SAFETY RULES

1) Know the location of all safety equipment such as fire extinguishers, eye-wash fountains, safety showers, and fire blankets.

2) ALWAYS WEAR EYE PROTECTION. Safety glasses or other eye protection must be worn in lab at all times. Chemicals used in Biochemistry, for the most part, are not hazardous. However, we will use strong acids and bases as well as oxidizing and reducing reagents, which can be very dangerous. When called for, goggles or face shields will be made available for protection against UV lamps and more hazardous compounds.

3) Never mouth pipet or taste anything in lab.

4) No smoking, eating, or drinking is allowed in the laboratory.

5) Long hair should be tied back out of the way of flames and apparatus.

6) **No open-toed shoes are allowed in the laboratory**

7) **No above-knee skirts, dresses, OR SHORTS are allowed in the laboratory unless you wear them underneath a lab coat**

8) Extra clothing, books, and bags should be stored on the shelves provided.

9) Report all injuries, no matter how trivial.

10) Never work in the laboratory alone or perform unauthorized experiments.

11) An untidy or dirty laboratory is an inherently unsafe laboratory. You will be held responsible, both as individuals and as a class, for ensuring that your lab bench, the common-use areas of the lab, and all pieces of apparatus are clean and well-organized when you leave the lab each day.

12) Protective gloves will be available to wear when handling potentially caustic materials. It is recommended that you don’t wear expensive clothing to lab in case of spills, or that you wear a laboratory coat.
LABORATORY RECORDS

You will be working in small groups for the semester. Each person in a lab group may do different parts of a larger experiment. To ensure that each member of a group has equal access to all the data collected by the group, notebooks will be provided in which ALL data collected by the group is to be placed. The notebooks will be three ring. Your group will need to provide notebook paper.

The data collected in the notebooks are the raw data and any commentary that you wish to add. The data should be well organized, neatly recorded and labeled with your group identification, the name of the person who collected the data, the experiment title and subtitle, and the date the data were collected. If your handwriting is a bit on the unreadable side you MUST recopy the data you collect and place it in the group’s notebook during the same lab period in which the experiment is done. If you choose to recopy any notes, you must of course include ALL original data records in case an error is made in copying the data.

Notebooks are NEVER to leave lab except for photocopying. Anyone who removes a notebook from lab for an extended period of time without permission from their lab partners and/or the instructor will earn a failing grade for the lab assignment in question.

The lab notebooks will be collected and evaluated twice during the semester: once during an oral interview with your lab instructor, and once at the end of the term. The score your group receives on the organization and presentation of the data in the notebook will be included in your laboratory grade. If you have any questions about how to record your data in the group’s notebook, please speak with the instructor.

LABORATORY ASSIGNMENTS

There will be two laboratory investigations during the semester:

1) Purification and Characterization of Yeast Cytochrome c.
2) Kinetic Analysis of an Enzyme, Tyrosinase.

Several assignments will be required as a part of each investigation. See the rest of the lab manual and subsequent handouts for specific instructions. At the end of the first project there will be a written lab quiz. At the end of the second project your group will present a poster on your work kinetic analysis of the enzyme tyrosinase. More detailed instructions on this aspect of the lab will be provided later in the semester.

SPECIAL NOTE: Although data will be collected in small groups, some assignments must be synthesized and written individually, and all group members are expected to be responsible for all the material at all times. The level of responsibility that you show in your lab group will be reflected in your subjective assessment score.
EXPERIMENT I
PURIFICATION AND CHARACTERIZATION
OF YEAST CYTOCHROME C

LAB ONE

Objectives:  Understand the Big Picture
Understand how to make measurements in the biochemistry laboratory
Introduction to this manual
Make buffer

Pre Lab - overview of purification and characterization procedure

Procedure:
Make approximately one liter of phosphate buffer - 100 mM NaPi, pH 7.0
LAB TWO

Objectives: Construct protein concentration standard curve
Pour gel filtration column, and determine void volume

Procedure:

Part I. Construct protein concentration standard curve

Total protein concentration will be determined using the Bio-Rad protein assay dye kit. Full instructions can be found in Appendix B of this manual.

Part II. Pouring a gel filtration column, and measuring the void volume

1. Obtain an 88 ml column (40 x 1.7 cm) and clamp it onto a ring stand with two clamps. Check that the column is vertically true, then attach a stopcock to the outlet.

2. Obtain enough Sephadex G-50-150 slurry to fill your column.

3. With the stopcock closed, pour approximately 25 mL of NaPi buffer containing 3 mM 2-mercaptoethanol into the column. ALWAYS WORK WITH THE NEAT 2-MERCAPTOETHANOL IN THE HOOD, NOT AT YOUR BENCH. Be sure that no air bubbles are trapped in the column or the stopcock. You can drain some buffer through the stopcock to remove any bubbles.

4. Fill your column with the slurry and let the column pack by gravity until a bed of at least 1 cm is evident. Once this packed bed is established, open the stopcock and continually add slurry until the column is packed to about 3 cm below the colored column top. Once it is packed, rinse the column for 15 min with buffer at a flow rate of approximately 1 mL/min.

HELPFUL HINT: So you don’t have to baby-sit the column as it rinses and to maintain an even flow rate for future runs you will want to set up a top tubing connector. This is done by cutting a 200 µL micropipet tip with a razor blade so that the large end fits snugly over the column cap inlet port. To the pipet tip is connected a length of small, clean Tygon tubing. Buffer is then drawn up into the tubing, and while maintaining a filled tube the column top is replaced on the column. The other end of the tubing is kept in a flask of buffer, and the flask is raised or lowered to the appropriate height to maintain a 1 mL/min flow rate. When the column is stored, the stopcock is closed, and Parafilm should be wrapped around the top and bottom of the column. The column will then be ready for use in the future. You are strongly advised to check your column every few days in order to ensure that there is always a layer of buffer on top of the column to prevent it from going dry. If your column runs dry, you’ll have to re-pour it, and repeat any measurements you have made with it up to that point in the semester.

5. Follow the instructions in Appendix C for running your Gel Filtration Column.
   Note: You will only need 30 glass tubes for this run.
Your sample will be a 1 mg/mL solution of blue dextran. Because your sample is strongly colored, you'll know when it has eluted. There's no need to continue collecting fractions after all the blue dextran is off the column. Because Blue Dextran is blue, it can be quantified by the absorbance of light at a wavelength of about 600 nm.

6. Graph your results and determine the void volume of your gel filtration column. Record this value in your notebook.

**Assignment:**
1) Use a spreadsheet program (such as Excel) to generate a standard curve for determining protein concentration. In the spreadsheet should be the original data (µg of lysozyme and related absorbance in triplicate).
2) Using the equation of the standard curve, calculate the amount of protein in each standard sample and list this in the spreadsheet. In other words, calculate the value indicated by the absorbance of that sample using the equation of the straight line, as opposed to the amount measured out from the stock lysozyme solution.
3) Turn in this well-organized and labeled spreadsheet. Pay special attention to the number of significant figures you are reporting; do not report more digits than are justified by your data.
4) Turn in a labeled and titled graph of your standard curve.
5) Suppose that one of your classmates ran a standard curve and computed an equation for the line of \( y = 0.065 x + 0.34 \). If this student also measured the absorbance of an experimental sample, combined with dye in the same manner as the lysozyme standards, and found an absorbance value of 0.45, how much protein was in her sample?
LAB THREE

Objectives: Yeast lysis - done for you prior to lab
Homogenate preparation
Bulk cation exchange

WARNING - This lab may run longer than the assigned time so please plan your schedule accordingly. You will get this time back later in the semester when we have some short labs. If you have a commitment that requires that you leave lab by 4:20 pm, please make arrangements in advance with your instructor and your lab partners.

