BIOD21H
Molecular Biology Laboratory I
Host, Vectors and Cloning
Course outline Summer 2013

Instructor:
Dr. Shelley Brunt
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PHILOSOPHY: Molecular techniques have evolved rapidly over the last 20 years, however, the fundamental principles remain unchanged. This course combines the opportunity to perform commonly used older techniques that provide the foundation for all the newly developed techniques while also introducing the newer techniques. Significant attention is paid to the principles behind the techniques which will allow you to gain familiarity with the techniques and most importantly gain insight into the theory behind the techniques to aid in troubleshooting.

The goal of this course is to provide you with the tools and the background to go forward to find a technical position in a research, biotechnology or pharmaceutical laboratory or go forward into graduate school in any research field that requires the use of molecular biology techniques. In today’s research that involves all biology fields. Therefore the emphasis is on the laboratory component of this course. We attempt to mimic the experience of a fourth year research project.

I encourage you to ask questions during lecture. If you have a question about the material, whether it is lecture or laboratory material I encourage you to talk to me during the laboratories as I will be around for much of the lab period or to visit me in my office. It is not feasible to give detailed answers to questions regarding material covered in the lecture or laboratory via email. Therefore I have an open door policy and in addition I hold scheduled office hours

Please use E-mail (UTORONTO ACCOUNT ONLY) when it is critical you get in touch with me, and you are unable to see me in person

Office hours: Remember I have an open door policy in addition to set office hours when I will definitely be in my office
  Tuesday 10 to 11 and 3 to 4 pm
  Wednesday 4 to 5 pm
  Thursday 11am to noon
  or email me and we can arrange an alternative time
**Accessibility:**
Students with diverse learning styles and needs are welcome in this course. In particular, if you have a disability/health consideration that may require accommodations, please feel free to approach me and/or the AccessAbility Services Office as soon as possible. I will work with you and AccessAbility Services to ensure you can achieve your learning goals in this course. Enquiries are confidential. The UTSC AccessAbility Services staff (located in S302) are available by appointment to assess specific needs, provide referrals and arrange appropriate accommodations (416) 287-7560 or ability@utsc.utoronto.ca.

**Academic integrity/plagiarism**

Academic integrity is essential to the pursuit of learning and scholarship in a university, and to ensuring that a degree from the University of Toronto is a strong signal of each student’s individual academic achievement. As a result, the University treats cases of cheating and plagiarism very seriously. The University of Toronto’s Code of Behaviour on Academic Matters ([http://www.governingcouncil.utoronto.ca/policies/behaveac.htm](http://www.governingcouncil.utoronto.ca/policies/behaveac.htm)) outlines the behaviours that constitute academic dishonesty and the processes for addressing academic offences. Potential offences include, but are not limited to:

(source: [http://ctl.utsc.utoronto.ca/home/integrity](http://ctl.utsc.utoronto.ca/home/integrity))

**In papers and assignments:**

- Using someone else’s ideas or words without appropriate acknowledgement.
- Submitting your own work in more than one course without the permission of the instructor.
- Making up sources or facts.
- Obtaining or providing unauthorized assistance on any assignment.

**On tests and exams:**

- Using or possessing unauthorized aids
- Looking at someone else’s answers during an exam or test.
- Misrepresenting your identity.

**In academic work:**

- Falsifying institutional documents or grades.
- Falsifying or altering any documentation required by the University, including (but not limited to) doctor’s notes.

All suspected cases of academic dishonesty will be investigated following procedures outlined in the Code of Behaviour on Academic Matters. If you have questions or concerns about what constitutes appropriate academic behaviour or appropriate research and citation methods, you are expected to seek out additional information on academic integrity from myself as your instructor or from other institutional resources (see [http://www.utoronto.ca/academicintegrity/resourcesforstudents.html](http://www.utoronto.ca/academicintegrity/resourcesforstudents.html)).

**Text and required materials:**

**Text:** Introduction to Biotechnology
Thieman and Palladino third edition 2013
Pearson

**Lab manual/handouts:**
- The lab manual will be posted on-line each lab will be posted at least a week in advance.
- Any additional laboratory material including reference material will be posted on blackboard.

**Lecture material/ data files** will be provided for the lecture. Please check blackboard for any postings of lecture material.

Lecture material will come in two forms. A more detailed PDF file and a more bare bones powerpoint. The word file will help you supplement you lecture material. You are responsible for what I cover in lecture and the related material in the posted the PDF files. Use the material I presented in lecture as a guide to what I emphasize.

