibiting drug cycloheximide; and *pf14* has straight and paralyzed flagella so it cannot swim.

Each person should:
A) Place 18 µL of *Chlamydomonas* on a clean (use a Kimwipe) glass microscope slide and gently cover with a coverslip. (If you press down on the coverslip you will crush the cells.) Place the slide on the stage of the microscope and use the 10X objective lens to observe the cells swimming around. Start with brightfield, then try darkfield and phase-contrast.
   1. Can you see the flagella?
   2. Which form of illumination allows you to see them the best?
   3. What is the total magnification you are using with a 10X objective lens and the 10X oculars?

B) Increase the magnification by using the 20X objective lens. Again, view the cells in brightfield, darkfield, and phase-contrast. Remember to use lenses with the red ring for phase and the 20X without the red ring for brightfield. Also view the cells at the highest magnification, either 40X or 60X.
   1. What is your total magnification now?
   2. Can you see the flagella? Which form of microscopy is the best for seeing flagella?
   3. Can you see any other organelles in these cells? (Try all three forms of illumination.)
   4. How can you see flagella better without staining them?
   5. What colors do you see in *Chlamydomonas* cells?
   6. What structures are responsible for the colors?
   7. How do you calculate the total magnification you are using on a microscope?

C) On a new slide (or on your original slide if there is space), place 10 µL of cells into 10 µL of Lugol's
Fixative. Lugol's fixative is a dye that stains the sugars that are covalently bound to the proteins (sugar-coated proteins are called glycoproteins) on the surface of the flagella. Cover with a coverslip. Examine this preparation of stained cells under the microscope. View the cells at all three magnifications with each form of illumination.

What structure(s) can you see better with fixed cells than with live, unstained cells? Give two possible reasons here:

**Goals for this Session:**

During this session, you will collect data on the regeneration of flagella on *Chlamydomonas*. These cells will be deflagellated, and you will immediately measure regrowing flagella over a 90-minute time period. You will learn to use TCapture software to capture images of the cells and ImageJ software to measure flagella within your images.

This set of experiments requires teamwork and efficiency. Assign tasks ahead of time (see next page) but make observations for yourself because some test questions may cover lab work. Be organized and ready to start immediately when the cells are delivered to you, because cells will already be starting to regenerate their flagella.

You will measure the length of the flagella as a function of time. At each time point, 50 µL of cells will be "fixed" with 50 µL of Lugol's fixative, an iodine-based substance that kills, stains, and immobilizes cells. Of that mixture, 18 µL will be placed on a slide. You will capture images of fixed cells and then measure the flagella on the image files. For each time point, you should measure the length of 15 flagella (total of 15 flagella on 15 cells).

**Prepare a spreadsheet to compile the following data (DON'T ENTER DATA HERE):**

*Chlamydomonas untreated with acid* (retaining their original flagella, to allow you to determine normal flagellar length. You must fix cells with Lugol's to immobilize and allow for visualization.)

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| **Deflagellated Chlamydomonas** (cells must stay alive until the specified time point, at which you treat with Lugol's to immobilize and allow for visualization.)

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Detailed Protocol:

A) About $5.0 \times 10^{12}$ *Chlamydomonas* cells will be deflagellated and delivered to you.

B) In a four person group:

- **Person #1** should fill microfuge tubes with 50 µL Lugol’s fixative, labeling with time points 0, 15, 30, etc. Then, for most of the time (with help from person #2) measure flagellar lengths on ImageJ, after images are received from persons #3 and #4. Record all the data and make sure that no time points are missed.
- **Person #2** should, at time 0 and every 15 minutes thereafter, add 50 µL of deflagellated *Chlamydomonas* to the appropriate microfuge tubes already containing Lugol’s fixative. Move the container of live cells back to the light shelf each time. Make a slide using 18 µL of each fixed sample; to avoid dried out slides, do not make the slides until persons #3 and #4 are ready to measure them. In your extra time, help person #1 measure flagellar lengths and record data.
- **Persons #3 & 4** should capture images of fixed cells on the computer using TCapture. For each slide, capture an image for each of the first 15 cells that you see whose full periphery is visible (i.e. not clumped). If flagella are visible, make sure that the longest flagellum for a given cell is in focus. Transmit the image files via flash drive or Bio111 shared folder (see instructions on the next page) to persons #1 and #2 so that they can make measurements.