VERY IMPORTANT NOTE: If you do not remove a sample for analysis after each step of the purification process your entire study and all its results will be meaningless. You should also accurately measure the volume of all samples resulting from the various purification steps. Again if you do not do this your data cannot be interpreted fully.

Procedure:

Part I. Yeast homogenization (done prior to lab)

Approximately 200 g of wet packed *Saccharomyces cerevisiae* (Baker’s yeast, Fleischmann’s Yeast Company) was weighed out and suspended in an equal volume of 100 mM NaPi, pH 7.0, containing 1.0 M NaCl, 3 mM 2-mercaptoethanol, and a protease inhibitor cocktail (see Appendix D for details). An equal volume of glass beads was added and the yeast were homogenized for 4 min on ice (alternating 30 s on/30 s off) in a Bead Beater (Biospec Products, Bartlesville, OK).

You should keep this homogenate and all subsequent fractions on ice at all times.

Part II. Homogenate clarification

To get rid of insoluble cell contents (what might these include?) centrifuge the lysate at 4,000 x g for 10 min at 4°C. Collect the supernatant, measure its total volume and set aside 5 mL for use in determining total protein concentration and cytochrome c concentration. Keep the unused portion of the sample at -20°C for later analysis by SDS-PAGE. Note: you will be held responsible for all samples stored at -20°C in this and subsequent steps, so take the necessary steps to insure that they are labeled and secure.

Part III. Bulk cation-exchange

1. Equilibrate ~10 g of Bio-Rex 70 cation exchange resin (100-200 mesh, Na+ form; from Bio-Rad) with two washes of 100 mM NaPi, pH 7.0.

2. Dilute the homogenate with six volumes of distilled deionized water (ddH₂O) containing 3 mM 2-mercaptoethanol, and add the suspension to the equilibrated resin.

3. Stir the mixture at 4°C for at least 30 min. Be sure that the stir bar is rotating evenly and
at a moderate speed -- just enough to create a small vortex and evenly suspend the resin.

4. Set up a small syringe column by cutting a Teflon frit to the proper size and inserting it into a 20 ml syringe (smooth side of frit towards the resin). You can use a cork borer to cut the frit, then clamp the column onto a short ring stand. The instructor or lab assistant will help you assemble this column.

5. When the 4°C incubation is complete, turn off the stirbar and let the resin settle to the bottom of the beaker. Remember to make observations as to color, cloudiness etc. at all points. Decant the supernatant into a separate beaker. Rinse the resin into your syringe column and let the column run dry.

6. Rinse the resin with one column volume of 100 mM NaPi, pH 7.0, containing 3 mM 2-mercaptoethanol.

7. Elute the cytochrome c with a minimum of 100 mM NaPi, pH 7.0 containing 0.8 M NaCl and 3 mM 2-mercaptoethanol. Your goal should be to capture all of the pink liquid in one tube, but as little unpigmented liquid as possible. Accurately measure the volume of the eluate. Keep a portion (~50 µL) of this eluate at -20°C for SDS-PAGE analysis.

   If you are out of time at this point, you may postpone the analysis of your samples until next week.

8. Determine the total protein content in the three samples you have collected (clarified yeast homogenate, cation exchange decant, cation exchange eluate) using the BioRad protein assay. You must also generate a new standard curve (triplicate samples) as done in Lab Two. You will have to dilute your experimental samples (with NaPi buffer) some unknown amount in order for the values to register on the standard curve. This is a trial and error process, but you can try several dilutions at once, as described in Appendix E. Calculate the quantity of protein in each sample (total volume) remembering to figure in any dilution factors that were made.

   HELPFUL HINT: To avoid interference by particulates in your samples, microcentrifuge the samples for 2 min at maximum rpm prior to analysis using the Bio-Rad protein concentration kit and UV/vis spectroscopy.

9. The same samples must now be examined for cytochrome c concentration. Using an extinction coefficient of 95.0 mM⁻¹ cm⁻¹ for cytochrome c at 410 nm you should be able to calculate the quantity of cytochrome c in each sample.

10. Store eluate at 4°C and other samples you wish to keep for SDS-PAGE analysis at -20°C (this should include a 50µL sample of your eluate).
NOTE: It is a good idea to take some time during this lab period to calculate your total protein and cytochrome c concentrations and determine how they are used in constructing the purification table. After generating a first set of numbers you may find that some results are erroneous. If this is the case you may need to repeat some measurements and/or explain the results in a convincing way. That is, your lab instructor must agree with your interpretation. If you leave without first going over these data you may be left in a tight situation when it comes to the assignment if you find that you cannot calculate the numbers correctly or if some mistake or misinterpretation has been made.

**Assignment:** Fill out the sheet in the appendices entitled "Ten Questions" (Appendix F).
LAB FOUR

Objectives: Begin dialysis of sample
Learn how to construct a purification table

Procedure:

Part I. Sample dialysis

1. Equilibrate a piece of dialysis tubing by soaking for 30 min (without stirring) in ddH₂O containing a pinch of EDTA. The tubing should be cut to hold approximately 2 - 3 times the volume of your eluate sample.

2. Place the eluate from the cation exchange into the dialysis bag, with one end sealed, then seal the other end leaving enough room for influx of buffer during the dialysis. Try to minimize air bubbles trapped in the dialysis bag. The instructor or lab assistant will help you with this procedure. Dialyze at 4°C against one liter of 100 mM NaPi, pH 7.0 containing 3 mM 2-mercaptoethanol, with gentle stirring. The dialysis buffer should be replaced with one liter of fresh buffer at least twice over the next week with no less than 5 h between changes.

Part II. A Purification Table

1. Your instructor will walk you through the construction of a purification table, using data for a hypothetical cytochrome c purification.

Assignment: Construct a purification table for cytochrome c using the table of data distributed in class (the format of your table should follow the example given) and answer the following questions.

Which step in the sample data purification table purifies the cytochrome c the greatest amount?
Which purifies it the least amount?

Briefly explain your answers.

NOTE: you may want to check on your stored gel filtration column during this class period to ensure that it is still sealed and properly preserved for next week’s lab.
LAB FIVE

Objectives: Concentrate sample
Standardize gel filtration column

Procedure:

Part I: Sample concentration

1. Using a disposable plastic pipet, remove your sample from the dialysis bag.

2. Clarify the sample by centrifugation.

3. Determine the levels of total protein and cytochrome c in this sample as before. You must repeat the standard curve determination (in triplicate) as in Lab Two.

