WELCOME TO GENETICS!
In this course we consider many important areas of genetic study including eukaryotic patterns of inheritance, genetic mapping, mutation and use of genetics for understanding gene function, chromosome organization and mutation, and Recombinant DNA technology and genome analysis.
To take this course you need to have successfully completed BIOB10H, BIOB11H (or BIOB10Y) and either a Calculus course that included probability and statistics, or PSYB07 or STAB22, or an equivalent course. Prerequisites are enforced for your benefit and that of the other students in the course.
The knowledge you learn in this course is valuable for the practice of health or veterinary-related science, genetic counseling, genetic diagnostics, the use of genetics and molecular technologies to better understand complex biological processes and systems. It also will help you be a better science-literate citizen in a complex world. If you keep up with the learning activities of this course it also can be fun!

INTERACTION TIMES AND COMMUNICATION METHODS

Lectures for this course are in SY110 Wednesdays 9-11am (there are two in-class tests at this time) and Fridays 9-10am.

Students and their TA meet once per week in SW248 or SW250 at times indicated in the timetable. Students organize into teams; each team has some lab work outside of the normal practical meetings in the term’s weeks 3-9.

Dr. H’s regular office hours are held in SY246 on Mondays and Tuesdays 4-6 (except no office hours on Labor Day Sept 5th, and Thanksgiving (Oct 10th).

email questions are welcome (hasenkampf@utsc.utoronto.ca), but allow two working days for response time.

Mr. Peilu Gan is the Head TA and lab coordinator; his email is p.gan@mail.utoronto.ca He is the person to contact regarding lab matters.

Dr. H is available for all questions about lectures and test content and is also able to provide advice about the fly crosses, and general program and post-graduation planning.

LEARNING GOALS FOR GENETICS STUDENTS

1. Students will be able to interrelate chromosome behavior during meiosis with the key rules of inheritance: segregation of alleles, independent assortment, sex linkage, linkage, and maternal inheritance.

2. Students will combine their knowledge of probability theory with the rules of inheritance to do pedigree analysis and accurately predict genetic outcomes. Additionally students will be able to interpret pedigrees and phenotypic ratios to determine if genes likely are autosomal or sex-linked, linked or sorting independently, and genotypes of parents.

3. Students will develop an appreciation of how genes work within organisms and will be able to use this knowledge to understand and predict phenotypic ratios. Also they will be able to interpret phenotypic ratios to identify the number of genes, allelic relationships, dominance relationships, and types of interaction gene interactions in biological pathways.
4. Students will do genetic crosses using the model organism *Drosophila melanogaster* to deduce the genotype, mode of inheritance, dominance relationship(s) and recombination frequency map distances and interference of several Drosophila genes. These classical genetic techniques will provide students with first hand experience with the rigor and precision needed to perform biological experiments, and with the difficulties, frustrations and errors that can occur.

5. Students will gain experience working collaboratively as part of a team to accomplish the work of a set of experiments. This includes learning to distribute a task’s workload equitably and to give each other productive feedback in a professional and constructive manner.

6. Students will evaluate their data and create a group oral and an individually, personally written report of their experiment that effectively communicate complex ideas to colleagues.

7. Students will efficiently search the primary literature for an article related to their experiment, and provide a concise analysis of key findings of the research article.

8. Students will examine the cellular processes that combat DNA damage and replication errors, and will be able to analyze how mutations can be used to explore biological processes, genome structure and evolution.

9. Students will be able to characterize the types of DNA- and chromosomal mutations and will understand the origins and consequences of these mutations.

10. Students will be able to describe the key molecular technologies that led to the sequencing of the human genome (and other model organisms) and will understand how entire genome sequencing has led to systems approaches to understanding biochemical and developmental pathways, human diseases and related therapeutic approaches.

11. Students will practice communicating as a professional in all communications for this course: emails to TAs and instructor, organization of lab notebook, and oral and written assignments.

To do well in this course it is vital to keep up with the work on a weekly basis: understanding the key concepts, completing assigned activities to develop skill in using the concepts, and doing weekly lab work on time. For example, by Monday of week 2 you will need to understand and be able to use, the material of week 1, etc throughout the course. To help you make a strong start with this learning pattern there will be two online quizzes and a bonus mark opportunity (you can work cooperatively with friends for the online quizzes and bonus mark but you must each complete the quiz, and you each must be a thinker in the final solutions!)