C) In a three person group:

- **Person #1** should fill microfuge tubes with 50 µL Lugol’s fixative, labeling with time points 0, 15, 30, etc. Then, for most of the time, measure flagellar lengths on ImageJ after images are received from person #3. Record all the data and make sure that no time points are missed.
- **Person #2** should, at time 0 and every 15 minutes thereafter, add 50 µL of deflagellated *Chlamydomonas* to the appropriate microfuge tubes already containing Lugol’s fixative. Move the container of live cells back to the light shelf each time. Make a slide using 18 µL of each fixed sample; to avoid dried out slides, do not make the slides until person #3 is ready to measure them. In your extra time, either help person #1 with measurements or help person #3 with image capturing, depending on who is running behind.
- **Person #3** should capture images of fixed cells on the computer using TCapture. For each slide, capture an image for each of the first 15 cells that you see whose full periphery is visible (i.e. not clumped). If flagella are visible, make sure that the longest flagellum for a given cell is in focus. Transmit the image files via flash drive or shared folder (see instructions on the next page) to person #1 so that he/she can make measurements.

D) Enter **RAW DATA** into your Excel file. A good scientist keeps all data in her/his notebooks, not just the averaged results. It is always better to take too many lab notes than too few. In the real world, scientists are required to keep on hand (for future scrutiny) all raw data that underlie any averages published. Otherwise, the scientists could be accused of fraud.

E) When you are done, clean your area, turn off the microscopes, throw away any trash, and return equipment to where you found it. It is good lab etiquette to clean up and return things properly when many people share equipment and space.
Instructions for using BIO111 shared folders on the Louise Public server*
ONE computer in each group initially creates the folder. The OTHER computer in each group simply finds the folder once it is made.

(*Or you are welcome to use any other file sharing method your group selects such as DropBox or GoogleDrive.)

On Mac:
Open a Finder window, and at left under Shared, click Louise. Open folders in the following path: Louise>Public>Biology>111 Shared Files. Then find your group’s pre-existing folder (or create a new one with a unique name if no one in your group has done that yet).

To make an alias/shortcut for this folder, select the folder and hit command-L. Drag the alias icon to the desktop.

On Windows:
Open a window, and at left click Network. Open folders in the following path: Louise>Public>Biology>111 Shared Files. Then find your group’s pre-existing folder (or create a new one with a unique name if no one in your group has done that yet).

General Information about Chlamydomonas Flagella
As it so happens, Chlamydomonas is very sensitive to changes in its environment. To remove Chlamydomonas flagella we use the pH shock method. If we manipulate the pH of the growth medium by adding acetic acid until the pH decreases from about 7.2 to 4.5, the cells shed their flagella. Scientists have investigated why this event happens. We know that cells will not shed their flagella if there is no calcium in the growth medium. (Calcium can be removed from any solution by adding in a compound commonly referred to as EGTA. EGTA has a very high affinity for calcium and acts as a chelator, like a molecular sponge, to absorb ionically all the calcium, which means Chlamydomonas cells can not use or sense any calcium ions if EGTA is present.) Other researchers have shown that if one can experimentally elevate the level of calcium in the cytoplasm of Chlamydomonas cells, they shed their flagella. Hypothesize what is going on when Chlamydomonas cells shed their flagella when pH shocked in the presence of calcium. Can you devise an experiment to test your hypothesis?