4. Keep a portion (50 µL) of this sample at -20°C for analysis by SDS-PAGE.

5. In order to load your entire sample onto your gel filtration column it must be concentrated to a volume of < 2 mL. To do this you will be using a vacuum concentrator. Your sample should be placed in a series of microcentrifuge tubes and labeled with your group's name. The samples will then be placed in the apparatus and concentrated. You will be shown how to correctly operate the instrument. Once concentrated to the appropriate volume, the sample can be covered and stored at 4°C. Note that concentrating a large sample down to < 2 mL may take several days, so plan your group's time accordingly.

Part II: Standardize the gel filtration column

1. Follow the instructions in Appendix C for running your Gel Filtration Column.
   Note: You will need 95 glass tubes for this run.
   The standard solution contains 3 mg each of aprotinin (6.5 kDa), bovine cytochrome c (12.5 kDa) and carbonic anhydrase (29 kDa) in a total volume of 2 mL.
   Continue collecting fractions until the A_{280} profile is flat and roughly equal to the background absorbance level.

2. Analyze all fractions for protein (absorbance at 280 nm) and cytochrome c (absorbance at 410 nm) levels using the plate reader. Think about why your data display the trends they do, and be prepared to explain these trends to your instructor next week.

Assignment: Graph the results of your gel filtration standardization so that you can discuss your data with your instructor in lab next week. There are instructions in Appendix G that show you how to extract three clean peaks from your absorbance data. Please bring a graph similar to that shown for step 4 of the process to lab next week.
LAB SIX

Objectives: Gel filtration of concentrated sample
Plan SDS-PAGE gel

Procedure:

Part I. Run sample on gel filtration column

1. Follow the instructions in Appendix C for running your Gel Filtration Column.
   
   Note:
   You will need 95 glass tubes for this run.
   Before loading your sample on the column, bring the volume to 2.0 mL with NaPi buffer, then add 5 μL of 2-mercaptoethanol and allow the sample to sit for five minutes. Microcentrifuge it for 2 min, carefully remove the supernatant (be sure not to disturb the pelleted debris) and load it onto your column using a plastic pipet. Continue collecting fractions until the $A_{280}$ profile is flat and roughly equal to the background absorbance level.

2. Analyze all fractions for protein (absorbance at 280 nm) and cytochrome c (absorbance at 410 nm) levels using the plate reader.

3. Graph the gel filtration results, and combine the fractions that represent each of the peaks of protein, being particularly cautious with the peak of cytochrome c. Most of your cytochrome c will be found in the fractions with absorbance levels greater than 50% of the peak value. Be sure to record the total volumes of each of these combined fractions.

4. Perform total protein and cytochrome c content assays on the pooled fractions as previously described. Save these pooled fractions at -$20°C$ for SDS-PAGE analysis.

Assignment: Once you have determined the levels of total protein and cytochrome c in your pooled gel filtration samples, you should prepare a final version of your purification table. This should be done keeping three things in mind; 1) neatness, presentation, and organization count, 2) significant figures are extremely important in the values you give, and 3) although you need only hand in one table for your group, each person must know how to do all of the calculations because there will be some aspects of these calculations on the lab quiz!

Part II. Plan samples for SDS-PAGE

1. Determine what samples you wish to run on your SDS-PAGE gel. There will be ten lanes per gel, one of which must be a molecular mass standard (provided by instructor) while a second must be reserved for a sample of pure cytochrome c (1 μg/μL; also provided by the instructor). You will have two gels per group; thus you have as many as sixteen lanes for your samples.
2. Consider whether you wish to run multiple dilutions of any of your samples on your SDS-PAGE gel to ensure good results. The method of protein visualization that you will use -- Coomassie Brilliant Blue dye -- typically requires 0.1 to 1 µg of protein per band for a strong signal.
LAB SEVEN

Objectives:  SDS-PAGE

Procedure:

Part I: SDS-PAGE

1. Set up two gels per group. General instructions for running SDS-PAGE gels are provided in Appendix H. Your instructor or lab assistant will help you set up the gel apparatus.

2. As one group member prepares the apparatus the others should be preparing the samples.

NOTE: It takes most groups at least 60 min to set up the gel, prep the samples, and load the gel. It takes 60 min to run the gel and 60 min to go through the first staining step. This totals 3 h. Thus you must be organized and efficient to ensure that the lab is completed within a reasonable amount of time.

3. Remove 50 µl of your sample and place in a small microcentrifuge tube that contains 50 µl of 2x sample buffer (provided by instructor). This will provide enough volume to run three amounts of each sample -- 20 µL in one lane, 5 µL in another, and 2 µL in a third -- and still leave enough to run a second gel if something goes awry.

4. Heat all samples, except the molecular weight standards, for 3 min at 100°C in the heating block. After heating, microcentrifuge the samples for 2 min at the highest spin setting. The samples are then ready for loading.

5. Load your samples. Save any extra sample at 4°C until you are sure that the gel was a success.

6. Run the gel at ~150 V for ~45 minutes, until the dye front is within 1 cm of the bottom of the gel.

7. Stain and destain the gel according to the directions in Appendix H. This will take several days. Next lab period we’ll take digital photographs of the gels for use in your final assignment.
Assignment:
1) Gel filtration analysis.

From your elution profile of the gel filtration standards (absorbance: y-axis vs. elution volume: x-axis) generate a gel filtration standard curve (log MW: y-axis vs. v_e/v_o: x-axis).

From the standard curve and the sample gel filtration graph (absorbance: y-axis vs. elution volume: x-axis) calculate the molecular weight of your cytochrome c sample.

Turn in an elution profile of your gel filtration standards, a gel filtration standard curve graph, and an elution profile of your experimental sample, and provide the measured molecular weight of your cytochrome c sample on the standard curve graph. All three graphs should be properly presented with appropriate labels and titles.

2) SDS-PAGE analysis.

From the migration of your molecular weight standards, generate an SDS-PAGE standard curve (log MW: y-axis vs. R_f: x-axis).

From the standard curve and the migration of your purified cytochrome c pool, calculate the molecular weight of your cytochrome c sample.

Turn in the SDS-PAGE standard curve (including the measured weight of your cytochrome c sample) and a professional-quality figure (with legend) for your SDS-PAGE results. This figure should be executed in the style of a figure in a Journal of Biological Chemistry article.

3) Answer the following questions.

A. Did the purification work? Compare the results of your purification table and your gel to answer this question. Do the numbers from your purification table agree with your visual assessment of your SDS-PAGE results? Which is a more accurate representation of the true purity of the sample? Explain briefly.

B. Which purification step worked best, and which worked least, as determined by overall analysis? Explain briefly.

C. Do you believe your data? In other words, are there anomalous values that need to be explained?

D. What is the apparent molecular weight of your cytochrome c as determined by SDS-PAGE and how does it compare to your value from gel filtration? If they are different, which is more credible and why? (More points will be given for a more rigorous answer.)
LAB ONE

Objectives: Understand the Big Picture
Determination of pH optimum for tyrosinase activity

Pre Lab - overview of tyrosinase activity, general kinetics overview, and spec use.

