# Introduction to Methods in Computational Biology and Genomics

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#### Overview

- Course
  - Course Intro
  - Teaching Philosophy
- 2 In-Class Activity
- 3 Computing and Genomics
  - Computer Requirements
  - Genomics: Why We're Here

## Today's Goals

- Get familiarized with course format
- Meet each other and myself
- Understand course expectations
- Know my teaching methods and reasons for offering this course

## Course Objectives

#### Objective 1

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### Objective 2

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#### Objective 3

Combine 1 and 2 to test biological hypotheses with RNAseq data and present those results in an "IMRD" format (pronounced EM-rod)

# Course Syllabus

## Course Grade

#### Course Policies

• Group Work vs. Own Work

#### **Course Policies**

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- No Direct Attendence Policy
- Interact and Contribute in Class

#### Potential Course Policies - TBD

• Latest pre-class email time (5pm? 9pm?)

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- Slack?

Bring a laptop

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- $\bullet \ \, \mathsf{Thursdays} = \mathsf{Lab}$

# Project

# Teaching Philosophy

- Clear Goals
- Active Learning
- Student Driven

#### Clear Goals

- Presented before each class
- Concepts or skills to focus on
- Call me on it if I forget them

## **Active Learning**

- Student Participation
- A different style than lecture-based courses (flipped courses fall into this category)
- Natural fit for smaller class size, advanced material, and learning skills

#### Student Driven

- Student Participation Required
- Work through examples in class and apply to your own question
- Many skills need to be practiced, not taught
- Grand Bargain

Pair up \*randomly\*

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  - Feel free to expand (see my slide for reference)

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  - Two answers from "Whimsy
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- Introduce partner to class

## Computer Requirements

- Bring your computer to class
- Run some software locally
  - How much storage do your computers have?
- Connect to cluster

#### Research Tools

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- github
  - Website for version control as well as script and code storage

## Programming Languages

- R
  - free statistical software
- bin/bash
  - unix language, automate many tasks, interact with cluster computers

## Software

• R -¿ R-Studio

### Software

- R -¿ R-Studio
- github -¿ SourceTree

#### Software

- R -¿ R-Studio
- github -¿ SourceTree
- data analysis software -¿ fastqc/trimmomatic/kallisto

## **Cluster Computing**

- Duke Computing Cluster
- SLURM workload manager

#### Hand Raising Request

• Student Driven - Active Learning

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- Student Driven Active Learning
- Raise your hand if you're confused

#### Hand Raising Request

- Student Driven Active Learning
- Raise your hand if you're confused
- Provides me with helpful feedback

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  phenomena such as heterosis (hybrid vigour), epistasis (effect of one
  gene on another), pleiotropy (one gene affecting more than one trait)
  and other interactions between loci and alleles within the genome.
- In contrast to genetics, which refers to the study of individual genes and their roles in inheritance, genomics uses high throughput DNA sequencing and bioinformatics to assemble, and analyze the function and structure of entire genomes.

## Brief History of Sequencing

- Allozymes
  - Electrophoresis separates different proteins by amino acid makeup
- Sanger Sequencing
  - Determines the sequences a single piece of DNA up to 500bp
- NGS Next Generation Sequencing
  - Reads sequence of many pieces of DNA many billions of times

## Brief History of Sequencing

- Allozymes
  - 1960's
- Sanger Sequencing
  - 1977
- NGS Next Generation Sequencing
  - 2000

## Next Generation Sequencing vs. Sanger

- Output for Quality Tradeoff
  - NGS = High Output / Lower Quality
  - $\bullet \ \, \mathsf{Sanger} = \mathsf{Low} \ \, \mathsf{Output} \ \, / \ \, \mathsf{High} \ \, \mathsf{Quality}$

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- NGS Methods and Machines
  - IonTorrent
  - Illumina
  - PacBio

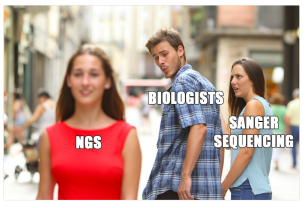
## Next Generation Sequencing vs. Sanger

- Output for Quality Tradeoff
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Following

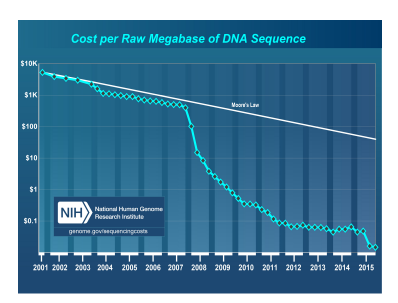
This meme is everywhere, so I thought I'd add a biology twist to it.



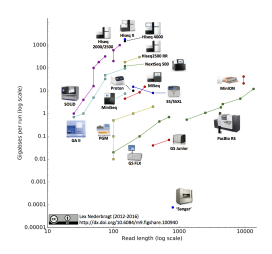
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it.

#### Sequencing Cost



# Sequencing Output



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- How does this change the scope of research?
  - (hint: Sanger is good for studying what?)

#### Course Motivations

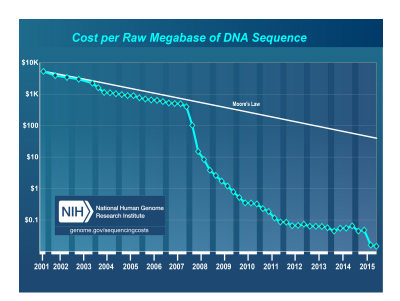
- Biologist increasingly need to be programmers
- Topics often aren't introduced to undergraduates
- Cheap and Free Data

#### Computational Need

- Biologist increasingly need to be programmers
- Many hypotheses are tested by generating piles of data
- Regardless of your future plans, programming and hypothesis testing are skills that all STEM students should have

# Cheap/Free Data

#### Cheap/Free Data



## Cheap/Free Data

- Data is cheap and often free
- Computation is getting faster
- Combined, this means student projects are feasible

#### Personal Experience

- Worked in Duke IGSP/CHGV Sequencing Core for 4 years
  - Ran Illumina GA, GA2, HiSeq2000
- PhD Thesis on Mouse Lemur Genomics
  - Whole Genome Sequencing and Mutation Rate
  - Rates of Sperm Gene Evolution

# The End