Lab #1 BioPython

BioPython

- * Sequence format conversion
- * Sequence maniplulation
- * Interface to common programs/databases
 - * BLAST, Clustalw, EMBOSS, SCOP, SwissProt, ...
- * PubMed & Medline access
- * Simple GUI programs
- * BioSQL integration
- * http://biopython.org/DIST/docs/tutorial/Tutorial.html

Simple SwissProt Example

```
from Bio import ExPASy
from Bio import SeqIO
handle = ExPASy.get_sprot_raw("A0LR17")
seq_record = SeqIO.read(handle, "swiss")
handle.close()
```

Note: help() and vars() come in handy here...

BLAST

```
from Bio.Blast import NCBIXML

result=NCBIWWW.qblast("blastp","swissprot",
"MAKMIAMADEAARDALEDCMNOLADAWKUMICDKCDNYWIEK
```

from Bio.Blast import NCBIWWW

```
"MAKMIAMADEAARRALERGMNQLADAVKVTLGPKGRNVVLEK
KWGAPTITNDGVSIAKEIELEDPYEKIGAELVKEVAKK")
blast_record = NCBIXML.read(result)
result.close()
blast_record.alignments
```

Pubmed

```
from Bio import Entrez
from Bio import Medline

# Always tell NCBI who you are
Entrez.email = "sludtke@bcm.edu"
handle = Entrez.esearch(db="pubmed", term="Ludtke SJ[Author]",
retmax=500)
record = Entrez.read(handle)
print(record)
ids=record["IdList"]

handle=Entrez.efetch(db="pubmed",id="22696402",rettype="medline")
records = list(Medline.parse(handle))
```

Medline Terms

Affiliation [AD]

Article Identifier [AID]

All Fields [ALL]

Author [AU]

Author Identifier [AUID]

Book [book]

Comment Corrections

Corporate Author [CN]

Create Date [CRDT]

Completion Date [DCOM]

EC/RN Number [RN]

Editor [ED]

Entrez Date [EDAT]

Filter [FILTER]

First Author Name [1AU]

Full Author Name [FAU]

Full Investigator Name [FIR]

Grant Number [GR]

Investigator [IR]

ISBN [ISBN]

Issue [IP]

Journal [TA]

Language [LA]

<u>Last Author [LASTAU]</u>

Location ID [LID]

MeSH Date [MHDA]

MeSH Major Topic [MAJR]

MeSH Subheadings [SH]

MeSH Terms [MH]

Modification Date [LR]

NLM Unique ID [JID]

Other Term [OT]

Owner

Pagination [PG]

Personal Name as Subject [PS]

Pharmacological Action [PA]

Place of Publication [PL]

PMID [PMID]

Publisher [PUBN]

Publication Date [DP]

Publication Type [PT]

Secondary Source ID [SI]

Subset [SB]

Supplementary Concept[NM]

Text Words [TW]

Title [TI]

Title/Abstract [TIAB]

Transliterated Title [TT]

UID [PMID]

<u>Version</u>

Volume [VI]

Lab 1 - Homework 5

- Download the starter program from the class website. The starter program queries PubMed for articles written by a specific author, and prints the number of publications and number of coauthors. Try the program and understand what it's doing Use this program as a starting point for the following exercise. Each person should select and complete one task (any of the 4). They are ordered from hardest to easiest. You may work separately, or in small groups. If you work in groups, you must each write your own code, but are free to discuss methods and how the program works. You may use any online resources you like, and please ask questions any time. Submit your answer like homework.
- 1. Rather than simply counting the number authors in the retrieved records, tabulate the number of times each coauthor name appears, then print a) the number of coauthors and b) the top 10 coauthors with their publication count.
- 2. Similar to 1), tabulate all of the words present in the title of all of the retrieved records. Eliminate common words like "the" and "and", and print an ordered list of the top 20 occurring words.
- 3. Very often it is impossible to identify authors uniquely by their PubMed names. "Ludtke SJ" could be "Ludtke S". "Chen F" could represent dozens of different authors. While there is no perfect solution to this problem, to help assess the impact of the first issue, count the number of unique AU values, and also count the number of unique family names. Print the difference between these counts as the number of possible overlaps.
- 4. Modify the program to query based on a provided keyword rather than an author's name. Pubmed does not like overly large queries, so be sure to restrict the number of retrieved records to a reasonable number (no more than 1000). Be sure to print a warning if this limit was reached.