By now you may be wondering why anyone would care about the dynamics of ‘pond scum’ flagellar regeneration. Chlamydomonas is a model organism for studying flagella and much of our understanding of cilia and flagella is due in large part to our understanding of Chlamydomonas flagella. As you read in your textbook, the flagella are comprised of many (~200) different proteins, but the predominant protein is tubulin. Each Chlamydomonas flagellum is built upon the 9+2 structure of microtubules (see the figure below). The outer nine microtubules are “doublets”, consisting of a complete circle of 13 protofilaments fused with a partial circle of 10 protofilaments. The two centrally located microtubules consist of a “singlet” of 13 protofilaments. Therefore, each flagellum contains (9x(13+10)) + (2x13) protofilaments. Each protofilament is composed of dimers of α tubulin and β tubulin. Each monomer of a globular tubulin molecule has a 4 nm (4 x 10^{-9} meters) diameter and is comprised of 450 amino acids. Therefore, the α/β dimer has a diameter of 8 nm and is made of 900 amino acids. Although there are multiple genes for tubulin, for the sake of simplicity let’s assume a single gene for each form of tubulin (one for α and one for β). These prototypical genes are about 1800 bases long. Remember that Chlamydomonas is haploid so one allele = one gene = one locus.
**Diagram of tubulin structure in flagella.** Compare with Figs 5.17 (page 96) & 5.20 (page 98) in your text.

**Study questions:**

1) Based on your results for flagella regeneration, calculate how many amino acids are being polymerized per minute into the tubulin component of the regenerating flagella.

2) Assume that all of the mRNA needed for this process is being synthesized **de novo**, from scratch. How many mRNA bases (assume no introns) must be transcribed per minute if every mRNA is translated only once? What if each mRNA is translated 100 times?

3) If RNA polymerase can travel no faster than 2500 bases per minute, is it possible for all of the RNA to be transcribed **de novo**? Explain your answer.

*Note: You will not be able to answer these questions with your experimentally determined rate because that rate will not be determined for another two weeks. However, make the assumption that flagella regenerate at 0.17 µm per minute. When you have determined the rate in your experiment, you should try the calculations again.*

**NEWS ITEM:** How can studying the flagella of pond scum be useful for understanding human health? A team of researchers found that mutant Chlamydomonas lacking the IFT88 gene appear normal except for the striking absence of flagella. IFT88, also called polaris, encodes for a protein involved in intraflagellar transport, the process of moving cargo up and down the microtubule tracks within flagella and cilia. Humans and mice have a gene called Tg737 that is orthologous to the Chlamydomonas IFT88 gene. Knockout mice, engineered to have defects in both copies of Tg737 die shortly after birth from polycystic kidney disease (PKD). Humans also die from PKD where kidneys enlarge due to overproliferation of kidney epithelial cells. Scientists now think that malfunctioning cilia on the side of the epithelial cells urine flows past bend when urine flows and this bending sends a signal to the epithelial cells cell that inhibits cyst formation. In patients with PCK, their kidney epithelial cells have abnormal cilia and their kidneys are filled with cysts that ultimately compromise the kidney’s ability to function. [HHMI Bulletin 18(2): 33]
**Measuring Chlamydomonas flagella lengths digitally**

**TCapture Instructions—for the ‘Taking Pictures’ computer**

BOOT COMPUTER IN WINDOWS. Make sure the camera is connected to the computer, and turn on both the microscope and the camera (the blue light on top will illuminate). Find the TCapture icon at the bottom of the viewing area. Double click this icon to launch the program. TCapture is the tool that allows you to capture images from the microscope.

Put the stage micrometer on the microscope stage. Use brightfield optics to view the micrometer with the 10X objective. Pull out the lever (sometimes called a stop) that sends light up to the digital camera. Find the object and focus the microscope. Now change the objective lens to the 40X or 60X power (depending on how your microscope is equipped) and refocus using only the fine focus knob. When the stage micrometer is centered in the field of view and in focus, you will see something similar to the image below. Notice that you can distinguish the long and short lines in this image. Make sure you can see the difference in the long and short lines in your image. Move the stage as needed.

Click the Capture button; in the dialog box, enter a file title, and select the destination folder for the file. You should use .tif images for your analysis. Do not include punctuation marks, underscores, or spaces in your file names.

If you have not already done so, figure out a file sharing method with your lab partners on the other computer. You can use the shared public folders on the College’s Louise drive (see box on p. 64) or else DropBox, Google Drive, OneDrive—whatever works for instantaneous file transfer to the other computer in your group. From here on out, the people taking pictures should save image files into the shared space, from which the people doing image analysis can directly retrieve the images for measuring.
At the same high magnification but with phase contrast optics, capture images of fixed Chlamydomonas cells each with a flagellum sharply in focus (from each time point, etc., as instructed elsewhere). For example, in the cropped image at right, the upper cell has both flagella in sharp focus, while the lower cell has one flagellum almost in focus and one out of focus. You could use this image to measure one flagellum from the upper cell. You would have to refocus and take another picture to use the lower cell.