Procedure:

Part I. pH Profile of Tyrosinase Activity

1. Solutions (provided)

<table>
<thead>
<tr>
<th>Buffer</th>
<th>5.0</th>
<th>6.0</th>
<th>7.0</th>
<th>8.0</th>
<th>9.0</th>
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</thead>
<tbody>
<tr>
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<td>2.50-x</td>
<td>2.50-x</td>
<td>2.50-x</td>
</tr>
<tr>
<td>L-dopa</td>
<td>0.50</td>
<td>0.50</td>
<td>0.50</td>
<td>0.50</td>
<td>0.50</td>
</tr>
<tr>
<td>Tyrosinase</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
</tbody>
</table>

*units are milliliters

NOTE: the value "x" in the table below refers to the proper amount of tyrosinase, which will be determined by your instructor prior to each lab. This value will be written on the white board in the front of the room.
Assignment: Generate a figure and legend for the data you collected on the pH profile of tyrosinase with L-dopa. Examples of such a figure can be found in many *Journal of Biological Chemistry* articles. For this figure you should provide both the level of detail and the quality of presentation that you would expect to find in a professionally produced figure and figure legend. Think carefully about your choice of graph form, and be sure that your figure legend includes the optimal pH value you determined.
LAB TWO

Objectives:  
K_M determination  
Inhibitor test #1

Today you will use the general procedure in Appendix I to measure the activity of tyrosinase at several concentrations of substrate, in the presence and absence of an inhibitor. You’ll calculate the K_M and V_max for tyrosinase under these conditions by inserting your initial velocity values into the Michaelis-Menten equation. Laptops will be available for you to process your data on the fly in lab.

Part I. Control K_M Determination for Tyrosinase

Using the quantity of tyrosinase that yields linear product formation for 3 min. (“x” below, determined by instructor) use the following table to prepare your solutions for determination of initial velocities. You will have to make the appropriate buffer, the one you determined in Lab 1 to have the pH that supports optimal tyrosinase activity. Feel free to make arrangements with your instructor to come in early to make the appropriate buffer.

<table>
<thead>
<tr>
<th>Assay #</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buffer</td>
<td>2.90-x*</td>
<td>2.80-x</td>
<td>2.60-x</td>
<td>2.40-x</td>
<td>2.20-x</td>
<td>2.00-x</td>
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<tr>
<td>L-dopa</td>
<td>0.10</td>
<td>0.20</td>
<td>0.40</td>
<td>0.60</td>
<td>0.80</td>
<td>1.00</td>
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<tr>
<td>Tyrosinase</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
</tbody>
</table>

*units are milliliters

Part II. Inhibitor Analysis

Having established the kinetic parameters for tyrosinase activity in the absence of an inhibitor, you will now test its activity in the presence of sodium benzoate. To do this, you will use 100 mM sodium benzoate in the appropriate buffer.

1. First determine the amount of inhibitor necessary to reduce the enzyme activity to approximately 20% of the uninhibited reaction with 0.40 mL of L-dopa (the volume of buffer plus inhibitor should equal 2.60-x mL).

2. Once you have determined how much inhibitor to use, you should follow the general procedure outlined above for the determination of K_M, except that another substance (the inhibitor) will be in the assay mixture, added into the blank, as shown in the table below. Note that you will be determining K_M apparent and V_max apparent -- the kinetic parameters for the enzyme under the influence of the inhibitor.

NOTE: the values x and y in the table refer to:
  x) the tyrosinase quantity necessary to give an absorbance change of ~0.2 units per min. (determined by the instructor)
y) the quantity of inhibitor necessary to reduce the rate of activity to ~20% of the uninhibited rate (from step 1)

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Assay #</th>
</tr>
</thead>
<tbody>
<tr>
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<tr>
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<td>2.90-x-y*</td>
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<tr>
<td>L-dopa</td>
<td>0.10</td>
</tr>
<tr>
<td>Inhibitor</td>
<td>y</td>
</tr>
<tr>
<td>Tyrosinase</td>
<td>x</td>
</tr>
</tbody>
</table>

*units are milliliters

NOTE: If you finish this lab with time to spare you may continue on to LAB THREE, analyzing the effect of the second inhibitor. If you do both inhibitors today, you only need to run the control $K_M$ determination once.

**Assignment:**

1) Generate a professional-caliber figure and legend for the data you collected for the $K_M$ determination of tyrosinase with L-dopa, with and without the inhibitor sodium benzoate.

This assignment requires you to produce a Lineweaver-Burk figure with two lines on one graph. Examples of such figures can be found in many *Journal of Biological Chemistry* articles. Please provide both the level of detail and the quality of presentation that you would expect to find in a professionally produced figure and figure legend. In your figure legend, be sure to give $K_M$ and $V_{max}$ values you determined (paying close attention to significant figures) as well as the type of inhibition determined by an analysis of the lines on the graph.

2) Based on your graph, what kind of inhibitor do you think sodium benzoate is? Please explain briefly. Be sure to take into account the uncertainty in your numbers when you make your analysis.

3) Can you rationalize your answer to #2 based on the structure of the inhibitor? Please explain briefly.
LAB THREE

Objectives:
- $K_M$ determination
- Inhibitor test #2

Procedure:

Perform this experiment exactly as you did in Lab Two. The only difference is that you will use a 100 mM solution of Thiourea in place of sodium benzoate.

*NOTE*: The $K_M$ determination must be repeated because the variation between tyrosinase samples and substrate solutions may cause errors in comparing data collected on different days.

LAB FOUR

Objectives:
- A control $K_M$ determination and your first independently-designed experiment

This experiment **MUST** be laid out before coming to lab. You are strongly encouraged to discuss the details of this experiment with your instructor during labs two and three.

LAB FIVE

Objectives:
- A control $K_M$ determination and your second independently-designed experiment

This experiment **MUST** be laid out before coming to lab. You are strongly encouraged to discuss the details of this experiment with your instructor during labs two through four.

**Assignment:** The results from these last two weeks of analysis of tyrosinase will be presented next week (the last week of classes) as a poster presentation. Guidelines are presented in Appendix J. Think about your presentation and prepare ahead of time since you will have many other assignments from this and other classes to fit into your schedule during the last week.
APPENDIX A

Procedure for using the Hanna pH 211 Microprocessor pH Meter

If the pH meter is turned off...

- Plug the pH meter into the wall.
- Plug the electrode into the pH meter.
- Remove the protective cap. Don't be alarmed if salt deposits are present. This is normal and they will disappear when rinsed with water. Immerse probe in the storage buffer solution.
- Switch the instrument on by pressing ON/OFF. The meter automatically defaults to pH measurement mode.

If the meter is turned on...

- Rinse the probe thoroughly with DI water whenever moving from one solution to another.
- For calibration:
  - Immerse the probe approximately 1 ½ in into the pH 7 solution and stir gently. Press CAL.
  - Check that BUFFER pH 7.01 appears on the screen. The “NOT READY” indication will blink until the reading has stabilized.
  - When the reading is stable and close to the selected buffer, the “READY” indication will appear and “CFM” will blink. Press the CFM button to confirm the calibration. (If the reading does not stabilize, ask your instructor for assistance.)
  - Rinse the electrode and move it into a second calibration buffer solution (either pH 4 or pH 10) and stir gently.
  - The LCD display should display the second buffer value. If necessary, use the arrow keys to select a different buffer value.
  - When the reading is stable, press CFM to confirm the calibration.
  - The meter stores the reading and returns to the normal mode.
- To measure the pH of a solution, submerge the tip of the electrode 1 ½ in. into the sample. Allow time for the electrode to stabilize. The pH reading is displayed on the primary LCD.
- When the electrode is not in use, be sure that it is immersed in storage solution.

