



**STEM MUSE + SAGES PRESENT
STEM MENTORSHIP PROGRAM!**

SCAN THE QR CODE TO SIGN UP AS A
MENTEE AND/OR MENTOR



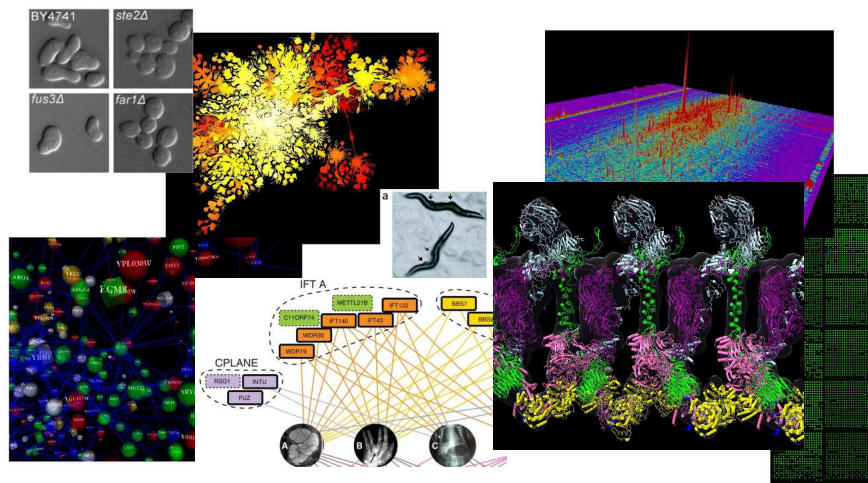
- Serves **women & underrepresented-gendered people in STEM**: undergrads, grad students, research staff, postdocs
- **One-on-One mentorship**: **Mentees choose mentors**
- Opportunity to **connect with STEM companies** & mentors
- **Networking and professional development** opportunities: group mentorship, workshops, socials, and resume/CV-bolstering events
- **Flexible scheduling**: pairs set schedule at kick-off (1 mtg./mo. for 6 months from February to July)
- Option for **in-person or virtual** interactions
- **Conversation guides** facilitate productive conversations

Testimonials: "10/10 would recommend", "best and most organized", "helped me figure out where I wanted to go and how to get there"

Mentors from more than 60 companies participated in our past programs (2020-2022), including:



BCH394P/BCH364C Systems Biology & Bioinformatics
(course # 55425 / 55330)
Spring 2023 Tue/Thu 11 – 12:30 PM WEL 2.110



Instructor: Prof. Edward Marcotte marcotte@utexas.edu

Zoom office hours: Mon 4 – 5

TA: Matt McGuffie mmcguffie@utexas.edu

Coding/problem set help hours:

Wed 3 – 4/Thu 12:30 – 1:30 in MBB 1.448BA

or by appointment on zoom

After hours Q/A, discussion: Canvas

The class zoom channel will be posted on Canvas.

It will be the same zoom for class and office hours.

Probably the most important slide today!

Course web page:

**[http://www.marcottelab.org/
index.php/BCH394P_BCH364C_2023](http://www.marcottelab.org/index.php/BCH394P_BCH364C_2023)**

This is a graduate student class!

It is open to a small # of upper division undergrads in natural sciences and engineering.

UG prerequisites: Biochemistry 339F with a grade of at least B; Computer Science 303E and Statistics and Data Sciences 328M (or Statistics and Scientific Computation 318M, 328M) with a grade of at least C-; and *consent of the instructor*.

An introduction to systems biology and bioinformatics, emphasizing quantitative analysis of high-throughput biological data, and covering typical data, data analysis, and computer algorithms.

Topics will include introductory probability and statistics, basics of Python programming, protein and nucleic acid sequence analysis, genome sequencing and assembly, proteomics, synthetic biology, analysis of large-scale gene expression data, data clustering, biological pattern recognition, and gene and protein networks.

Note: it's NOT a course on practical sequence analysis or using web-based tools. We'll use these occasionally, but the focus will be on learning the underlying algorithms, exploratory data analyses, and their applications, esp. in high-throughput biology.