Be organized and systematic with your file titles to keep straight what time point each image is from and which cell within an image is the one to measure. Put all images into the shared space for your lab partners to measure. One note about the community folders and about your files: if you use the Louise public folders for file sharing, save ONLY into your group’s folder and be careful not to alter any other group’s data.

**Image J Instructions—for the ‘measuring’ computer**

ImageJ runs in either Windows or Mac. The instructions below are for Mac, so Windows users will have to explore a bit to find equivalent commands (it is pretty similar). Find the ImageJ icon on the desktop (could look like either icon at right) or search for the app on the computer. After ImageJ is launched, then you can drag and drop image files onto the software window to load them.

We first calibrate the software so it can apply real, physical dimensions to the number of pixels on the computer monitor.

Load your micrometer image in ImageJ. Now draw a line on your calibration image. To do this, select the line tool from the ImageJ menu window.

**Hold the line tool button down (or right click) and select the straight line option.** Draw a line perpendicular to the micrometer markings from the edge one long bar to the same edge of the next long bar.

Notice that the yellow calibration line barely touches the ends of the four short bars. This placement helps ensure that the calibration line is fairly straight, even though the image is at an angle. Also notice that the yellow calibration line goes from the left edge of the long bar on the left to the left edge of the long bar on the right. It is fine to go from the right edge to the right edge instead. (Think about why it would not be accurate to draw a calibration line from the middle of one bar to the left or right side of the other bar.)

Tell ImageJ that the line you have just drawn is exactly 50 µm (micrometers or microns) long. To do this, select “Set Scale…” from the Analyze menu. You will see a dialog box similar to the one at left. **DO NOT CHANGE the ‘distance in pixels’ box on your computer ONLY CHANGE the ‘known distance’ and ‘the unit of length’** This action converts pixels on the monitor to microns. When you have calibrated ImageJ for this session at a given magnification, you do not need to recalibrate. **(Make sure to check the “global” box so that your calibration will apply to all subsequent images.)**
Test your calibration by drawing a line from the middle of one short bar to the middle of the next short bar. Then type “zM” or select “Measure” from the Analyze menu. A new window should appear that shows you the length of the line you just drew. (What should be the approximate length?)

However, notice there are three columns that are irrelevant (Mean, Min, & Max). To get rid of these three columns, while the Results are still displayed, choose “Set Measurements…” from the Edit menu. Deselect all the options as shown:

Close the Results window by clicking on the red dot in the top left corner. Then type “zM” again. If you do not see the Results window, choose Results under the Window menu. Your data should just appear as length in microns, as you have calibrated ImageJ.

Now you are almost ready to measure flagella. From the shared folder online, get the images that your lab partners have already taken of fixed Chlamydomonas cells at different time points. One by one, load each image into ImageJ and ZOOM IN the same amount each time so that you get better resolution to draw the lines to measure flagella. Zooming does not alter the pixels per flagellum and therefore does not affect your ultimate measurements, but it allows you to trace distance and shape of the flagella more accurately.

Because flagella are not straight, you will need to pick a different line-drawing option in ImageJ. Go back to the line tool, and from its pull-down menu, select the free hand option.

Now trace the flagellum from one end to the other, and select Measure (or type “zM” on the Mac). Your data will accumulate in the “Results” window. To be safe, write the numbers down and/or enter them into Excel as you go, in case ImageJ crashes.

You will need to capture at least 15 images of flagella per sample per time point. Your instructor will tell you how many flagella to measure per time point. Count only one flagellum per cell and measure only the longer of the two. Each time you capture the image in TCapture, you will need to measure it in ImageJ. Repeat this process until all your measurements are collected. Luckily, ImageJ can perform some calculations for you. When you have all your measurements collected, choose “Summarize” from the Edit menu. You will find the mean, standard deviation as well as the maximum and minimum lengths you measured. Note that we will learn more about these statistical parameters soon.