For long-term storage...

- To prevent damage to the electrode, remove the pH electrode from the solution before turning the meter off. If the meter is off, detach the electrode from the meter before immersing the electrode in storage solution. Replace the solution in the protective cap with a few drops of storage solution and affix the protective cap.
APPENDIX B

Bio-Rad/Bradford assay for total protein  (modified from Bradford (1976), Analytical Biochemistry 72:248)

Preliminary notes:

i. The Bio-Rad protein assay dye kit is a commercial implementation of the tried-and-true Bradford procedure. The molecular basis of the procedure involves the binding of the dye Coomassie Brilliant Blue G-250 to basic and aromatic amino acids in proteins. Upon binding to the protein, the absorbance characteristics of the dye molecules change; for an acidic solution, the $\lambda_{\text{max}}$ shifts from 465 nm to 595 nm and the absorbance at 595 nm primarily reflects protein-bound dye molecules rather than free dye molecules.

Unfortunately, because the binding of Coomassie to proteins is mediated by aromatic and basic amino acids (especially Arginine), and because different proteins have different numbers of basic and aromatic amino acids exposed to the dye, different proteins bind different numbers of dye molecules. For instance, equal amounts of two of the most commonly used "generic" proteins, lysozyme and albumin, give $A_{595}$ values that differ by a factor of about 2.

However, to a first approximation, the binding of dye (and thus the absorbance at 595 nm) is proportional to the number of amino acids and thus to the mass of the protein. Always include a blank (no protein) to determine the level of absorbance resulting from the free dye, and if you want to know the true range of error of the assay, use multiple proteins as standards.

ii. As with other absorbance-based assays, it is critical to develop a good standard curve to relate protein concentration to dye binding (and $A_{595}$). Because of different amino acid compositions of different proteins, it would be ideal to use the protein that you are studying to generate your standard curve, but this is usually impractical, so egg white lysozyme and bovine serum albumin (BSA) are commonly used as generic protein standards. The procedure is simple but it is very important to keep accurate records of each sample prep and analysis. For the most reliable results, prepare every sample in triplicate, and run a new standard curve for every set of experimental samples.

iii. When making up dilutions of standards and samples, it is vital that
   - buffer used for the standards reflect the buffer used for the experimental samples
   - attention be paid to compounds that interfere with the Bradford assay. A detailed list is included in the Bio-Rad assay kit instructions. SDS above 0.001% may affect linearity; above 0.002%, definitely.
   - tubes (or microplates) be clean (this is NOT the time to save money by reusing tubes!)

iv. The Bio-Rad dye reagent contains Coomassie, phosphoric acid, and methanol. The Coomassie dye will stain skin and clothing and neither the phosphoric acid nor the methanol is good for your health. Treat it with appropriate care.
The Assay:

1. Prepare dilutions of standards and experimental samples; the total volume of each well should be 200 µL

   A. Standards. You will use egg white lysozyme (0.1 mg/mL in 100 mM NaPi, pH 7.0) as your protein standard. Prepare samples in triplicate according to the following table. Add the lysozyme and buffer, then proceed to step #2.

<table>
<thead>
<tr>
<th>SAMPLE</th>
<th>µL buffer</th>
<th>µL lysozyme stock</th>
<th>µL dye</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blank</td>
<td>160</td>
<td>0</td>
<td>40</td>
</tr>
<tr>
<td>0.5 µg</td>
<td>155</td>
<td>5</td>
<td>40</td>
</tr>
<tr>
<td>1 µg</td>
<td>150</td>
<td>10</td>
<td>40</td>
</tr>
<tr>
<td>2 µg</td>
<td>140</td>
<td>20</td>
<td>40</td>
</tr>
<tr>
<td>3 µg</td>
<td>130</td>
<td>30</td>
<td>40</td>
</tr>
<tr>
<td>4 µg</td>
<td>120</td>
<td>40</td>
<td>40</td>
</tr>
<tr>
<td>5 µg</td>
<td>110</td>
<td>50</td>
<td>40</td>
</tr>
<tr>
<td>7.5 µg</td>
<td>85</td>
<td>75</td>
<td>40</td>
</tr>
</tbody>
</table>

   B. Experimental samples. In order to keep the final volume equivalent to that of your standards, all samples should have a volume of 160 µL (before addition of dye). Depending on the situation, this may be pure sample, or may be a mix of sample and buffer. See Appendix E for an example of how to set up dilutions of experimental samples collected during cytochrome c purification.

2. Add 40 µL dye concentrate to all microtiter wells (see notes on timing under step 4).

3. Pipet up and down gently but very thoroughly. Good initial mixing is extremely important for a successful assay, but be sure to avoid creating bubbles in the wells, since this will adversely affect absorbance readings.

4. Incubate for at least five minutes at room temperature (for best results, don't go over an hour). Color development is somewhat time sensitive, so all samples should be incubated approximately the same amount of time from addition of dye to spectrophotometry.

5. Read absorbance at 595 nm in the microplate reader. Instructions are posted by the instruments.

6. Use a spreadsheet program to generate a standard curve from your lysozyme concentration standards. To do this, plot quantity of protein on the x-axis versus absorbance at 595 nm on the y-axis.

7. Compute the amount (or concentration) of protein in experimental samples based on the standard curve. To a first approximation, the uncertainty in any protein determination derived from your standard curve will equal the computed amount of protein in your experimental sample times the fractional uncertainty of the slope of your standard curve (the uncertainty on the slope divided by the magnitude of the slope. That is,

   \[
   \frac{\text{uncertainty in } \mu\text{g protein}}{\mu\text{g protein}} = \frac{\text{uncertainty in slope}}{\text{slope}}
   \]
APPENDIX C

General Instructions for Running your Gel Filtration Column

1. Prepare enough glass tubes to collect all the fractions you anticipate needing (30 should suffice for void volume, up to 95 for other experiments) and place them in the fraction collector. Connect a piece of tubing to the outlet of the stopcock and the arm of the fraction collector (use a pipet tip as before). The instructor or lab assistant will help you set up your fraction collector.

2. Rinse your column for 15 min with buffer (100 mM NaPi, pH 7.0, containing 3 mM 2-mercaptoethanol). Measure the flow rate of your column (in mL/min) and check to see that it agrees with your previous flow rate determination (unless this is your first run). If it does not agree, adjust the height of the buffer reservoir to correct the flow rate. Make sure that you are measuring the flow rate through the tubing attached to the fraction collector.

3. Remove the buffer inlet from the column top and open the stopcock to let the buffer drain until it is just at the bed level then quickly close the stopcock.

4. Carefully load 2 mL of your sample onto your column using a plastic transfer pipet, being careful not to disturb the bed surface.

5. Open the stopcock and begin collecting 1.0 mL fractions (collected by elution time).

6. When the sample has entered the column resin carefully add 2.0 mL of buffer using a plastic pipet (continue to collect fractions throughout this step).

