### **BIOL5705: Gene Sequence Analysis**

### **Title**

Computational methods for detecting gene homology and phylogenetic tree construction.

### Lecture Layout

There are 4 lectures and each lecture is 2 hours in length. I will use 1 - 1.5 hours for lecture and you can work on the assignment and ask questions during the remaining time.

#### **Lecture Location & Time**

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Jan.16 - 1:30 - 3:30 in 4202 Mona Campbell Building
Jan.23 - 1:00 - 3:00 Rose Room 9th floor of Tupper Building
Jan.30 - 1:00 - 3:00 Rose Room 9th floor of Tupper Building
Feb.6 - 10:00 - 12:00 Andrew Roger's Lab Meeting Room (8B-1 Tupper)
```

### **Grading/Evaluation**

Grading will be based on 4 equally weighted assignments.

Assignments will be due at the start of the next lecture (or one week later for the last assignment). Students are encouraged to ask each other and me for help, but assignments should be done individually.

Assignments are to be emailed to <a href="mailto:morgan.g.i.langille@gmail.com">morgan.g.i.langille@gmail.com</a>

#### Course Materials

All lecture materials including presentations and assignments are available online at: <a href="http://morganlangille.com/biol5705.html">http://morganlangille.com/biol5705.html</a>

#### **Outline**

## 1st lecture (Jan. 16<sup>st</sup>)

- What is homology?
- Orthologs, paralogs, etc.
- Local vs global alignment
- E values, bit scores, "coverage", identity vs similarity
- Different BLAST flavours (blastn, blastp, tblastn, etc.)
- Databases (nr, refseq, etc.)
- Blast (Web)

## 2<sup>nd</sup> Lecture (Jan 23<sup>rd</sup>)

- HMMs (HMMer & PFAM website)
- PSSMs (PSI-BLAST)
- RNA secondary structures (stochastic free grammars)
- Genome Alignment (Mauve)
- Mappers (Bowtie)
- Assemblers
- Running BLAST locally

## 3<sup>rd</sup> Lecture (Jan. 30<sup>th</sup>)

- Multiple Alignment (Clustalw, Muscle, T-COFFEE, HMMAlign)
- Editing Alignments (Gblocks)
- Viewing Alignments (Jalview)

# 4<sup>th</sup> Lecture (Feb. 6<sup>th</sup>)

• Why we build trees?

Tree file formats

Parsimony

Neighbour Joining (distance based)

ML (PhyML, RAxML, FAST-Tree)

Bayes (Mr. Bayes, Beast)

http://www.phylogeny.fr/