By the end of the course, you'll know the fundamentals of important algorithms in bioinformatics and systems biology, be able to design and run computational studies in biology, and have performed an element of original computational biology research

Books

Most of the lectures will be from research articles and slides. For sequence analysis, there will be an **Optional text**:

Biological sequence analysis, Durbin, Eddy, Krogh, Mitchison, Cambridge Univ. Press (available from Amazon, used & ebook)

For biologists rusty on their stats, *The Cartoon Guide to Statistics* (Gonick/Smith) is very good (really!).

We will also be learning intro Python programming.

The course web site lists some recommendations to help you out, such as the free web course **Practical Python Programming**
<https://dabeaz-course.github.io/practical-python/>

**Important: There are bi-weekly coding/problem set help sessions.
Plan to attend at least one per week!**

Grading

No exams. Grades will be based on:

- **Online programming homework**
(10 points each and counting 30% of the final grade)
- **3 problem sets**
(15 points each and counting 45% of the final grade)
- **A course project** that you will develop over the semester & present in the last 3 days of class (25% of final grade)

The course project will consist of a research project on a bioinformatics topic chosen by the student (with approval by the instructor) containing an element of independent computational biology research (e.g. calculation, programming, database analysis, etc.) turned in as a web URL (20%) and presented in class (5%).

The project will be emailed as a web URL to the TA & I, developed through the semester and finished by 10 PM, April 12, 2023.

The last few classes will be spent presenting your projects.

Late policy

- All projects and homework will be turned in electronically and time-stamped.
- No makeup work will be given.
- Instead, all students have 5 days of free “late time”.
This is for the entire semester, NOT per project, and counting weekends/holidays just like any other day.
 - For projects turned in late, days will be deducted from the 5 day total (or what remains of it) by the # of days late.
 - Deductions are in 1 day increments, rounding up
e.g. 10 minutes late = 1 day deducted.
 - Once the 5 days are used up, assignments will be penalized 10% / day late (rounding up), e.g., a 50 point assignment turned in 1 ½ days late would be penalized 20%, or 10 points.

Online homework will be via *Rosalind*: <http://rosalind.info/faq/>

Enroll specifically for BCH394P/364C at:

<https://rosalind.info/classes/enroll/fb013a1910/>

Rosalind SALIND About ▾ Problems ▾ Statistics ▾ Glossary search f t My Classes ▾ edward marcotte Log out

BCH394P/364C (Spring 2023) Systems Biology/Bioinformatics

[Edit class info](#) [Edit problems](#) [Enroll link](#) [Grade sheet](#) [Assistants](#) [Print all problems](#) [Announcements](#) [All classes](#) [Delete](#)

by Edward Marcotte at University of Texas at Austin

An introduction to systems biology and bioinformatics, emphasizing quantitative analysis of high-throughput biological data, and covering typical data, data analysis, and computer algorithms. Topics will include introductory probability and statistics, basics of Python programming, protein and nucleic acid sequence analysis, genome sequencing and assembly, proteomics, synthetic biology, analysis of large-scale gene expression data, data clustering, biological pattern recognition, and gene and protein networks.

Num	Title	Solved By	Cost	Due Date	Questions	Solutions
1	Installing Python	0	2	Jan 18, 2023	📄	📄
2	Variables and Some Arithmetic	0	2	Jan 18, 2023	📄	📄
3	Strings and Lists	0	2	Jan 18, 2023	📄	📄
4	Conditions and Loops	0	2	Jan 18, 2023	📄	📄
5	Working with Files	0	2	Jan 18, 2023	📄	📄
			10			

[Found a typo?](#) [Suggest a new problem](#) [Take a tour](#)

The first homework will be due (in Rosalind) by 10 PM, Jan 18

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Installing Python

Problem 1 @ BCH394P/364C (Spring 2023) Systems Biology/Bioinformatics ↗

Dec. 7, 2012, 12:42 p.m. by Rosalind Team Topics: [Introductory Exercises](#), [Programming](#)

Why Python? [click to expand](#)

Problem

After downloading and installing Python, type `import this` into the Python command line and see what happens. Then, click the "Download dataset" button below and copy the Zen of Python into the space provided.

