# Lecture 01 Intro to practical bioinformatics

#### Practical Bioinformatics

Biol 4220 @ WUSTL

Course: Practical Bioinformatics (BIOL 4220)

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## Practical bioinformatics

Practice foundational computing skills for everyday biological research

We all have different backgrounds, research interests, goals, etc.

## Practical bioinformatics

#### Broad goals:

- Learn new computer skills for biological data analysis
- Translate research ideas into code
- Solve problems independently
- Communicate in technical terms

## Practical bioinformatics

#### Specific skills we'll develop:

- Write and debug programs
- Build your own analysis pipeline
- Test hypotheses with pipelines
- Make reproducible research
- Communicate research findings

## Instructors

Instructor

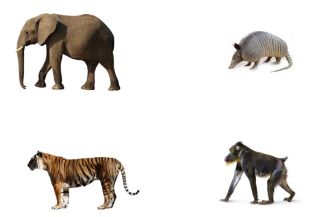
Michael Landis

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https://landislab.org

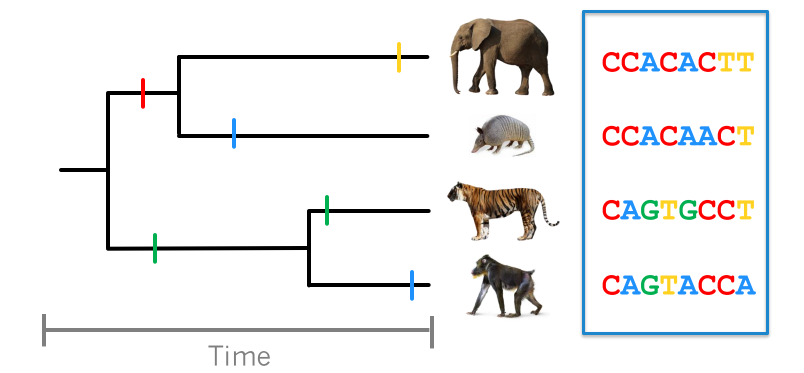
Teaching assistant
Hannah Chay
<a href="mailto:chayh@wustl.edu">chayh@wustl.edu</a>

## Statistical phylogenetics



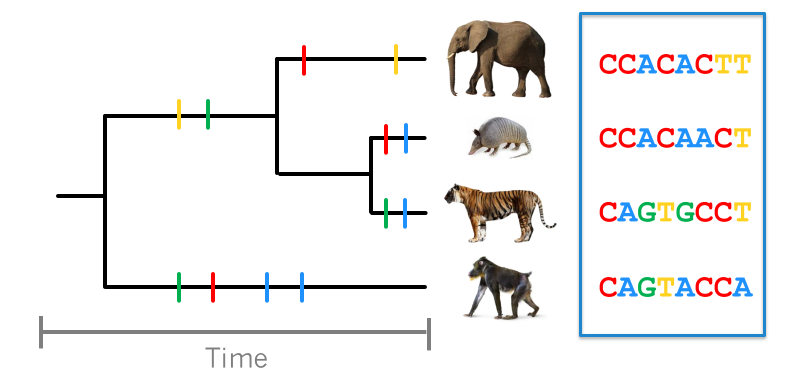
How species related? How old are they? How did their traits evolve?

#### DNA



A likely history (six mutations)

#### DNA



A less likely history (twelve mutations)

## Lecture 01 outline

Why bioinformatics?
Biol 4220 overview
Biol 4220 logistics
Brief intro to Unix

# Why bioinformatics?

Computers are essential to modern biological research

- \$1K human genome
- global biodiversity health
- human brain connectome
- tracking SARS-CoV-19
- identifying genetic diseases
- reconstructing tree of life