When you are ready to measure another round of flagella, you can choose “Clear Results” from the Edit menu to clean out the results table.
Preparing for Next Week’s experiments investigating the role of gene expression and protein localization in the building of a cellular structure:

Before leaving lab today each laboratory group will formulate an hypothesis and design an experiment to test that hypothesis. To formulate an hypothesis, you might just wonder aloud, “What if we...?” For instance, what if we prevent the cells from transcribing any new RNA? What if we prevent the cells from translating any new proteins? What if these plant cells are put in the dark? What would happen in the presence of added ATP? caffeine? glucose? amino acids? EGTA? When your lab team has agreed upon a question of interest, you then should use your knowledge of molecular and cellular biology to formulate a probable answer to your question. For example, if we block translation with the drug cycloheximide, then you might hypothesize that flagella will not regenerate at all. This hypothesis is a good one because it can be tested. A bad hypothesis might be, “Chlamydomonas cells do not like to have their flagella removed and are happier when the flagella are regenerate.” This hypothesis is practically impossible to test for a measurement of Chlamydomonas happiness is not known. Formulate your hypothesis so that you can design an experiment to test it. To help you formulate a testable hypothesis consider the reagents that we can make available to you in lab next week in the table below. Your team must sign up for an option before leaving lab today.

<table>
<thead>
<tr>
<th>FUNCTION</th>
<th>DRUG/REAGENT(S)</th>
<th>STOCK Concentration</th>
<th>FINAL/WORKING Concentration</th>
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<tr>
<td>translation inhibitor</td>
<td>Cycloheximide</td>
<td>2 mg/mL in ethanol</td>
<td>10 µg/mL</td>
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<tr>
<td>transcription inhibitor</td>
<td>Actinomycin D</td>
<td>5 mg/mL in ethanol</td>
<td>50 µg/mL</td>
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<tr>
<td>phosphodiesterase (PDE) inhibitor</td>
<td>Caffeine</td>
<td>66 mM in water</td>
<td>6.6 mM</td>
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<td>elevated and/or reduced Ca++</td>
<td>Calcium</td>
<td>100 mM in water</td>
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<td>EGTA (ion chelator)</td>
<td>100 mM in water</td>
<td>1 mM</td>
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<tr>
<td>disrupts IP₃ production</td>
<td>lithium chloride</td>
<td>1 M in water</td>
<td>20 mM</td>
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<tr>
<td>prevent photosynthesis</td>
<td>darkness (use foil)</td>
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<tr>
<td>alter metabolism</td>
<td>temperature variation</td>
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Remember to include good controls in your experimental design. A good control is an experimental condition that will give you a standard or predictable result against which you can compare the results of the condition you are actually interested in studying. For example, if you wanted to see the effects of disco music on the regeneration of flagella, you would design an experiment that had two experimental conditions:

1) Cells regenerating their flagella in the presence of disco music
2) Control cells regenerating their flagella in the presence of non-disco music at the same volume, beat, etc. (Notice the difference between the control and experimental is only one variable - the presence or absence of disco music.)

Your hypothesis might be that disco music will prevent flagella from growing. This hypothesis is a testable hypothesis because you can measure the length of the flagella in the two situations (with and without disco music). Let’s look at a hypothetical set of results. When the cells are subjected to disco, the flagella did not regenerate. Flagella in the presence of normal music did not grow either. How should these results be interpreted? Did disco prevent the regeneration? What do the results of your
control condition tell you? Why must every experiment have good controls?

Each group should decide upon a question to ask next week, formulate an hypothesis, design the experiment, and discuss the protocol with your instructor. This meeting will give us a chance to answer any major questions you might have and order the reagents you will need. Next week’s lab will be very busy — so careful preparation this week is particularly critical to your success next week.

**Your protocol must be written today** and it should be specific enough so that next week, you can come into the lab and begin immediately by following your own directions. Your instructor will give you feedback on your protocol. Check to see how your instructor will review your protocol — you may need to submit a written protocol.

**You also should have the following in your lab notebook:**
1) The data from today’s experiment.
2) Answer all the questions asked of you in the protocol above (with the exception of the optional question regarding the moon).
3) You should note any observations you think note worthy - use your best judgment.