Time limit You'll have 5 minutes to upload the answer.

[Download dataset](#) You may make an unlimited number of attempts without being penalized.

[Questions](#)

[Found a typo?](#) [Suggest a new problem](#) [Take a tour](#)

Rosalind SALIND About ▾ Problems ▾ Statistics ▾ Glossary search f t My Classes ▾ edward.marcotte Log out

Installing Python

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Why Python? [click to collapse](#)

Rosalind problems can be solved using any programming language. Our language of choice is Python. Why? Because it's simple, powerful, and even funny. You'll see what we mean.

If you don't already have Python software, please [download and install the appropriate version for your platform](#) (Windows, Linux or Mac OS X). Please install Python of version 2.x (not 3.x) – it has more libraries support and [easier installation guides](#).

After completing installation, [launch IDLE](#) (default Python development environment; it's usually installed with Python, however you may need to install it separately on Linux). You'll see a window containing three arrows, like so:

Rosalind uses Python version 2, but we'll use version 3

Rosalind uses the "vanilla" installation of Python. You're welcome to do it this way, but I recommend Anaconda/Jupyter as a nicer option

→ New Window from the IDLE menu. You can now type code as you would

```
print "Hello, World!"
```

Select File → Save to save your creation with an appropriate name (e.g., `hello.py`).

To run your program, select Run → Run Module. You'll see the result in the interactive mode window (Python Shell).

Congratulations! You just ran your first program in Python!

Problem

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Time limit You'll have 5 minutes to upload the answer.

[Download dataset](#) You may make an unlimited number of attempts without being penalized.

[Questions](#)

Click here to turn in your answer

Installing Anaconda/Jupyter

My recommendation for a good, all-round Python installation is **Anaconda**, available free to individuals here:

<https://www.anaconda.com/products/individual>

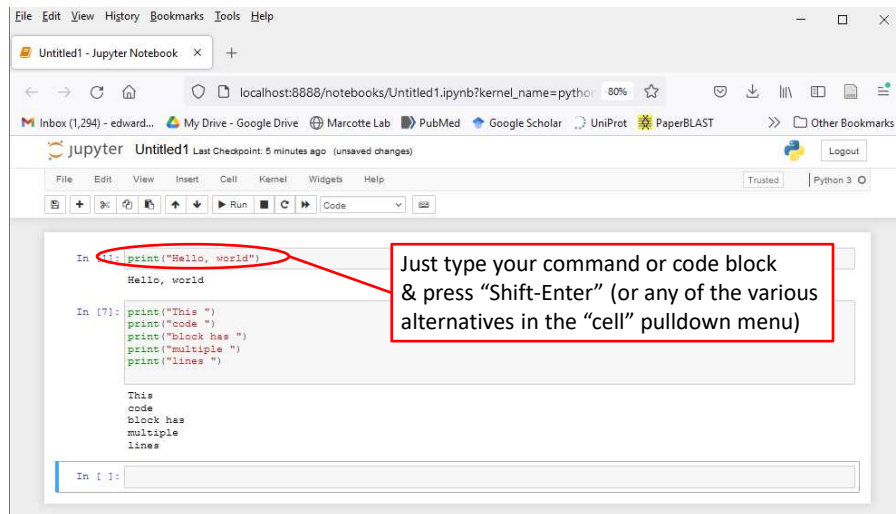
*****Get the latest Python 3 version*****

(currently 3.9 in Anaconda, but any version >3 is probably fine)

Anaconda is a general management system for the various Python libraries and packages you might need, with >7,500 data science, visualization, and machine learning packages

Anaconda also provides multiple Python interfaces. For this course, I recommend using **Jupyter Notebook**, which can be launched directly from the main Anaconda navigation window.

Jupyter is an interactive Python interface that shows your code & its output in successive entries in a shareable, archivable notebook viewable in any web browser, e.g.



It's widely used in bioinformatics and data visualization.

Back to Rosalind, for those of you that are a bit more advanced:

If you're feeling restless/adventurous...

Rosalind navigation bar: About Problems Statistics Glossary search [f] [t] My Classes edward.marcotte Log out

Installing Python

Problem 1 @ BCH394P/364C (Spring 2023) Systems Biology/Bioinformatics ↗

Dec. 7, 2012, 12:42 p.m. by Rosalind Team

Topics: Introductory Exercises, Programming

...there are quite a few good bioinformatics problems in the archives.

Rosalind navigation bar: About Problems Statistics Glossary search [f] [t] My Classes edward.marcotte Log out

Problems

Bioinformatics Stronghold List Tree

Rosalind is a platform for learning bioinformatics and programming through problem solving. Take a tour to get the hang of how Rosalind works.

Last win: charlotte.hui.wang vs. "Dictionaries", 12 minutes ago Problems: 264 (total), users: 91799, attempts: 1516775, correct: 638920

ID	Title	Solved By	Correct Ratio	Questions	Solutions	Explanation
DNA	Counting DNA Nucleotides	53370				
RNA	Transcribing DNA into RNA	47647				
REVC	Complementing a Strand of DNA	43218				
FIB	Rabbits and Recurrence Relations	25082				
GC	Computing GC Content	25001				
HAMM	Counting Point Mutations	20161				
IPRB	Mendel's First Law	16649				
PROT	Translating RNA into Protein	22086				
SUBS	Finding a Motif in DNA	22395				
CONS	Consensus and Profile	12281				
FIBO	Mortal Fibonacci Rabbits	10554				
GRPH	Overlap Graphs	9925				
IEV	Calculating Expected Offspring	9620				
LCSM	Finding a Shared Motif	8609				
LIA	Independent Alleles	5051				
MPRT	Finding a Protein Motif	5290				
MRNA	Inferring mRNA from Protein	8135				
ORF	Open Reading Frames	6245				
PERM	Enumerating Gene Orders	10953				
PRTM	Calculating Protein Mass	10650				
REVP	Locating Restriction Sites	6619				
SPLC	RNA Splicing	7393				
LEXF	Enumerating k-mers Lexicographically	6060				
LGIS	Longest Increasing Subsequence	2690				



Expectations on working together

Students are welcome to discuss ideas and problems with each other, but **all programs, Rosalind homework, problem sets, and written solutions** **should be performed independently,**

→ *except* the final presentation.

tl;dr: study/discuss together
do your own programming/writing/project
collaborate on the final presentation



THE UNIVERSITY OF TEXAS AT AUSTIN

Student Judicial Services

Office of the Dean of Students

What is Academic Dishonesty?

In promoting a high standard of academic integrity, the University broadly defines academic dishonesty—basically, all conduct that violates this standard, including *any act designed to give an unfair or undeserved academic advantage*, such as:

- Cheating
- [Plagiarism](#)
- [Unauthorized Collaboration / Collusion](#)
- Falsifying Academic Records
- Misrepresenting Facts (e.g., providing false information to postpone an exam, obtain an extended deadline for an assignment, or even gain an unearned financial benefit)
- Any other acts (or attempted acts) that violate the basic standard of academic integrity (e.g., [multiple submissions](#)—submitting essentially the same written assignment for two courses without authorization to do so)

<https://deanofstudents.utexas.edu/conduct/academicintegrity.php>

- By submitting *as your own work* any unattributed material that you obtained from other sources, you have committed plagiarism.
- Copying homework solutions from other students or internet sources (e.g. CourseHero) is cheating, collusion, and/or plagiarism.
- Software and computer code are legally considered in the same framework as other written works. Copying code directly without attribution is plagiarism.

- Any materials found online (e.g. CourseHero) that are associated with you, or any suspected unauthorized sharing of materials, will be reported to Student Conduct and Academic Integrity in the Office of the Dean of Students. These reports can result in sanctions, including failure in the course.

See the university's official policy on plagiarism here: <https://catalog.utexas.edu/general-information/appendices/appendix-c/student-discipline-and-conduct/>

- You can use the internet to get *ideas*, programming *suggestions* and *syntax*, but **downloading completed answers to assigned questions and submitting these as your own work is cheating/plagiarism.**
- **Copying entire programs** verbatim from marked repositories offering Rosalind homework solutions **is cheating and plagiarism.**

Consequences of Academic Dishonesty Can Be Severe!