**Students also require:**

a) a lab coat (no exceptions)

b) at least one permanent marker with a fine tip (Black). The best choice is a Sanford “Sharpie” fine point or extra fine point. (I would suggest you buy yourself two markers (a fine point and extra fine point))

c) a book for recording your work (your laboratory log book). This book can be hard covered and bound, or a binder with paper added.

**Course schedule:**

Class meets 2 days a week: Wednesday noon to 4 pm (3 to 4 pm is used when I can’t complete lecture during lab periods) and Thursday 2 to 5 pm. I lecture at different times during this seven hours. On average there is two hours of lecture per week. Some
Attendance is mandatory. You will be carrying out a laboratory exercise every Wednesday and Thursday, starting May 8, 2013. Missing a laboratory will be equivalent to missing a midterm. Therefore, the procedure for missed laboratories/quizzes/term tests is as follows:

A UTSC medical certificate filled in by a Medical Doctor will be required. Lab work cannot be made up, but should you provide a proper medical note I will provide a makeup assignment for the missed laboratory.

If you miss the term test contact me within 48 Hr. To write a makeup a UTSC medical note is required. However, it will not be the same exam.

A reminder, if you miss the final exam, I cannot give a makeup exam. You must deal with the registrar, fill out the appropriate forms to defer the exam.

- If you have one unexcused absence you will forfeit all of your participation grade and grades associated with that lab
- Two unexcused absences leads to a loss of all laboratory grades associated with those days including the lab report required
- More than two unexcused absences and you will forfeit all grades associated with the laboratory, which means you will not pass the course

**Lectures:**

BIOD21H is a laboratory course. The lecture material covered will relate to the laboratory techniques carried out throughout the course. It will include in depth explanations of methodologies, the theory behind the methodology, and discussions on how to apply the methodology to studies in molecular genetics. The time and length of lecture will vary week to week. The course outline gives you a general idea of the length of each lecture, but exact times may vary. On a weekly basis I will give you an update as to any changes in the schedule for the following week. In this way you will be able to keep ahead in your reading, and will be prepared for the laboratory/lecture in the upcoming week. Lecture is an ideal time to ask questions if you have a question likely another student has the same question.

**Laboratories:**

As I have stated above the emphasis of this course is on the laboratory, and the lecture complements the laboratory. Think of this course as a supervised study course. It is your responsibility to carry out the experiments correctly and within the time frame of the laboratory schedule. You will be graded on how you work in the laboratory, whether you are prepared, and how well you keep a log of your experiments, detailing exactly what you did and what you observed (diagrams/tables are excellent additions). In research you MUST have excellent notes on your daily work,
as REPRODUCIBLE DATA is an absolute must. Your mental mistakes and oversights will be reflective of how well you prepared and will be considered when you are graded. Therefore, simply showing up to the lab will not ensure you any success in this course. You must arrive well informed and prepared to carry out the laboratory exercises. Since each week builds on the previous week of work, you will often be preparing the materials you need for subsequent experiments. The intent of this course is to introduce you to how you would work within a research, industry or government laboratory, where you are producing materials you need for subsequent experiments.

Log books for laboratory

Every class you should arrive with an introduction (paragraph) written in your log book (pages should be numbered) that describes in general what your goals are for the day. This introduction will be followed by a flow chart/outline that will diagrammatically describe how you will carry out the procedures within the exercises. Include all relevant information (for example incubation times, volumes to use). If two experiments are ongoing then indicate within this flow chart when you might be carrying out certain steps of the various exercises. You should be able to use at the flow chart/outline to carry out the experiment without constant referral to your manual. This will ensure you are prepared for the laboratory and will help you formulate any questions before starting your work.

This preparation is required and will be checked each day. I will record whether the preparation was done and to what level (unacceptable/acceptable/good/excellent).

During the course of the experiment you will record a detailed log of what you did. Each step you carry out will be written down (use past tense). Indicate volumes used, time of incubation (write the actual times). Describe exactly what you did and what you observed. If a step was carried out by your partner and not you indicate that in your log book.

For example:

Thursday Sept 23:
1) An isolated white colony from plate number 1, containing E. coli strain D21-1, was aseptically transferred to a 5 ml aliquot of sterile LB containing 100 μg/ml of ampicillin. This tube was assigned the number 1.
2) The culture was incubated at 37°C with constant shaking in a water bath shaker. The cultures were grown overnight (if you know the exact time indicate it) and placed at 4°C the next morning by the teaching technician.