NEWS ITEM: In lab we study the flagella of the volvocine green algae Chlamydomonas reinhardtii. The multicellular volvocine Volvox is like 1,000-5,000 Chlamydomonas all hooked up in a sphere with their flagella sticking on the outside. This arrangement is rudimentary multicellularity and introduces new hurdles including the fact that the organism (without a circulatory system) needs to figure out how to get nutrients to all of those cells. Diffusion isn’t fast enough. A study published in 2006 showed that Volvox have evolved a mechanism that coordinates flagella beating. In this manner not only can the organism move within its environment but it can also move its environment (i.e., food) across more of its surface area and facilitate feeding. [PNAS 103:8315]

**Lab Notes**
Parameters That Affect *Chlamydomonas* Flagellar Regeneration: Gene Expression and Protein Localization

**Review Reading**
- p 161-165 “How are enzymes…….” to end of chapter
- p 352-354 “How is Eukaryotic……” to “Other Proteins”
- p 361 “Translation of mRNA….” to end of page

**✓ Special Note**
- You will be working with iodine so you may want to wear old clothes.
- This is a particularly busy day in lab and requires exceptionally organized and efficient teamwork.
- You must have your solutions mixed and be ready to go shortly after the start the lab period.

You will perform the experiments you have designed and record your results in Excel spreadsheets modeled after the tables on the following page. If any modifications were made to your written protocols, you should talk to your instructor before you begin your experiment. Today will be a busy session of performing your experiment and collecting data. Do not worry about analyzing your data today. You will spend the next lab session analyzing data.

**Your protocol:**

Note – You may not need to use all the cells in these data tables – check with your instructor to determine how many flagella you should measure and/or for how long you should conduct your experiment.

**Untreated *Chlamydomonas*** (a.k.a. *Chlamydomonas* that were never deflagellated - so you can see what normal flagellar lengths are)

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LM 71
Make Excel spreadsheets modeled after these tables.

### Control Data

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The number of experimental conditions depends upon what you are testing. Each test condition will require its own time course and, therefore, its own spreadsheet like this one.

### Experimental Data

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A Beginner’s Guide to Descriptive Statistics

Statistics
When you have collected a large set of data as you did last week for flagellar regeneration, you need to use some descriptive statistics to convey the important aspects of the distribution of your data. Two features of the distribution that you should describe include:

1) The central tendency
2) The spread of your data

Mean (a.k.a. average)
A simple measure of the central tendency of the data is the mean (or average):

\[
\text{Mean} = \frac{\text{sum of all the data values}}{\text{sample size}}
\]

For example, with the data set (1,1,1, 5), \(n = 4\); the mean is \(8 ÷ 4 = 2\)

Range
The simplest measure of the spread of your data is the range, which tells you the distance between your most extreme data values, but does not address the issue of how frequent these extreme values are. The formula for calculating the range is:

\[
\text{Range} = \text{value of maximum data point} - \text{value of minimum data point}
\]

For example, with the data set (1,1,1, 5):
the range is 5 - 1 = 4.

Variance (Var)
The variance of your data is a measure of spread that will take into account both the deviations of your data (away from the mean) and how frequently these deviations occur. The formula for calculating variance is:

\[
\text{Variance} = \frac{\text{the sum of (each data point - the mean)}^2}{\text{sample size}}
\]

For example, with the data set (1,1,1, 5):
the mean = 2
the variance = \(((1-2)^2 + (1-2)^2 + (1-2)^2 + (5-2)^2) / 4 = 3

Standard Deviation (SD)
The standard deviation of your data is the square root of the variance, and therefore it reflects both the deviation from the mean and the frequency of this deviation. Standard deviation often is used instead of the variance because the scale of the variance tends to be larger than the scale of the raw data, while the standard deviation is on the same scale as most of the data. The formula for standard deviation is:

\[
\text{Standard deviation} = \sqrt{\text{variance}}
\]

For example, with the data set (1,1,1, 5):
the standard deviation is the square root of 3, which is 1.73.
**Standard Error of the Mean (SEM)**

The standard error of the mean is another common way to describe the deviation from the mean and the frequency of this deviation, but it also takes into account the size of your data set. The formula for standard error is:

\[ \text{Standard error} = \sqrt{\frac{\text{variance}}{n}} \]

For example, with the data set (1, 1, 1, 5):
the standard error = the square root of \( \frac{3}{4} \) = 0.866.