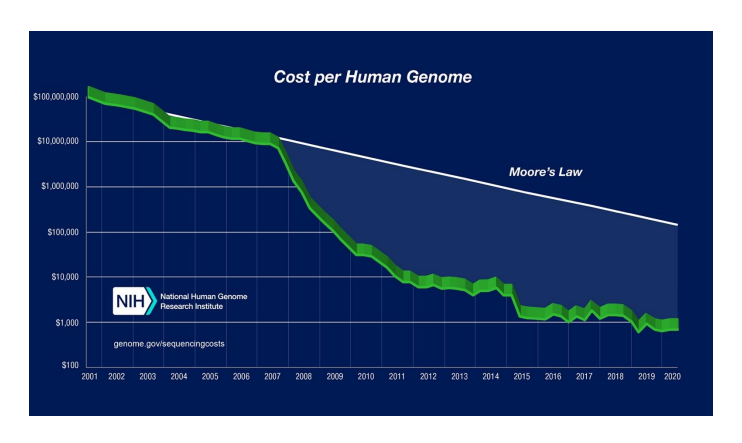
# Why bioinformatics?

Different biological disciplines face similar computational challenges

### Every year

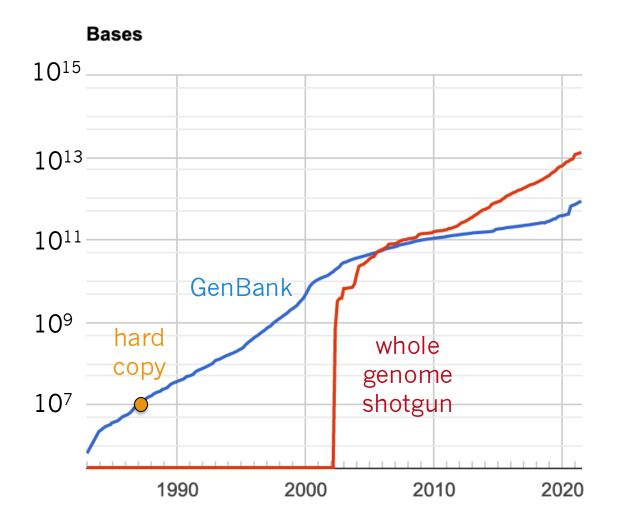
- more data samples
- higher dimensional data
- methods grow more resource-intensive
- methods are more scalable
- methods are more interconnected
- need for reproducibility increases