7. When the buffer has entered the column resin carefully fill the column with buffer using a plastic pipet (continue to collect fractions throughout this step).

8. Replace the column top without the tubing attached.

9. Attach the buffer-filled tubing to the column top. The column should now flow at a rate of about 1 mL/min as you previously determined, and you can collect fractions until you are sure that all analyte has eluted from the column.

10. After all components have eluted, stop the flow of the column by closing the stopcock, and carefully store your column on its ring stand on the side bench of the lab (be sure to label it and to wrap Parafilm carefully around the top and bottom).

11. Measure accurately the volume of your fractions (one fraction in the middle should be enough) using your pipetman. Record this volume for later use in graphing.

12. Transfer aliquots of all fractions to a microtiter plate for spectroscopic analysis at the appropriate wavelength(s).
Protease Inhibitor Cocktail for general use

Catalog Number: P2714
Storage Temperature: −20 °C

Product Description
Crude cell extracts contain a number of endogenous enzymes, such as proteases and phosphatases, which are capable of degrading the proteins present in the extract. The best way to improve the yield of intact proteins is to add inhibitors of those enzymes known to be present. This protease inhibitor cocktail has been optimized and tested for general use. It is a mixture of water-soluble protease inhibitors with a broad specificity for the inhibition of serine, cysteine, and metalloproteases.

Specific inhibitory properties of the components are:
- AEBSF – [4-(2-Aminoethyl)benzenesulfonyl fluoride hydrochloride] – serine proteases, e.g., trypsin, chymotrypsin, plasmin, kallikrein and thrombin
- Aprotinin – serine proteases, e.g., trypsin, chymotrypsin, plasmin, and kallikrein; human leukocyte elastase, but not pancreatic elastase.
- Bestatin hydrochloride – aminopeptidases, e.g., leucine aminopeptidase and alanyl aminopeptidase.¹,²,³,⁴
- E-64 – [N-(trans-Epoxysuccinyl)-L-leucine 4-guanidinobutylamide] – cysteine proteases, e.g., calpain, papain, cathepsin B, and cathepsin L.
- EDTA – metalloproteases
- Leupeptin hemisulfate salt – both serine and cysteine proteases, e.g., plasmin, trypsin, papain, and cathepsin B.

Recommended Usage
One ml of the cocktail solution is recommended for the inhibition of endogenous enzymes equivalent to 1 mg of USP pancreatin. One bottle is recommended for the inhibition of proteases present in a maximum of 20 g of cell extract.

Note: Not all extracts contain the same levels of endogenous enzymes, and it may be necessary to adjust the volume of cocktail required.

Reagent
Supplied as a lyophilized powder

Precautions and Disclaimer
This product is for R&D use only, not for drug, household, or other uses. Please consult the Material Safety Data Sheet for information regarding hazards and safe handling practices.

Preparation Instructions
One bottle prepares 100 ml of cocktail solution. The powder dissolves quickly in a minimal volume of water or buffer, and may be prepared as a concentrate and diluted as needed.

The product is packaged in a 10 mL serum vial with a crimp cap. Dissolve the contents of the vial in 10 ml of water, then transfer to another container for dilution to 100 ml.

Storage/Stability
Store the lyophilized powder at −20 °C. The product, as supplied, is stable for 4 years when stored at −20 °C.

References

AP,NDH,PHC 02/06-1

Sigma brand products are sold through Sigma-Aldrich, Inc.
Sigma-Aldrich, Inc. warrants that its products conform to the information contained in this and other Sigma-Aldrich publications. Purchaser must determine the suitability of the product(s) for their particular use. Additional terms and conditions may apply. Please see reverse side of the invoice or packing slip.
APPENDIX E

Dilution series for Bio-Rad/Bradford assay and cytochrome c quantification

1. **For each sample** (e.g., yeast homogenate, cation-exchange decant, cation-exchange eluate),
   a. put a 100 uL aliquot in a 1.5 mL centrifuge tube. Add 900 uL phosphate buffer (with βME) and mix gently but thoroughly. Mark this tube "1/10".
   
b. Take 100 uL of your 1/10 dilution, transfer to a new 1.5 mL tube, and add 900 uL buffer as before. Mix gently but thoroughly. Mark this tube "1/100".
   
c. Repeat step (b) to generate dilutions of 1/1000, 1/10\(^4\), and 1/10\(^5\).

2. Centrifuge all samples to precipitate suspended debris.

3. To determine total protein and cytochrome c content, set up the following samples in a 96-well plate:
   - lysozyme dilutions for a standard curve, PLUS buffer, PLUS 40 uL Bio-Rad dye, as you did in week two (in triplicate)
   - a row for each sample: 20 uL undiluted and 20 uL of each dilution PLUS 140 uL of buffer PLUS 40 uL Bio-Rad dye
   - a second row for each sample: 200 uL undiluted and 200 uL of each dilution

For example:

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>assay blank plus buffer and dye</td>
<td>assay blank plus buffer and dye</td>
<td>assay blank plus buffer and dye</td>
<td>20 uL lysate plus buffer and dye</td>
<td>20 uL decant plus buffer and dye</td>
<td>20 uL eluate plus buffer and dye</td>
<td>200 uL lysate</td>
<td>200 uL decant</td>
<td>200 uL eluate</td>
</tr>
<tr>
<td>B</td>
<td>0.5 ug lysozyme plus buffer and dye</td>
<td>0.5 ug lysozyme plus buffer and dye</td>
<td>20 uL lysate 1/10 plus buffer and dye</td>
<td>20 uL decant 1/10 plus buffer and dye</td>
<td>20 uL eluate 1/10 plus buffer and dye</td>
<td>200 uL lysate 1/10</td>
<td>200 uL decant 1/10</td>
<td>200 uL eluate 1/10</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>1 ug lysozyme plus buffer and dye</td>
<td>1 ug lysozyme plus buffer and dye</td>
<td>20 uL lysate 1/100 plus buffer and dye</td>
<td>20 uL decant 1/100 plus buffer and dye</td>
<td>20 uL eluate 1/100 plus buffer and dye</td>
<td>200 uL lysate 1/100</td>
<td>200 uL decant 1/100</td>
<td>200 uL eluate 1/100</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>2 ug lysozyme plus buffer and dye</td>
<td>2 ug lysozyme plus buffer and dye</td>
<td>20 uL lysate 1/1000 plus buffer and dye</td>
<td>20 uL decant 1/1000 plus buffer and dye</td>
<td>20 uL eluate 1/1000 plus buffer and dye</td>
<td>200 uL lysate 1/1000</td>
<td>200 uL decant 1/1000</td>
<td>200 uL eluate 1/1000</td>
<td></td>
</tr>
<tr>
<td>E</td>
<td>3 ug lysozyme plus buffer and dye</td>
<td>3 ug lysozyme plus buffer and dye</td>
<td>20 uL lysate 1/10(^4) plus buffer and dye</td>
<td>20 uL decant 1/10(^4) plus buffer and dye</td>
<td>20 uL eluate 1/10(^4) plus buffer and dye</td>
<td>200 uL lysate 1/10(^4)</td>
<td>200 uL decant 1/10(^4)</td>
<td>200 uL eluate 1/10(^4)</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>4 ug lysozyme plus buffer and dye</td>
<td>4 ug lysozyme plus buffer and dye</td>
<td>20 uL lysate 1/10(^5) plus buffer and dye</td>
<td>20 uL decant 1/10(^5) plus buffer and dye</td>
<td>20 uL eluate 1/10(^5) plus buffer and dye</td>
<td>200 uL lysate 1/10(^5)</td>
<td>200 uL decant 1/10(^5)</td>
<td>200 uL eluate 1/10(^5)</td>
<td></td>
</tr>
</tbody>
</table>

6. Read your plate twice, once at 595 nm for the Bradford assay, and then once at 410 nm for the cytochrome c.
APPENDIX F

Ten Questions for week three

1) What are two characteristics of a proper buffer that, when not taken into consideration, are likely to affect protein stability during the purification of a protein?