**COMMUNICATION INFORMATION**

Course announcements, communications and lecture notes will be available on the BIOC15 Blackboard course site. Be sure you have a Utorid and know how to access the Blackboard sites. Check the Blackboard site regularly for important, time-sensitive announcements.

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 web site. Your TA will give you instructions on how to submit your assignment to TurnItIn in advance of the submission deadline.
Lectures notes typically will be posted within 24 hrs AFTER the relevant class meeting. But creating a class with effective learning is a two-way exchange; I need your feedback and interactions to understand how well I am getting the ideas across, so I need many students attending class! If class attendance drops off too much the posted lecture videos will stop.

Learn the name of your TA and interact with your TA and fellow students with respect. They are your important learning partners! Questions about lab work and assignments, lab quizzes etc should be addressed to your TA, first.

Course evaluations for this course and all UTSC courses will be done on-line; please participate. Your assessments and insights are important to me in particular and the university in general!

ABSENCE POLICY
If you are prevented from attending or completing a course requirement (test, lab, or assignment), due to any illness or other circumstance of a grave nature, contact Dr.Hasenkampf by email within three days of the missed requirement, clearly stating the reason. This explanation should be accompanied by a completed ‘Verification of illness or injury’- form or other official documentation of the grave circumstance. These documents will be used to determine eligibility to recover any lost marks.

The ‘verification of illness or injury’ form can be found at http://www.illnessverification.utoronto.ca/getattachment/index/Verification-of-Illness-or-Injury-form-Jan-22-2013.pdf.aspx A copy of the form is provided on the last page of the syllabus. Acquaint yourself with its content such that in case of an emergency you can obtain the essential information required, even in the absence of the official form.

Remember this course is about learning Genetics. When you are well, work hard and engage! If you are sick, take care of yourself, do what you can from home and get back into the swing of the course once you are well. Feel free to contact me for help strategizing for getting caught up.

IMPORTANT ACQUISITIONS
The textbook for this course is Genetics from Genes to Genomes, 1st Canadian edition. It is packed as a bundle with the Solutions manual. The textbook/solutions manual bundle may be purchased at the bookstore.

The lab manual is posted on the Blackboard course site. You need to download your copy and bring the relevant part to each lab.

INFORMATION ON TERM TESTS AND COURSE ASSIGNMENTS
There will be two, IN CLASS term tests (on a Wednesday in October and November) and a comprehensive final exam. The final exam is held during the examination period.

Because there are three major assessments during the term (two term tests and the oral & written Lab Report) there will be no time for makeup test for the two, term tests. Anyone missing either of the term tests (and who has a valid and documented medical or serious personal problem to miss that test) will have access to that test and have access to the answers (for self-assessment), but will recover the lost marks by having to take a final exam which has the relevant portion of the comprehensive final exam more heavily weighted and hence the final exam will be worth more of their final grade.
Anyone without an acceptable (and documentable) reason for missing a test (or assignment or lab) will receive the grade of zero for the relevant work. To be eligible to recover the marks missed on a test, etc. you are required to contact Dr Hasenkampf, within 72 hours of the test or assignment and provide the appropriate medical or other type of documentation within one week of the test.

Test content, and the level of detail of tests and the exam will be as covered in the posted lecture class notes, assigned problems, assigned independent reading, the figures from the textbook (as assigned in the lecture class notes) definitions in bold in the lab manual and the questions and answers posed within the lab manual. In our class meetings I try to highlight the most important and/or most challenging concepts and applications, but the posted lecture class notes are the definitive source for lecture content that might be included on the two in class tests or the final exam.

Doing well is not just about having a good set of notes! To convert the information in the lecture notes to your own working knowledge of genetics you need to consider the content compared to what you already know, and how you can use the content to solve real world genetic problems. One good way to assess how well you are doing this is to come to class and to fully engage with the learning activities of BIOC15.

Best wishes for an exciting learning journey!

**FINAL EXAM**
The final exam in this course, as in the challenges we face in life, is comprehensive in nature. For BIOC15 this means the exam will assess your working knowledge of our topics from throughout the course. The Final exam will be held in the examination period. Anyone absent from the Final exam must petition the registrar’s office to take a deferred exam.