# More data samples



Genome cost fall by >50% every two years

## More data samples

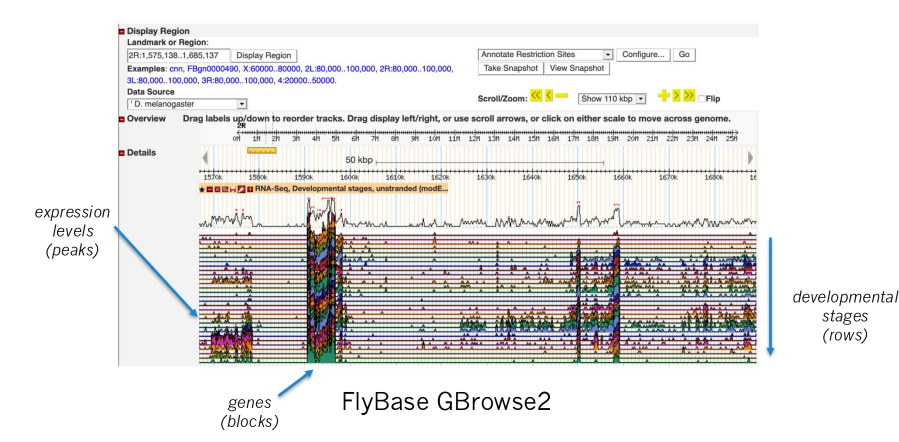


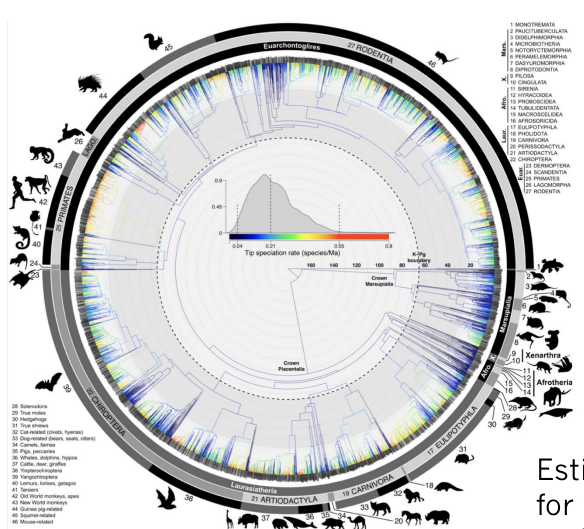


1987 hard copy of GenBank + EMBL

## More data dimensions

# Gene expression for fly level x gene x developmental stage





More resource-intensive methods

Estimating the phylogeny for 6000+ mammal species took 120+ computer years

## More scalable methods

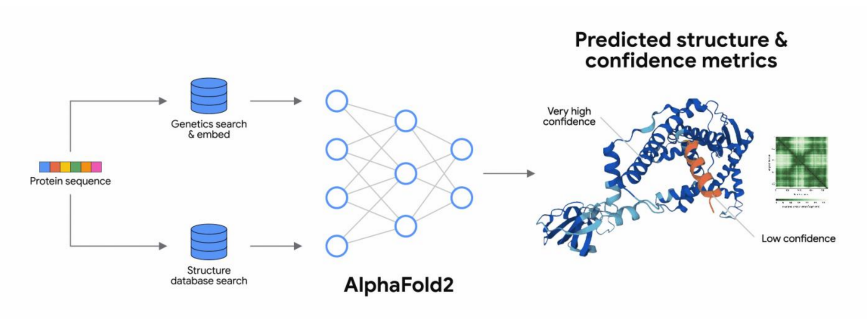
25 million 35-bp reads per hour 3.2 Gbp in human genome



**Bowtie** is an ultrafast, memory-efficient short read aligner. It aligns short DNA sequences (reads) to the human genome at a rate of over 25 million 35-bp reads per hour. Bowtie indexes the genome with a Burrows-Wheeler index to keep its memory footprint small: typically about 2.2 GB for the human genome (2.9 GB for **OSI certified** paired-end).

Bowtie: short-read alignment software

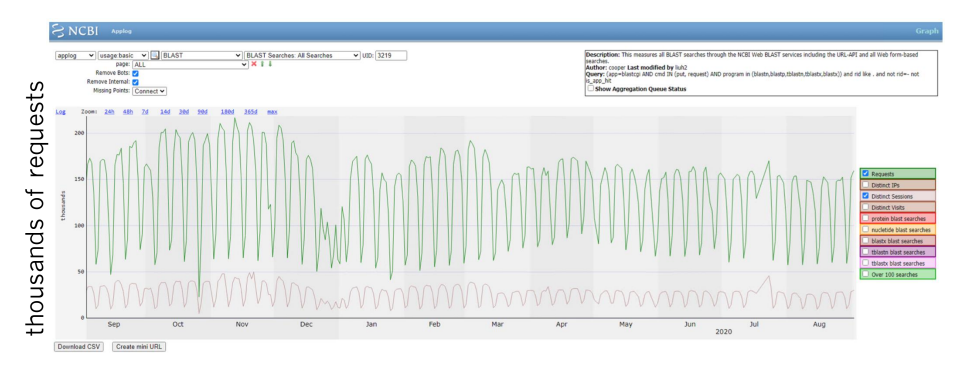
# More detailed predictions



(trained with massive clusters and datasets)

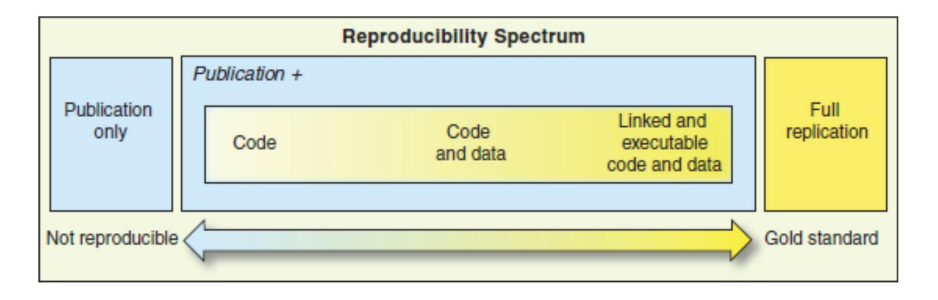
## More interconnected methods

NCBI: 150k+ BLAST requests each weekday



## Greater need for reproducibility

Computational methods allow exact reproduction of published results



from "Reproducible research in computational science" in Peng (2011, Science)

# What interests *you* in biological research?

#### You wrote:

- genetics and genomics
- virology
- phylogenetics
- gene expression
- agriculture
- cancer biology
- population genetics
- immunology

# Why are *you* interested in bioinformatics?

#### You wrote:

- curious about field
- learn new tools and techniques
- difficult research project
- improve coding skills
- develop confidence
- learn about operating systems
- preparation for future career
- build an analysis pipeline

# Biol 4220 topics

### Computational skills

- Unix-based operating systems
- Python and shell scripts
- scientific computing libraries
- version control software
- bioinformatics pipeline design

### Biological problems

- sequence processing
- molecular phylogenetics
- hypothesis testing

# Course page

#### All course info is centralized here:

github.com/WUSTL-Biol4220/home

#### Contains links to:

- syllabus
- lectures
- labs
- course project
- GitHub Classroom

## Labs

Each lab focuses on a new set of skills

One lab is assigned per class

Labs will be submitted using an online tool called GitHub Classrooms

# Free Writing

Reinforce understanding

Submissions earning high grades have these qualities:

- demonstrates understanding of recent material
- explains their strategy for solving recent lab assignment
- describes why they found recent material interesting or confusing
- suggests good quiz questions based on recent material

Each Monday after lecture for 15 min

# Quizzes

Assess understanding

Closed notes, pen and paper

#### Example

Q: Describe what each step does in this command:

```
ls *.txt | grep -v "draft" | wc -l
```

A: This pipeline (1) lists all local files ending in "txt", (2) filters out files that contain "draft" in the name (-v), and then (3) prints the number (-l for lines) of final matches

Each Wednesday after lecture for 15 min

# Course project

Design a pipeline to analyze genes

#### Example pipeline:

- download and align sequences
- summarize molecular variation
- build molecular phylogeny
- print and plot output
- add 2+ unique features

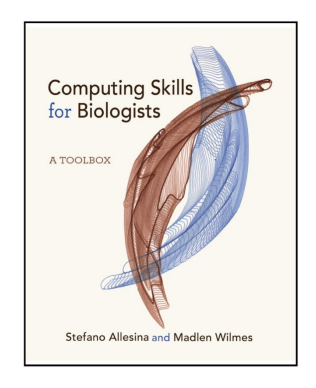
At the end of the semester, students will

- present their pipeline
- submit code, output, documentation

(more details later in semester)

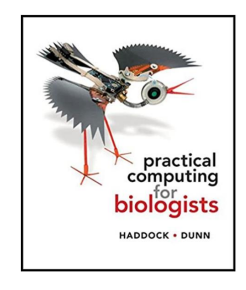
# Coding resources

#### Primary text



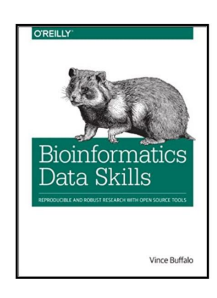
Allesina & Wilmes ISBN: 9780691167299 ~\$35

#### ...other useful texts



Haddock & Dunn ISBN: 0878933913

~\$60



Buffalo ISBN: 1449367372

~\$35

# Coding resources

You may use ChatGPT, StackOverflow, etc. for Lab and Project assignments

Important to use these tools responsibly, especially when learning the fundamentals

Comment code/text using these tools

- provide example of query or search terms
- describe any errors they produced
- describe motivation to use the tool (e.g. what was confusing?)