To see why standard error is a useful statistical description, let’s consider another data set where the variance equals 3 but \( n = 30 \).

\[ \text{Standard Error} = \text{square root of } \frac{3}{30} = 0.316. \]

The same variance of 3 in the two examples above gave different standard errors (if \( n = 4 \): variance = 0.866 versus if \( n = 30 \): 0.316) because of the difference in sample size.

If you look closely at standard error and standard deviation formulas, you will notice that standard error has taken the sample size into account twice. To some extent, this calculation is statistics at its worst. Standard error is a statistical analysis of one set of data treated as if you had actually repeated the same experiment many times and gotten a range of means. In other words, standard error is a statistical approach that attempts to look at the variance of this imaginary range of means and determine the variance of these means. Some scientists use standard error to make their data look better than they really are.

- \( \bar{x} \) indicates mean
- \( n \) indicates sample size
- SD indicates standard deviation
- SEM indicates standard error of the mean (sometimes abbreviated as SE for standard error)
Using Microsoft Excel for Statistical Analysis

Rather than calculating averages, standard deviations, etc. manually, we will have Excel calculate these values for us. As you already know, you also can use this program to graph the results.

Access the data file that you compiled last week. If you have not done so already, insert the formula for calculating the average into the appropriate column.

Put your cursor on the box in the lower right corner of the box that just calculated your average length at time zero. If you drag the cursor down to your bottom row and then let go, you will have calculated the averages for your remaining rows.

These data are not so easy to present in this tabular format. It is much more clear to present such data as a graph. To graph your data, pick Chart, and make subsequent selections as you’ve done before. Be sure to define the X axis values properly. After some additional formatting, your graph might look something like this image.

In its current form, your averaged data points do not include any indication of the variation in your data. By adding “y error bars” to the averaged data points you can indicate how “tight” or “broad” your averaged values were. Very small/short error bars indicate that the averaged values were very similar, while large/tall error bars indicate that values averaged were quite different.

Most often in published scientific papers you will see that the error bars indicate the standard error of the mean (SEM), which is related to standard deviation (SD). To calculate the SEM you just need to divide the SD by the square root of your sample size (n). You can set up two new columns in Excel to do these calculation on all your data.

First, create a new heading called Standard Deviation in the column next to your averages. Click on the first empty box in this column and insert a function called STDEV. Enter the coordinates where your data are found.
When you hit return, the standard deviation for your data will appear in that cell. Again, you can apply the formula to all the rows simply by highlighting the first cell and dragging down to the bottom row.

Now determine the square root of your sample size (scientific calculators are online if you don’t have one.) Add a new column called **SEM. In the first cell, type =W2/(square root of your sample size)**. This will divide the SD (from cell W2) by the square root of the sample size—the SEM. 

Copy the SEM formula down to all the rows.

Finally, to add **error bars**, double click on any one of the data points within the graph. You will get a window similar to the one shown below left. Select the Y Error Bars tab and pick “both” under the display options. Select the Custom option and click Specify Value. For the Positive Error Value, use the cursor to select all the cells on your worksheet containing the SEM values. Repeat this process for the Negative Error Value, making certain that you selected the very same cells. You will see that error bars have been added to the data points on your graph as illustrated in the graph below right. Remember that error bars in various situations can potentially represent SD, SEM, or even other values (that we haven’t discussed) so your figure legend should include a short sentence that indicates what exactly your error bars represent.

How do you interpret error bars? When SEM error bars from two data sets overlap, the population averages are statistically not different from each other. It’s a little more tricky when SEM bars do NOT overlap. To make it simple, we’ll say here that non-overlapping SEM error bars mean that the two population averages are indeed statistically significantly different. In reality, to meet the commonly accepted threshold for statistical significance ($p<0.05$), SEM error bars would have to not overlap AND be separated a bit further, by about 50% the width of the bars.
Insert your flagellar regeneration graph (with error bars) here: