

College of the Canyons: Introduction to Biotechnology: Custom Labs





- Lab notebooks are used to document all ideas stemming from lab • research (either academic or industrial).
- Lab notebooks provide a daily record of procedures performed. data gathered, and determined results for each lab.
- They also help provide legal proof of research activities (this would be especially important for cutting edge research).
- Lab notebooks are to be completed as the lab is being performed.
- In addition to documenting key lab activities, this lab book will serve as a tool to help assess student ability to write and think critically about lab exercises. Subsequently, the content and format is slightly novel to facilitate the educational process.
- The notebooks will be collected and graded periodically throughout the semester (see syllabus).

For more information on the College of the Canyons' Introduction to Biotechnology Course, contact, Jim Wolf, Professor of Biology/Biotechnology at (661)362-3092 or email: jim.wolf@canyons.edu

The following lab protocol can be reproduced for educational purposes only. It has been developed by Jim Wolf, and/or those individuals or agencies mentioned in the references.

I. Background:

Laboratory Notebook

In a biotechnology course, one of the primary objectives is to make the laboratory setting as realistic as possible. In research laboratories (either academic or industrial) the lab notebook is <u>the</u> source for all of the ideas stemming from research efforts. While the exact nature of a notebook is subject to lab specifics, individual idiosyncrasies, and familiarity with the subject matter, all good lab books have certain things in common. The following is a list of these attributes. Note that these attributes apply to notebooks in a research setting. Since the academic setting is substantially different from the research setting (i.e. in research, experiments are often repeated dozens to hundreds of times, while in a education setting, it is rare to repeat an experiment more than once) you should be aware of these differences. You should balance your lab notebook with an additional three ring binder-full of Standard Operating Procedures (SOPs) handed out in lab. The lab notebook and SOP binder *may (at my discretion)* be used during exams or quizzes or as a resource for reports. Consider the following "education specific" lab book (as seen on last pages of this handout.)

Education Lab book Idiosyncrasies:

1. Your lab book and SOP binder shall contain only materials relevant to lab. <u>You must</u> leave your lab notebook in class at all times, and you should keep you SOP binder with you at all times. No lecture notes, Xeroxed lecture related materials or other material clearly related to lecture shall be permitted in these books.

2. The detail in the lab book is largely up to your discretion (although a general format should be followed). Grading emphasis will be placed on completeness (see below.) Additional materials, while welcome; will not result in extra credit.

3. Strive to write the notebook in the following listed format, and you can use SOP's to provide additional detail (especially in the materials and methods section.)

4. Your lab notebook will have a spiral binding or solid spine (no loose leaf) and will have **carbon copy pages that are easily removed.**

5. FINAL NOTE: KEEP YOUR LAB NOTEBOOK CURRENT! WRITE YOU INTRODUCTION TO THE LAB DURING THE LAB AND ENTER ALL DATA DIRECTLY INTO THE LAB NOTEBOOK. WRITE YOUR CONCLUSIONS AT THE END OF THE FINAL LAB PERIOD. AGAIN... KEEP CURRENT, AS YOU WILL BE SUBSTANTIALY PENALIZED IF YOU LET YOUR NOTEBOOK LAPSE. Typical Components of A lab Note Book:

These criteria will be used in grading your notebook.

- 1. <u>In front of the notebook</u>, put identifying information including: Name, course, room number, Professor's name (PH # and other contact information as well.)
- 2. <u>Table of contents</u>: Keep Current!
- 3. <u>Title</u>, date, and room # and/or collaborating signatures/name of lab partners should be on each page of the lab notebook.
- 4. <u>Introduction</u>: For each project includes a brief introduction; 4-6 sentences per paragraphs and 1-2 paragraphs per subject. .

- 5. <u>Materials and Methods:</u> In many cases these points will be covered in your SOP. Be sure to reference the SOP in this portion of the lab notebook. Be specific in your reference of the SOP (i.e. SOP # 3, HPLC lab, pages 3-5.) and add any addenda noted in lab lecture. <u>This section is at most 1/2 a page long.</u> So, again, do not write out the entire materials and methods section. Just reference the SOP, and any addenda mentioned in the pre-lab lecture. The included example at the end of this lab included a M and M section FYI only.
- 6. <u>Data and results</u>: Collected directly from experiments, this section should include, but not necessarily be limited to: tables, graphs, procedural details, instruments used, calculations, computer printouts and gel images.
- 7. <u>Conclusion</u>: Include a brief interpretation <u>of all</u> relevant experimental outcomes. In addition you should answer the final questions listed at the end of the lab handout. This section will be from 1-4 paragraphs long and remember to answer the questions in essay format (no listing or numbering of answers.)
- **II. Format:** (The format of the lab notebook should be as follows):

Introduction: A 3-5 sentence introduction should be sufficient. Remember to BOTH introduce the subject in a general sense, and to say what the specific lab was addressing (this sentence is often called the Hypothesis being tested). Ex: Photosynthesis is the reaction where by plants utilize carbon dioxide and water, in the presence of light and chlorophyll to produce sugars and oxygen. (Put in a 3-5 extra introductory sentences after the first sentence). The purpose of this lab was to investigate the relationship of light intensity to the rates of photosynthesis in the aquatic plant <u>Elodea.</u> (Hypothesis being tested sentence used to close out the introduction).

<u>Materials and Methods</u>: Do not bother writing out all of the procedures in your notebook. Simply allude to the SOP's: "i.e. SOP's, Transformation Protocols, pgs 3-5," unless changes are made. In that case an addenda is necessary.

Data/Results: In this section you state what data you gathered. This may be presented textually, in a graph, table, or diagram. What ever method you choose, again ensure that you are both clear and to the point. Actual numbers, formulas, calculations, graphs or other results should be also included in this section. Be sure to fully label and graphs, gel images, or other data with relevant information. Avoid *explaining* your results or data in this section.

Conclusion: In the conclusion section comment on whether or not the experiment was successful, how it may be improved, and some closing comments on what was deduced. At the end of every lab handout there are several questions for you to answer. These questions should provoke thought and understanding of the labs. Answers to these questions are to be addressed in the conclusion section of the lab notebook and will be graded when the lab notebooks are turned in (see syllabus). Please note that the questions should not be answered directly but should be worded in such a way that the question is answered in your own words. Example:

Question: Why is polarity so important in the HPLC lab?

Entry in lab notebook: Separation of the Kool-Aid dye occurred as a direct result of polarity. The more polar the solvent used, the less of the non-polar dye was pulled through the HPLC column. Therefore, polarity served as the main basis for separation of the different dyes used in grape Kool-Aid. (NOT: Polarity was important in the HPLC lab because...)

And finally: KEEP CURRENT, KEEP CURRENT, KEEP CURRENT.....

III. <u>Example</u>:

Following is an example from a previous BioSci 230 student's lab notebook: This student received full credit for this lab notebook entry. Again, recall that the material and methods section in your lab notebook should be much briefer!

EXP. NUMBER	EXPERIMENT/SUBJECT		DAIT	
Г	Transformation		3-14-01	
NAME	~	LAB PARTNER	LOCKEP Drow No. 15 33 2	
Shane	Ramey	Ron Moat	- Biotech	230

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I. Intro! Cello are transformed when new DNA is insertis and thus changes that cell's descriptions and those of its offlepsing. Developmention revely happens is notice, but accessionably a cell dappenets to up and use a piece of DNA ther is adhift in the cell's environment. Usually, Rowene, when this dappens it is a falal event for the cell size the foreign DNA could change it cellibre to summer in mediately. We to Richard, though, have beened in the last thity years bent Auccessfully transform boateria and year cells.

In this lab <u>Cacherichia celi</u> (<u>C. celi</u>, a commen battenium in the human large intertine) is transformed from an ampleillin-servicine <u>E. celi</u> to sun ampleillin-relietant cells.

II <u>Materials (Mittled</u>s: Smithelly the <u>E. coli</u> allo must be used and so that they will mar neadily accept the foreign DNA. This is performed by adding <u>E. coli</u> colonies to 50mM ico-cold CaCL2 and putting on ico for at lease I minute. Next, the plasmid certaining anglicillin-resultant DNA is introduced to the purposed detection. This is accomplished by adding the plasmid to the locateria and coing for 18 minutes. Then the plasmid must be made to extend allow to the locateria and coing for 18 minutes. Then the plasmid must be made to extend allow to the locateria and coing for 18 minutes. Then the plasmid must be made to extend allow to reapin them publicly from the iso both to a 43°C water both for 90 ascerde. It allo are allowed to reapin their stampt buy adding them is hath and incubiting to a 37°C with both for 5 minutes. The allow on pelid made with and without added antibiotic and incubated of 37°C for 24 hours. Cire 50P's, Transformation Charetic Crainsering. Protected).

III. Data Resulto:





EXP. NUMBER	Transformatio	N	UATE 3-14	1-01	21
Shane	Ramey	LAB PARTNER Ron Moat	LOCKER/DESK NO.	COURSE & SECTION NO. Biotech 23	Ø

is alguna bimady - but bimady + alt tot may reianegaua lb by Un 001 parappened to two types of midia, are with and one without ampicillin. She plate are then incubated and evaluated. She planned will as as expected - they arow well on the amphellin and a measure ministras stale at the all to warp the above the stale and low of arouth is encider on the ang-free plate. The + plasmid collo also arow into a nice law on the amp-file glate. nowener, some allo form colonies on the plate with ampicultin. a have does not archist, but growth is orider. This growth electrotes the fact that some of the original tacteria was successfully transformed with the plasmid. By counting it ang-resistant colonies on it glats is determined that 780 of these colonies are growing. With this data is in further determined that 78,000 colonies use transformed per microop an of plasmid DNA. ataly the amp- resistant cells did not grow a dense law on the plate al containing the antibiotic, inducation that a loss percentury of the cells were actually for an interest bimally call the to the to the fact that the finally in the first of brital and results in low amp-resistant cell riceld, SIGNATURE James mall 3-14-01 3-14-01

NOTE: INSERT BACK COVER UNDER COPY SHEET BEFORE WRITING