Wed Sept 29:
1) Culture tube #1 was removed from the 4°C fridge. The cells had sedimented to the bottom of the tube. The pellet was resuspended by gently tapping the tube. The cell pellet dispersed and the culture was turbid.
2) One ml of E.coli strain D21-1 (XL1B cells containing the plasmid) was aseptically removed from the 5 ml overnight culture and placed in a 1.5 ml microfuge tube. The remaining culture was put back into the fridge to keep as a source of culture if needed.
3) The sample was centrifuged at 1000xg for 5 min at RT (room temperature). A small cream coloured pellet was observed at the bottom of the tube. The supernatant was clear.
4) The spent media was poured off and excess media drained from the tube by inverting the tube on a paper towel for 1 min.
5) 100 μl of solution 1 was added to the cell pellet and the pellet was resuspended by agitation using a vortex at speed 6. The sample was left on ice for 5 min. 2:20 pm-2:25 pm.
6) While I waited for the sample to resuspend I labelled my tubes for subsequent steps in the procedure.

another example

1) John prepared the DNA samples for Eco RI digestion, for specific details see John’s log book. I prepared the samples for Hind III digestion. See table below.

- At the end of the experiment there should be a summary of what you did, what you observed, and how this relates to the next experiment and DATA analysis
- an inventory table will also be kept at the back of your log book (explanation in class)

Your log book will be checked at some point during the day, usually in the first hour of lab. I or the TA will initial various pages. At the end of the year you will be assigned a final grade for your preparation, lab performance and record keeping (see mark breakdown). You will be required to keep you book up to date. You book will be graded on a regular basis.

The exercises will be carried out in pairs. Remember the success you have will depend on each of you being will prepared.

Each pair will be provided with in:
1) Sterile pipette tips for micropipettors
   - a box (blue) for a large volume pipettor (100 μl-1000 μl)
   - a box for small to mid range micropipettors (1μl to 200 μl)
2) a can containing sterile 1.5 ml microcentrifuge tubes
3) a bag of disposable gloves for each student (the size given to you will be determined in the first week). There will be enough gloves to last you the entire course. If you however use more than three pairs a day, you will run out. If this happens you will be required to buy any extra gloves you require. Going without gloves when they are required is not an option. This will introduce you to the cost of research. Extra gloves
will cost 50¢ a pair.

4) racks for tubes
5) sectioned box for microcentrifuge tube storage (2 boxes each pair).
5) you will be given a variety of solutions that you will keep in your locker, fridge or freezer over the course of the labs.
   • It is your responsibility to keep track of your supplies and tools

per pair of students you will be provided with a set of micropipettors. These will be numbered with your group number and placed in zip lock bag or plastic container. These are your responsibility for the term. You are responsible for properly using, and storing your micropipettors.

You will be responsible for returning the empty pipet tip boxes and picking up a new box.
Therefore it is up to you to properly look after your laboratory tools. Responsible use of your supplies and equipment is critical to obtaining good results in a research setting. **Misuse of equipment is very costly and will not be tolerated.**

**Grade Breakdown**

**Midterm** (includes lecture and lab (a minimum of 10% is directly lab/applied)
includes (In class Wed June 13, 2012, 1: to 3 pm) 19%

**Final Exam** (minimum of 20% is lab or applied) 33%
(cumulative, for applied): lab and lecture (theory based on material since midterm)

**Lab performance** 11%
preparation 3%
technical performance 3%
laboratory log including data analysis and summaries 5%

**Small assignments/presentations /data analysis/summaries/reflective practice/concept map. These may take place in both the lab and lecture** 8%

**Quizzes (4 x 1%)** 4%

**Assignments (3)** 25%
1) bioinformatics (6.5%)

2) formal lab report #1 (9%)

3) formal lab report #2 (9.5%)

*The content required for each assignment will be explained during the appropriate class. Assignments will be considered late if they are not handed in at the beginning of the lab on the due date. Late lab reports will not be accepted more than two days late with a 10% penalty a day. If you have an issue getting your lab report completed on time you will have to speak to Dr. Brunt, preferably before the lab is due. All labs must be submitted electronically to Turnitin and to the TA as well as in hard copy. Each page must have your name as a header and you must initial each page.