Visit <a href="https://it.wustl.edu/ai/">https://it.wustl.edu/ai/</a> for free ChatGPT access

# Participation

#### Communicate with others!

#### Examples

- asking and/or answering questions
- working in groups
- helping other students
- visiting office hours
- discussing research problems

Questions?

# Operating systems (OS)

**Operating systems** coordinate user commands with computational resources & hardware

#### Examples:

- Windows
- Mac OS X (Unix-based)
- Linux (Unix-based)

Most scientific computing uses *Unix*-based systems; we'll be using the Linux distribution, *Ubuntu* 

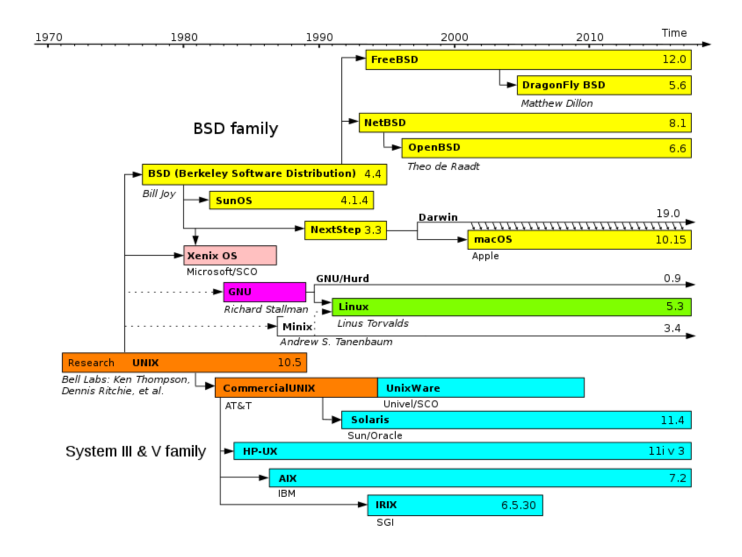
# Operating systems (OS)

What do operating systems do?

#### They manage

- *user interface* for computer input/output
- scheduled tasks across multiple users/resources
- *user interruptions* of scheduled tasks
- memory use in efficient manner
- filesystem organization on hard drive
- *user permissions* for resource security
- **network communication** with other devices
- custom software to interact w/ OS and hardware