You may see or hear of other students engaging in some form of academic dishonesty. If so, do not assume that this misconduct is tolerated. Such violations are, in fact, regarded very seriously, often resulting in severe consequences.

Grade-related penalties are routinely assessed ("F" in the course is not uncommon), but students can also be suspended or even permanently expelled from the University for scholastic dishonesty.

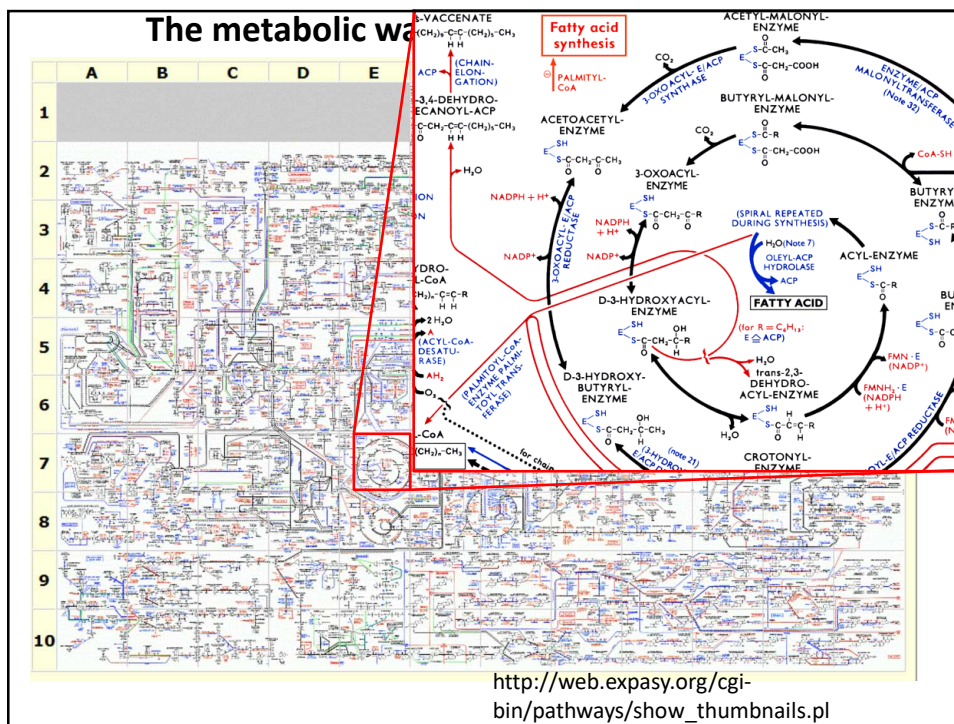
But there's always an exception!

Later in the semester,
we'll try co-programming
with AI using chatGPT,
where the goal is to make
the computer write the
code for you



Why are we here?

(practically, not existentially)

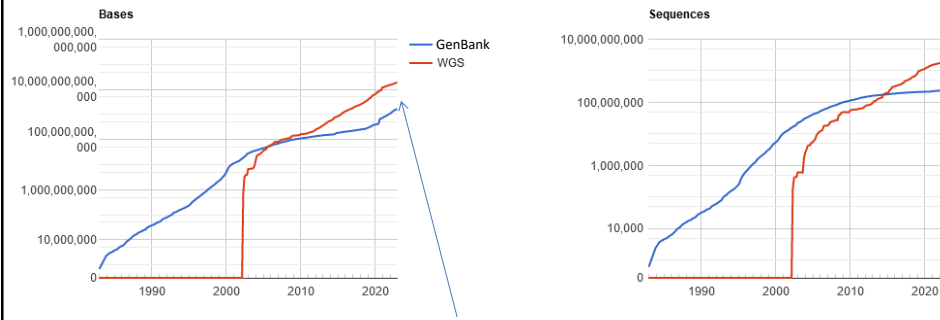


Our current-ish knowledge of human metabolism...