**LAB MEETINGS**
Labs start in the week of Sept 12-16th. Each of you is assigned to one practical section that you attend on a weekly basis. In addition to this time period, additional lab work will need to be done by your lab team; some members attend the genetics lab at additional posted times to create your genetic crosses and collect data. Attendance in labs is mandatory; absence will reduce your mark. The success of your genetic crosses requires coordinated teamwork. Please develop a good working relationship and effective communication system with your team members. The ability to work as part of a team is an important life skill; use this course as an opportunity to improve your abilities. The lab work is designed to reinforce the concepts and problems considered in the class meetings and assessed on the tests and exam. If you fully engage with the lab work it will improve your learning in genetics.

Mr Peilu Gan is the Head TA and lab coordinator; please direct general questions about the labs to him. His email is p.gan@mail.utoronto.ca

**MARKING SCHEME FOR THE COURSE**

<table>
<thead>
<tr>
<th>Component</th>
<th>Percentage</th>
<th>Description</th>
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<tbody>
<tr>
<td>Test 1 October 5th</td>
<td>18%</td>
<td>In class time (Classes 1-9)</td>
</tr>
<tr>
<td>Test 2 November 16th</td>
<td>18%</td>
<td>In class time (emphasis on Classes 11-17)</td>
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<tr>
<td>Online quizzes</td>
<td>6%</td>
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<tr>
<td>Lab engagement</td>
<td>12%</td>
<td>(attendance, participation, ability to work fairly and effectively as a team-member, lab quizzes, lab assignments.)</td>
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<tr>
<td>Oral report</td>
<td>2%</td>
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<tr>
<td>Written lab report</td>
<td>10%</td>
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<tr>
<td>Final exam</td>
<td>34%</td>
<td>(Comprehensive for all lectures) in final exam period</td>
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The University of Toronto is dedicated to fostering an academic community in which the learning and scholarship of every member may flourish, with vigilant protection for individual human rights, and a resolute commitment to the principles of equal opportunity, equity and justice. The instructor and Teaching Assistants of BIOC15 fully endorse this policy.

ACADEMIC INTEGRITY
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) outlines the behaviours that constitute academic dishonesty and the processes for addressing academic offences.

Potential offences in papers and assignments include 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 cheating includes using or possessing unauthorized aids, looking at someone else’s answers during an exam or test, misrepresenting your identity, or falsifying or altering any documentation required by the University, including (but not limited to) doctor’s notes.

Please avoid academic dishonesty, have confidence in your own ability to learn and grow academically by doing your own thinking and writing! I know you can learn a lot about genetics and yourself in this course.

ACCESSABILITY
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 SW302) are available by appointment to assess specific needs, provide referrals and arrange appropriate accommodations (416) 287-7560 or ability@utsc.utoronto.ca.

SCHEDULE OF LECTURE TOPICS AND IN CLASS TESTS. (Lab schedule is in the lab manual)
Please note this is the topic-order, but we occasionally get a little ahead or a little behind the posted dates.

September 2 (C1) Overview of BIOC15 and Modern Genetics (Chapter sections 1.1-1.4).

September 7 (C2) Inheritance – Mendel’s First and Second Laws & Probability Chapter sections 2.1-2.2), and Pedigree analysis (2.3)

September 9 (C3) Extensions of Mendelian Analysis: Dominance Relationships and Multiple alleles, wild type and mutant alleles (Chapter sections 2.4 and figures 7.22-7.24 and related text)