## Unix family tree

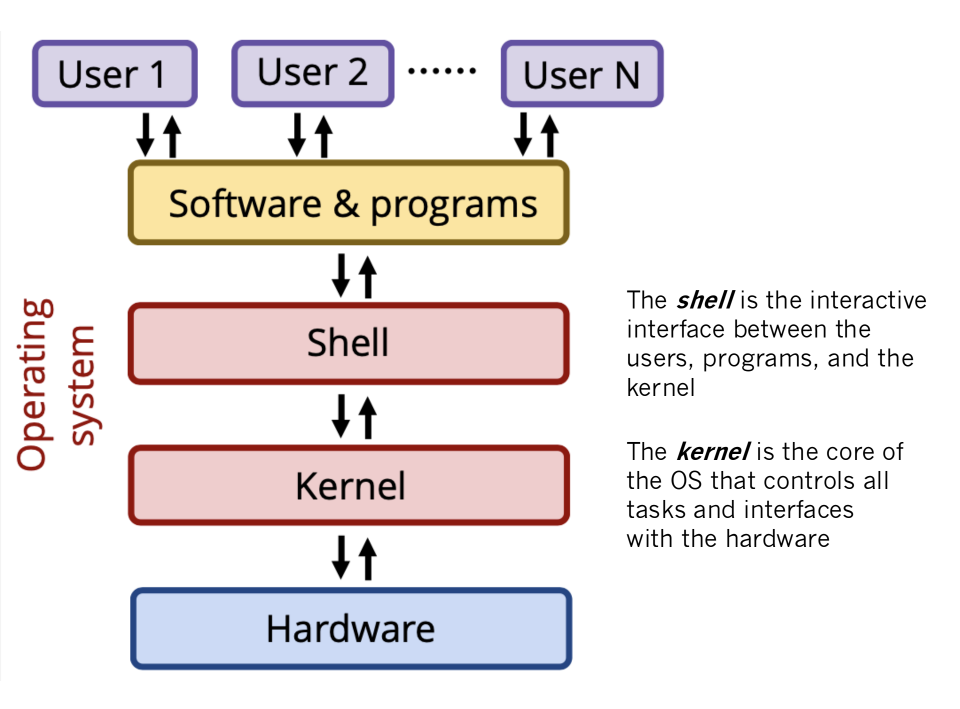


## Ubuntu

#### We'll use Ubuntu 24.04 LTS

- Reliable testing + release cycles
- Excellent tutorials
  - https://ubuntu.com/tutorials
- Extremely active support community <u>https://ubuntuforums.org</u>
- modified *Debian kernel* for stability
- popular bash shell by default





## Kernel vs. shell

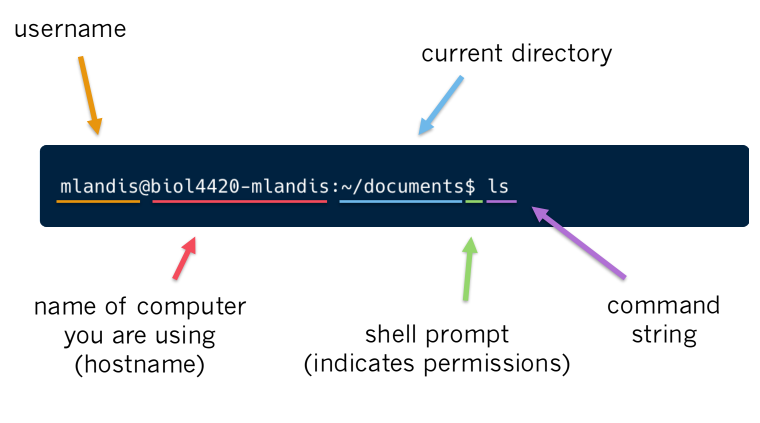
The *kernel* has control over all computer resources, including processors, memory, storage, devices, task management, etc.

The *shell* is a command line interface and scripting language that communicates user commands to the kernel for processing

- > # connect to my workstation
- > ssh mlandis@128.252.89.47

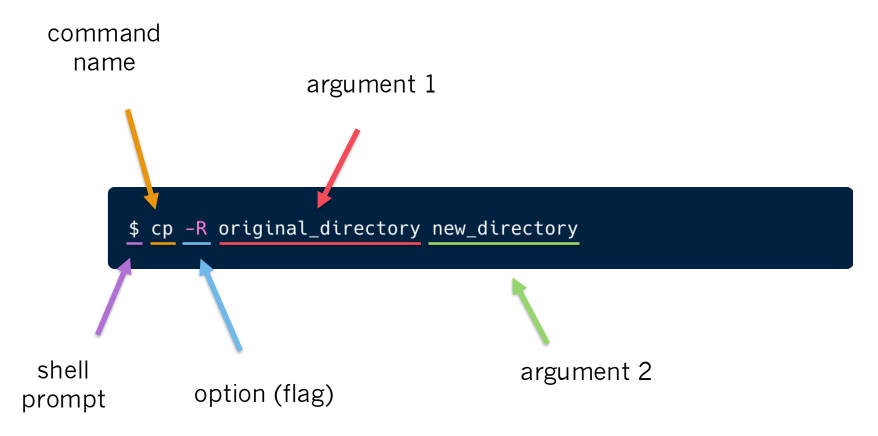
Example of Unix shell command

## Command line



The *command line* accepts commands provided by the user (you!)

# Command string



A *command* is applied against an *argument(s)* and its behavior can be modified by *option(s)* 

#### Computers are predictable

- accept input as data
- process that data
- output processed data

"On two occasions, I have been asked [by members of Parliament], 'Pray, Mr. Babbage, if you put into the machine wrong figures, will the right answers come out?' I am not able to rightly apprehend the kind of confusion of ideas that could provoke such a question."

Charles Babbage "father of the computer"



## Overview for Lab 01