BME 4985/6160 & CSE 4095/6800: Computational Genomics
Spring 2009

Time/place: TuTh 2-3:15pm / ITE C-80. ITE 125 (room change)

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Course outline: Started in 1995 by the completion of the first genome sequence of a free-living organism, *H. influenzae*, the genomic era has led to thousands of complete genome sequences deposited in public databases and many more genome projects at various stages of completion. The large-scale availability of genome data is revolutionizing biological and medical research, with data-driven computational approaches taking a central role. This course covers fundamental computational methods for genomic data analysis, with a main emphasis on statistical methods and current applications in genomics and genetic epidemiology.

Tentative list of topics to be covered: Basic probability theory and statistics; introduction to R; statistical modeling of biological sequences; EM and Gibbs sampling algorithms for DNA motif discovery; Markov chains; profile HMMs for representing sequence families; models of DNA and protein evolution; likelihood methods in phylogenetics, bootstrapping; basic principles of population genetics; genotype phasing and haplotype frequency estimation; computation of Mendelian likelihoods, Elston-Stewart and Lander-Green algorithms; case-control association tests. The list of topics may change according to progress and students' interest.

Textbooks: There is no required textbook for this course. Most of the covered material appears in the following books:


Prerequisites: Desired pre-requisites are an introductory course in bioinformatics and undergraduate-level courses in biology, programming, and statistics. However, required background in these areas will be provided as needed via lectures and supplementary readings.

Grading: Grades will be assigned based on in-class quizzes (10%), bi-weekly homework assignments (40%) and a final project (50%). Homework assignments will consist of both theoretical problems reinforcing the material covered in lectures and practical assignments using the R statistical package. The final project will give you the opportunity to study a computational genomics problem in more depth. You are encouraged to devise your own final project topic; suitable topics include surveys of computational genomics topics not covered in the lectures, design and implementation of novel algorithms, and empirical evaluation of existing methods. Project requirements will include submitting two intermediate progress reports and a written final
You will also be required to give a short presentation on your project at the end of the semester. When appropriate, projects can be done in teams of two students.

**HuskyCT:** We have a HuskyCT site for the class; you can access it by logging in with your NetID and password at [https://huskyct.uconn.edu/](https://huskyct.uconn.edu/). Please check this site regularly for class materials, assignment clarifications, changes in class schedule, and other class announcements.

**Academic honesty:** You are expected to adhere to the highest standards of academic honesty. All submitted solutions must be your own work. You may discuss ideas and concepts with other people, but *must not share written solutions or computer code*. Use of published materials is allowed, but the sources should be explicitly stated in your submission. Violations will be reviewed and sanctioned according to the University Policy on Academic Integrity.

**Students with disabilities:** If you have a documented disability for which you are or may be requesting an accommodation, you are encouraged to contact the instructor and the Center for Students with Disabilities or the University Program for College Students with Learning Disabilities as soon as possible to better ensure that such accommodations are implemented in a timely fashion.