2) Suppose that you have a purified sample of bovine cytochrome c in a test tube. The total volume is 1.0 mL. You remove 100 µL of this solution and add 400 µL of buffer. You then take the absorbance of the diluted sample and find it to be 0.105 at a wavelength of 410 nm. How many milligrams (mg) of bovine cytochrome c were in the original 1.0 mL sample?

3) What is the purpose of adding β-mercaptoethanol to the buffer when purifying cytochrome c?

4) In diluting a protein sample a student does the following. She first removes 900 µL of buffer from a stock solution using a P1000 pipetman and dispenses it into a labeled 1.5 mL microcentrifuge tube. She then removes 100 µL of the protein solution using the P1000 and adds it to the microcentrifuge tube. She caps the tube and mixes the solution. What lab sin did she commit?

5) Suppose that you need to measure the protein quantity in a 5.0 mL sample. You have already constructed a Bio-Rad protein standard curve with lysozyme as the standard. The equation of your best-fit line for the standard curve is $y = 0.0637 \times + 0.3257$. You remove 100 µL of sample and add 900 µL of buffer, and then use 20 µL of this dilution to find an absorbance of 0.455 at a wavelength of 595 nm. How many milligrams (mg) of protein were in the original 5.0 mL sample?

6) Yeast cytochrome c has a pI of approximately 9 and you used a cation-exchange resin to bind the protein during the purification. If a protein has a pI of 6.5, what type of resin would you use to bind it to help purify it?

7) Why don’t we use the more conveniently-disrupted organism Eschericia coli rather than the cell wall-enclosed Saccharomyces cerevisiae as a source from which to purify cytochrome c?

8) What two characteristics of blue dextran make it ideal for determining the void volume of a gel filtration column?

9) Imagine that using the Bio-Rad protein assay reagent you measure the absorbance of a solution containing exactly 10 µg of pure bovine cytochrome c. When you fit the absorbance to a well-constructed standard curve you find the apparent amount of cytochrome c in the sample is calculated to be 25 µg. What might explain such a large discrepancy between the actual value and that calculated from the standard curve?

10) Why is the use of a cation-exchange column so effective in purifying cytochrome c from yeast?
APPENDIX G

De-convoluting Gel Filtration Data

A wealth of information is hidden inside even a poorly resolved Gel Filtration peak profile. As we've discussed, a longer column will typically give greater resolution of proteins in a gel filtration experiment. One of the lab groups in 2005 had some technical problems and ended up with an unusually short column, and so I've used their data to illustrate how to extract good information from what initially looks like a completely uninformative peak profile.

Step One: The Raw Data
First, looking at the raw data (panel 1 below), both absorbance profiles seem to indicate that only one protein has eluted, though the broad peak of absorbance at 280 nm doesn't quite line up with the fractions showing maximal absorbance at 410 nm.

Step Two: Leveling the Field
To begin to extract the hidden information, I first rescaled the data so that the absorbance at 280 nm and the absorbance at 410 nm have the same range of intensities (panel 2). This is done by dividing all the $A_{280}$ values by the largest reading at 280, and separately dividing all the $A_{410}$ values by the maximal reading at 410. The resulting graph shows that there is a slightly higher hump, just barely visible in the raw $A_{280}$ data, that corresponds to the peak in absorbance at 410 nm.

Cytochrome c is a protein that contains aromatic amino acids as well as a heme group, so it absorbs light both at 280 nm and at 410 nm. Consequently, any fraction that contains cytochrome c should show absorbance at both wavelengths, in proportion to the amount of cytochrome c that's present. If only cytochrome c were present, then 100% of the $A_{280}$ and 100% of the $A_{410}$ would be attributable to cytochrome c. That's precisely what we're seeing where the blue line and the red line in panel 2 coincide -- at those points, all of the eluting protein is cytochrome c.

Step Three: Removing Interference
If the absorbance at 280 nm around $V_e = 40$ mL is due to the cytochrome c, then the $A_{280}$ to the left of the cytochrome c peak is probably due to carbonic anhydrase, and the $A_{280}$ to the right of the cytochrome c peak is probably due to aprotinin (since we know those are the only three proteins present). But because many fractions contain more than one protein we're not seeing three clean peaks in the $A_{280}$ profile. Remember, the raw $A_{280}$ values are the result of aromatic amino acid absorbance from all the proteins in solution, and therefore they are the arithmetic sum of the absorbances of each of the three protein species individually. If we could separate out the individual protein species, we could resolve -- the jargon term is "deconvolute" -- the messy $A_{280}$ profile into something cleaner.

Fortunately, we can do just that. As you know, the $A_{410}$ profile shows us where cytochrome c is. So what we're doing in this step is simply subtracting the cytochrome c
peak out from the mix, leaving the $A_{280}$ contributions of carbonic anhydrase and aprotinin. Up to this point, these had been pretty much hidden by the fact that the strong cytochrome c peak overlapped with them.

Computationally, this is done by making a new column of values of: (rescaled $A_{280}$) minus (rescaled $A_{410}$). The result is shown in panel 3 -- voila! -- there really ARE three clean peaks here!

*Step Four: Dressing it up*
Finally, I simply prettied up the results of step 3 to make the three peaks clearer (panel 4).

You can easily do the same thing with your data, and remove all ambiguity about where exactly each of the three standards is eluting. This ought to make your gel filtration standard curve more accurate.
Step 1: The raw data

Step 2: rescaled absorbance data

Step 3: the hidden peaks emerge...

Step 4: elution profiles of the three standards
APPENDIX H

SDS-PAGE FOR DETERMINATION OF MOLECULAR WEIGHT OF PROTEINS
(LAEMMLI SYSTEM)


REDUCING SAMPLE BUFFER:

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Volume</th>
<th>Conc. In 2X stock</th>
<th>Final conc.*</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.50M Tris-HCl, pH 6.8</td>
<td>6.4mL</td>
<td>0.125M</td>
<td>0.0625M</td>
</tr>
<tr>
<td>10% SDS</td>
<td>10.0mL</td>
<td>4%</td>
<td>2%</td>
</tr>
<tr>
<td>Glycerol</td>
<td>5.0mL</td>
<td>20%</td>
<td>10%</td>
</tr>
<tr>
<td>0.5% Bromphenol Blue</td>
<td>1.0mL</td>
<td>0.02%</td>
<td>0.01%</td>
</tr>
<tr>
<td>2-Mercaptoethanol</td>
<td>2.5mL</td>
<td>10%</td>
<td>5%</td>
</tr>
</tbody>
</table>

* Diluted with equal parts of sample or water

Store in brown bottle at 4°C. SDS may precipitate, but will go back into solution at room temperature. Boil samples 3 minutes before loading.