Total number of reactions	7,440
Total number of metabolites	5,063
Number of unique metabolites	2,626
Number of metabolites in extracellular space	642
Number of metabolites in cytoplasm	1,878
Number of metabolites in mitochondrion	754
Number of metabolites in nucleus	165
Number of metabolites in endoplasmic reticulum	570
Number of metabolites in peroxisome	435
Number of metabolites in lysosome	302
Number of metabolites in Golgi apparatus	317
Number of transcripts	2,194
Number of unique genes	1,789

Nat Biotechnol. 2013 May;31(5):419-25
Updated in Metabolomics 2016 12:109

Pales beside the phenomenal explosion of DNA sequencing:



Here are the latest statistics...

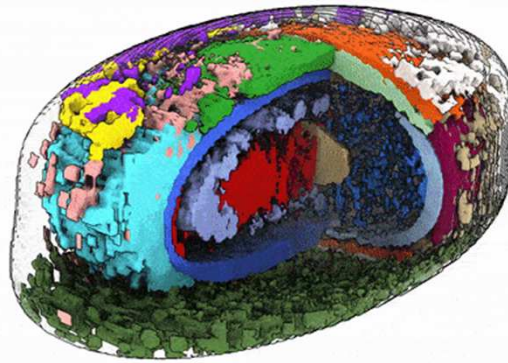
December 2022:
 1.6 trillion bp Genbank
 +
 19 trillion bp DNA
 whole genome
 shotgun sequencing

Which basically means GenBank is falling behind more every year!

<http://www.ncbi.nlm.nih.gov/genbank/statistics>



Just last week:



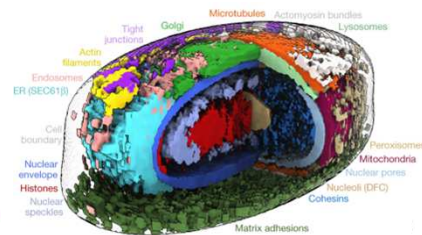
RESEARCH BRIEFINGS | 04 January 2023

Structural landscape inside cells mapped in detail

More than 200,000 human stem cells were imaged at high resolution and in 3D to make a reference data set that was used to create a generalizable computational framework. This enables cell shapes and the locations of internal structures to be measured and compared using rigorous statistical methods.



This is a summary of: [Viana, M. P. et al. Integrated intracellular organization and its variations in human iPS cells. *Nature* https://doi.org/10.1038/s41586-022-05563-7 \(2023\).](https://doi.org/10.1038/s41586-022-05563-7)



Why are we here? We have no choice!

- **Biologists are faced with a staggering deluge of data, growing exponentially**
- **Bioinformatics/comp bio tools and approaches help us understand these data and work productively, and to build increasingly powerful models of biological systems**
- **We'll learn important basic concepts in this field and get exposed to key technologies driving the field**

Specifically...

We'll cover the following topics, approximately in this order:

BASICS OF PROGRAMMING

Introduction to Rosalind
A Python programming primer for non-programmers
Rosalind help & programming Q/A

BIOLOGICAL SEQUENCE ANALYSIS

Substitution matrices (BLOSSUM, PAM) & sequence alignment
Protein and nucleic acid sequence alignments, dynamic programming
Sequence profiles
BLAST! (the algorithm)
Biological databases
Markov processes and Hidden Markov Models

GENOMES, PROTEOMES, & "BIG BIOLOGY"

Gene finding algorithms
Genome assembly & how the human genome was sequenced
An introduction to large gene expression data sets
Promoter and motif finding, Gibbs sampling
Clustering algorithms, hierarchical, k-means, self-organizing maps,
force-directed maps
Classification algorithms
Principal component analysis and data transformations

NETWORK & SYNTHETIC BIOLOGY

Biological networks: metabolic, signaling, graphs, regulatory
Deep homology and the evolution of traits
Designing, simulating, and building gene circuits
Genome design and synthesis

Plus, expert guest lectures on:

More advanced Python techniques
NGS best practices
Protein mass spectrometry / proteomics
Protein 3D structural modeling

THE FINAL COURSE PROJECT IS DUE by 10 PM, April 12, 2023

The last 3 class days will be for presenting your projects