"Normally, students will be required to submit their course essays to Turnitin.com for a review of textual similarity and detection of possible plagiarism. In doing so, students will allow their essays to be included as source documents in the Turnitin.com reference database, where they will be used solely for the purpose of detecting plagiarism. The terms that apply to the University's use of the Turnitin.com service are described on the Turnitin.com website".

If you wish to opt out of turnitin you must state this to Dr. Brunt in writing and submit and electron copy to Dr. Brunt

BIOD21 2011 Lab assignments and quizzes
Assignments (3) (24.5%)
1. Computational analyses of amino acid sequences provided to you (6.5%)
   - introduced week 3 and finished outside class. You will be asked to analyze and compare two protein sequences provided to you.
   Due Thursday June 6, 2013

2. Construction of cDNA library and cloning and characterization of cDNA clones
   - week 1 to 7 (9%)
   due Thursday July 18 2013

3. Construction of a genomic library/genomic southern and cloning and characterization of genomic clones (9.5%)
   - weeks 5-12
   due Tuesday August 6, 2013

You may be asked to hand in calculations or graphs for certain labs which will be
graded as Pass/Fail and will contribute to laboratory log data analysis grade.

Your log books will be looked at throughout the year. If you are not keeping up you mark will reflect this. The book will be collected on the day you hand in your final lab report just to look at the analysis of the last week of data. You can not fix missing material from previous checks.

**Quizzes (4x1%)**

1. Thursday May 16, 2013 (Growth, plasmid isolation restriction digests and gel electrophoresis)

2. Wednesday June 5, 2013 (cDNA library construction, probe preparation, screening of library)

3. Wednesday June 27, 2013 (Southern blot, PCR, yeast genomic DNA isolation)

4. Wednesday July 17, 2013 (Construction and screening of genomic libraries, characterization of genomic clones, RNA, RTPCR if done)