RUNNING BUFFER:

for one liter of 1X Tris-glycine:

- 3.0 g Trizma base (final concentration 0.025 M)
- 14.4 g Glycine (final concentration 0.19 M)

fill to 1000 mL with nanopure water
add 10 mL 10% SDS (final concentration 0.1%)

Store at room temperature. Use ~300 mL per gel run.

RUNNING THE GEL

The Bio-Rad mini-gel rigs take ~110 V-hrs to run the dye front to the bottom of gel in a 10% gel.

STAINING:

This stain may be mixed as a stock. It may be reused multiple times. Place the gel in the Staining Solution for 30-60 minutes. If you stain for a longer period of time, destaining will take longer as well.

Stain Stock:

- 2.00g Coomassie Blue R-250
  fill to 200mL with nanopure water
  stir and filter using buchner funnel with aspiration

Staining Solution:

- 62.5mL Stain Stock
- 250mL methanol
- 50mL acetic acid
  fill to 500mL with nanopure water
DESTAINING:
The procedure described above will stain the entire gel blue. To see protein bands, Coomassie that is not associated with proteins -- the "background" -- must be removed. During destaining, the background will lose dye before the protein bands, but incubating too long in Destain I can result in loss of too much stain, so proceed carefully.

To Destain:
1. Place the gel in Destain I with gentle agitation until the protein bands become clearly distinguishable from the background.
2. Destain the background with several changes of Destain II. Solution should be changed when the solution and the gel show the same intensity of blue color. The background should be almost completely clear after the destaining procedure.

To destain heavily stained gels or to destain the protein bands themselves, place the gel in Super Duper Destain.

Destaining can also be achieved by heating gel(s) in water in a microwave oven ~5 minutes (or until nearly boiling), pouring water off and repeating as many times as needed to achieve desired level of background stain.

Destain I:
- 500mL methanol
- 100mL acetic acid
  fill to 1.0L with nanopure water

Destain II:
- 50mL methanol
- 70mL acetic acid
  fill to 1.0L with nanopure water

Super Duper Destain:
- 75mL acetic acid
- 25mL acetone
APPENDIX I

Procedure for using the SQ-2800 Single Beam Scanning UV/Visible Spectrophotometer for Kinetic Experiments in Biochemistry Lab.

*Note that the spectrophotometer requires 25 minutes of warm-up and calibration before any samples can be measured.*

1. Turn on the spectrophotometer

2. After the 15 minute warm up period, select [yes] to calibrate.


5. To select units of measurement, press [F2] and select Abs, then press [enter].

6. To choose a wavelength, press [Set $\lambda$] and enter 475 nm.

7. In a 3 mL plastic cuvette, place all solutions *EXCEPT* the tyrosinase (all solutions should be at room temperature except the tyrosinase, which should be kept on ice). Wipe down the cuvette with a Kimwipe.

8. Place the cuvette into the spectrophotometer and press [0 Abs / 100% T] to zero the instrument.

9. Add tyrosinase to the solution, cover the cuvette with parafilm, invert three times *(DO NOT SHAKE)*, wipe with a Kimwipe, and quickly replace in the spectrophotometer. Press [Start].

10. When the time-scan stops and the spec has beeped 3 times, press [F3] and enter the start time and stop time of the linear region of the graph, pressing [Enter] after each command. You will also be prompted to enter a “factor”, at which you should enter your “magic number” and press [Enter].

11. The I.U. readout on the right side of the screen is the "factor" x ($\Delta A$/min).

12. The scale can be changed by the left and right arrow keys for the x-axis and the up and down arrow keys for the y-axis.
APPENDIX J

Independently-designed Projects on Analysis of Tyrosinase Activity

- by Monday, Nov. 12th, at the latest, your group should talk to your instructor about your plan for the next two weeks' lab periods. If you do not talk to us by Monday at the end of class, we can't guarantee that the reagents and equipment you need will be in place; even Monday may be too late depending on your needs. If you delay and the toys you need aren't in place in lab that week, you will still be expected to get your investigation done by December 6th.
- the results of your investigation will be presented at a combined poster session (for all three sections of Biochemistry) at 6 pm on Dec. 6th. You should prepare a brief (~3 minute) talk in which you can summarize your goals and your findings to your classmates and your instructor. You'll get a chance to check out the other groups' work and evaluate their results and presentations.
- your poster presentation should describe your experimental context, the hypothesis you were trying to evaluate, your results, and your interpretation of them. Your interpretation should include an attempt to make structure/function inferences about tyrosinase on the basis of your data and relevant published data. More guidelines and recommendations are available online at http://www.denison.edu/~kuhlman/courses/poster.info.html
- at lab check-out (or before, as specified by your instructor), your group should turn in an abstract of your poster and a summary data table, providing

<table>
<thead>
<tr>
<th></th>
<th>$K_M$</th>
<th>$V_{max}$</th>
<th>$K_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>control sample(s)</td>
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<td>not applicable</td>
</tr>
<tr>
<td>(multiple values if</td>
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<td>day to day)</td>
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<td>benzoate</td>
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<td>your conditions/</td>
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<td>inhibitors</td>
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(all of these values should be reported with measures of statistical significance)

- where relevant, posters that also present estimates of $k_{cat}$ and/or the specificity constant for your reactions will be favorably received.
APPENDIX K

Pipetter Accuracy Report Form

Summary
Pipetter type: ____________ (e.g., P20)
Pipetter serial number: ____________
Fractional error at low volume (95% CI/target) ____________
Fractional error at low volume (95% CI/target) ____________
Fractional error at low volume (95% CI/target) ____________

Raw data
Low volume target (µL) ____________
Mass reading #1 (mg) ____________
Mass reading #2 (mg) ____________
Mass reading #3 (mg) ____________
Mid-range volume target (µL) ____________
Mass reading #1 (mg) ____________
Mass reading #2 (mg) ____________
Mass reading #3 (mg) ____________
High volume target (µL) ____________
Mass reading #1 (mg) ____________
Mass reading #2 (mg) ____________
Mass reading #3 (mg) ____________

*   **  ***  ****  *****  ****  ***  **  *

Summary
Pipetter type: ____________ (e.g., P20)
Pipetter serial number: ____________
Fractional error at low volume (95% CI/target) ____________
Fractional error at low volume (95% CI/target) ____________
Fractional error at low volume (95% CI/target) ____________

Raw data
Low volume target (µL) ____________
Mass reading #1 (mg) ____________
Mass reading #2 (mg) ____________
Mass reading #3 (mg) ____________
Mid-range volume target (µL) ____________
Mass reading #1 (mg) ____________
Mass reading #2 (mg) ____________
Mass reading #3 (mg) ____________
High volume target (µL) ____________
Mass reading #1 (mg) ____________
Mass reading #2 (mg) ____________
Mass reading #3 (mg) ____________