September 14 (C4) Extensions of Mendelian Analysis: How Genes (& the environment) interact to determine phenotypes (Chapter sections 2.4-2.5) and using Mutational Analysis to understand how phenotypes are determined (Chapter section 8.3)
<table>
<thead>
<tr>
<th>Date</th>
<th>Class</th>
<th>Topic</th>
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<tbody>
<tr>
<td>Sept 15-20th</td>
<td></td>
<td>Online Quiz 1 Classes 1-4</td>
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<tr>
<td>Sept 16 (C5)</td>
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<td>Extensions of Mendelian Analysis: Genes’ Products Interactions and Continuous Traits (Chapter section 2, 2.5)</td>
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<td>Sept 21 (C6)</td>
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<td>Chromosome Theory of Inheritance- Chromosome organization (Chapter section 6.2); Mitosis &amp; Meiosis overview (Chapter sections 3.2-3.3)</td>
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<td>Sept 23 (C7)</td>
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<td>Meiosis (Chapter section 3.3) &amp; Karyotype Analysis (Chapter section 3.1)</td>
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<td>Sept 28 (C8)</td>
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<td>X-linkage: an exception to Mendel’s First Law, Sex Determination, (Chapter sections 3.5) &amp; nonNuclear Inheritance- another exception to Mendel’s First Law</td>
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<td>Sept 30 (C9)</td>
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<td>Lecture ‘catch up if needed and test review</td>
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<tr>
<td>Oct 5</td>
<td>(C10)</td>
<td>Test 1 Classes 1-9</td>
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<tr>
<td>Oct 7</td>
<td>(C11)</td>
<td>Linkage: A Exception to Mendel’s Second Law (genes don’t always sort independently) (Chapter section 4.1 &amp; 4.3)</td>
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<td>Oct 10-14</td>
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<td>Thanksgiving and Reading Week</td>
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<tr>
<td>Oct 19</td>
<td>(C12)</td>
<td>Chi-Square Test and Linkage analysis (Chapter section 4.2) and Mapping genes on chromosomes using recombination frequencies (Chapter section 4.4)</td>
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<td>Oct 21</td>
<td>(C13)</td>
<td>Special Mapping Techniques</td>
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<td>Oct 26</td>
<td>(C14)</td>
<td>Chromosome Mutations:Deletions, Duplications, Inversions (Chapter section 9.1)</td>
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<tr>
<td>Online quiz 2</td>
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<td>October 27-Nov. 1 Classes 11,12,13,14</td>
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<tr>
<td>Oct 28</td>
<td>(C15)</td>
<td>Chromosome Mutations:Translocations (Chapter section 9.1), Transposition(Chapter section 9.2), and Karyotype Evolution (bits of Chapter section 9.1 &amp; 9.3)</td>
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<tr>
<td>Nov 2</td>
<td>(C16)</td>
<td>Aneuploid and Polyploid Chromosome Mutations (Chapter section 9.4)</td>
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<td>Nov 4</td>
<td>(C17)</td>
<td>DNA damage, Gene Mutation and DNA Repair Mechanisms</td>
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<tr>
<td>Nov 9</td>
<td>(C18)</td>
<td>DNA damage, and DNA repair mechanisms, continued</td>
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<tr>
<td>Nov 11</td>
<td>(C19)</td>
<td>Lecture ‘Catch up’ and Test 2 review</td>
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<tr>
<td>Nov 16</td>
<td>(C20)</td>
<td>Test 2 Emphasis on Classes 11-19</td>
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<tr>
<td>Nov 18</td>
<td>(C21)</td>
<td>Molecular Biology and Recombinant DNA Technology</td>
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<tr>
<td>Week of November 21-25</td>
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<td>Oral reports done in lab practicals, written reports due Nov. 28&lt;sup&gt;th&lt;/sup&gt; details below.</td>
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November 23  (C22) Molecular Biology and Recombinant DNA technology, continued

November 25  (C23) Recombinant DNA Technology and The Human Genome Sequencing Initiative

November 30th  (C24) Post Sequencing Technologies and Genomic Approaches to Genetics

Online quiz 3  November 30- Dec 5th  Classes 21-24

Lab reports are due on Monday November 28th.  The report must be completely in your own words. A paper copies must be submitted to Dr. Hasenkampf and an electronic copy via TURNITIN; your TA will provide instructions on how to use TURNITIN. The paper and electronic copy must be identical and have the same deadlines.

Life happens; there is a grace period until Tuesday November 29th at 5 pm for both the paper copy and the electronic copy. Once the grace period has ended there is a 5pt per day penalty for each working day late.

• Submitted after 5pm Tuesday Nov. 29th minus 5 pts;
• Submitted after 5pm Wednesday Nov. 30th minus 10 pts etc.

Please use this writing assignment to improve your writing skills; do your own work. Also make all your hard work pay off with a good grade by turning the report in on time! Turning it in on time is also important so you can turn your efforts to assignments in other courses, studying etc.

Oral reports are a team effort, but you each must write your own unique lab report, which has been written completely in your own words. I know you can do it! Have confidence in yourself, and use the report writing as a way to solidify your understanding of inheritance and practice the writing skills scientists use to communicate with each other and the public. This approach will optimize learning and will prevent instances of academic dishonesty.

We are using TURNITIN software that is very powerful in detecting writing that originates from sources other than yourself. You can look at the similarity report score before you submit the final version to TurnItIn. Typically a report done entirely in your own words, with proper citations, will have a similarity score of less than 16%. We expect all reports to have less than 16% similarity. Once you have created the final version of your report submitted the final version to TURNITIN and submit an identical paper copy.

Bonus Review quiz will be posted toward the end of classes

😊  I look forward to exploring the marvels of Genetics with you this term! Welcome!