### Lecture and Laboratory Schedule for BIOD21H

<table>
<thead>
<tr>
<th>Date</th>
<th>Quiz</th>
<th>Lecture: Schedule is tentative</th>
<th>Laboratory exercise</th>
<th>assignment</th>
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<tbody>
<tr>
<td>May 8</td>
<td>-----</td>
<td>Lecture 12 to 2:30 pm Brief outline of course: first lecture Topic 1 and topic 2 on bacterial growth:</td>
<td><strong>Appendix week 1A/1B Exercise 2-inoculation of Individual colonies into LB/amp broth</strong></td>
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| May 10 | Introductory laboratory  
|        | - evaluate the results from yesterday |
|        | **Appendix week 1B:**  
|        | - completion of exercise 2.  
|        | - the completion of exercise 2 provides the cultures for Week 2 plasmid prep  
|        | Appendix week 1A |
| May 15 | **Start Topic 3:**  
|        | vectors used  
|        | In molecular cloning  
|        | Lecture time 3 to 4 pm  
|        | **Appendix week 2A:**  
|        | exercise 3- isolation of plasmid DNA (vector [pB] only and plasmid containing *Achlya* insert cDNA using rapid alkaline lysis  
|        | exercise 4- quantification of plasmid DNA using gel electrophoresis: digest set up (part 1) |
| May 16 | **Quiz 1**  
|        | **Topic 3:** vectors used in molecular cloning  
|        | Topic 4: DNA analysis: restriction endonucleases, gel electrophoresis 3:30 to 4:30 pm  
|        | **Appendix week 2B:**  
|        | exercise 4 (part 2)- load gel as soon as possible-no later the 2:45 pm  
|        | exercise 5 (part 1)- restriction analysis of pB and cDNA clone (part 1)  
|        | REFLECTIVE PRACTICE (either today or tomorrow) |
| May 22 | **topic 4**  
|        | Bioinformatics lecture followed by  
|        | **Appendix week 3A:**  
|        | exercise 5 (part 2)-load gels  
<p>|        | - data interpretation for |</p>
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<tr>
<th>Date</th>
<th>Topic</th>
<th>Exercises</th>
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<tbody>
<tr>
<td>May 23</td>
<td>data analysis and figure legend exercise and introduction of concept maps using sequence analysis</td>
<td>week2 class discussion come with your data -figure legend example/group discussion</td>
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<td>May 29</td>
<td>Topic 5: basic cloning Approx 2:30 to 3:30 pm Hand in a concept map of your Bioinformatics assignment</td>
<td>Appendix week 3B: exercise 6 - programs for computer analysis of nucleotide and amino acid sequences (computer lab) TBA</td>
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<td>May 30</td>
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<td>Appendix week 4A: exercise 7- transformation of XL1B cells with cDNA clone using rapid method (construction of cDNA plasmid library using Achlya cDNA)</td>
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<td>June 5</td>
<td>Quiz 2 topic 6: cDNA cloning 3 to 4 pm</td>
<td>Appendix week 4B: exercise 8 part 1- Screening of a plasmid library: colony lifts</td>
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<td>June 6</td>
<td>Discuss how to write an abstract</td>
<td>Appendix week 5A: exercise 9-random primer labelling using DIG system exercise 10 part 1-colony hybridization exercise A:part 1Yeast DNA isolation Exercise B: part 1 <em>Drosophila</em> DNA isolation</td>
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<td>Appendix week 5B: exercise 10- part 2 completion of colony hybridization exercise 11- enhanced Bioinformatic assignment due</td>
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<tr>
<td>Date</td>
<td>Appendix week 6A</td>
<td>Appendix week 6B</td>
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<td>June 12</td>
<td>exercise 12 part 1- isolation of plasmid DNA from putative cDNA: inoculation into LB/amp</td>
<td>Exercise 12 part 2 plasmid isolation and digests Exercise 13, part 2 precipitation of genomic DNA</td>
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<td>June 13</td>
<td>Quiz 3</td>
<td><strong>Topic 7:</strong> Southern and PCR Approx 2 pm to 3:30 Discuss the writing of a materials and methods</td>
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<td>June 26</td>
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<td>July 3</td>
<td><strong>Topic 7 complete</strong></td>
<td><strong>Appendix week 8A:</strong> exercise 15, part 4: construction of genomic library continued: transformation of <em>E. coli</em> XL-1 Blue with recombinant vectors containing genomic DNA exercise 13, part 5: Southern blot continued: prehybridization and hybridization</td>
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<td><em>Exercise 13, part 4:</em> Southern transfer <em>Exercise 14, part 2:</em> agarose gel analysis of PCR products from Ex 14 part 1  <em>Exercise 15, part 2:</em> analysis of genomic DNA digests carried out in Exercise 15 part 1 for genomic library construction <em>Exercise 15, part 3:</em> ligation of genomic DNA insert and pB vector for library construction</td>
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<td>July 4</td>
<td><strong>Appendix week 8B:</strong></td>
<td><em>Exercise 16 part 1:</em> Screening of a genomic library: colony lifts exercise 13 part 5 continued and part 6: Southern hybridization: washes and ECL exercise C RNA isolation from spores using qiagen plant kit</td>
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<td><em>Exercise 15, part 4:</em> construction of genomic library continued: transformation of <em>E. coli</em> XL-1 Blue with recombinant vectors containing genomic DNA exercise 13, part 5: Southern blot continued: prehybridization and hybridization</td>
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| July 10 | **Topic 8 genomic cloning**  
3 to 4pm (tentative based on progress, may dedicate it to presentations | **Appendix week 9A:**  
exercise 16 part 2: prehybridization and hybridization of colony lifts,  
data analysis presentations from cDNA cloning |
| July 11 |  | **Appendix week 9B:**  
exercise 16, part 2 continued and part 3: colony hybridization: washes and ECL  
-complete RNA isolation if needed |
| July 17, Quiz 4 | **Topic 9 : overview of advanced techniques**  
1 pm- 3 pm | **Appendix week 10A**  
exercise 17 part 1: isolation and characterization of putative positive genomic clones |
| July 18 |  | **Appendix week 10B**  
exercise 17 part 2: plasmid prep on putative positives  
exercise 17 part 3: restriction digests of plasmids  
exercise 17, part 4: RT-PCR analysis  
Formal lab 1 (week 1 through 7) |
| July 25 | **Topic 9** | **Appendix week 11A**  
exercise 17 part 5: agarose gel analysis of restriction digests of putative clones  
exercise 17 part 6: agarose gel analysis of, Exercise C RT-PCR products exercise 17,  
part 4: PCR on genomic clones based on |
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<th>Date</th>
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<th>Activity</th>
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<tr>
<td>July 26</td>
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<td><strong>Appendix week 11B:</strong> data analysis and PCR gel analysis: full class discussion, summarize your data</td>
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<tr>
<td>July 31 and Aug 1</td>
<td><strong>Topic 9</strong></td>
<td>Data analysis and completion of experiments class presentations</td>
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<td>Aug 6</td>
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<td>Lab report and books due</td>
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