# Lecture 10B: Biopython



Practical Bioinformatics (Biol 4220)

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### Lecture 10A outline

- 1. Overview of Biopython
- 2. Sequence objects
- 3. Alignment objects
- 4. Other features
- 5. Lab 10A overview



Biopython is an open source Python library that provides a wide range of bioinformatics utilities. *Features include:* 

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- reading and writing sequence data
- aligning sequences
- accessing and manipulating alignments
- parsing GenBank records
- handling BLAST calls
- accessing databases
- navigating and manipulating phylogenetic trees
- displaying genome architecture
- and more

### Biopython Tutorial and Cookbook

Many code snippets in this lecture are tutorial excerpts:

http://biopython.org/DIST/docs/tutorial/Tutorial.html

### 2.2 Working with sequences

Disputably (of course!), the central object in bioinformatics is the sequence. Thus, we'll start with a quick introduction to the Biopython mechanisms for dealing with sequences, the seq object, which we'll discuss in more detail in Chapter  $\underline{3}$ .

Most of the time when we think about sequences we have in my mind a string of letters like 'AGTACACTGGT'. You can create such seq object with this sequence as follows - the ">>>" represents the Python prompt followed by what you would type in:

```
>>> from Bio.Seq import Seq
>>> my_seq = Seq("AGTACACTGGT")
>>> my_seq
Seq('AGTACACTGGT')
>>> print(my_seq)
AGTACACTGGT
```

### Parsing a fasta file with native Python

```
> Species_A
AGTCCTAGCATGTTC
> Species_B
AGTCATAGCATGTTC
> Species_C
AGTCCTAGGATGTTC
> Species_D
AGTTCTGGCATGTTC

example.fasta
```

```
f = open('example.fasta', 'r')  # open file
d = {}
lines = f.readlines()  # get list of lines
entries = ''.join(lines).split('>')  # split into entries
entries = entries[1:]  # drop lst entry, ''
for s in entries:  # loop over entries
    z = s.split('\n')  # separate entry by \n
    species = z[0].strip()  # get species name
    seq = ''.join(z[1:])  # get sequence info
    d[species] = [ nt for nt in seq ]  # store as name:seq

f.close()  # close file
print(d)  # print dict
```

```
read_fasta_native.py
```

dictionary-of-lists

### Parsing a fasta file with Biopython

```
> Species A
AGTCCTAGCATGTTC
> Species B
AGTCATAGCATGTTC
                                             from Bio import AlignIO
> Species C
                                             d = AlignIO.read('example.fasta', 'fasta')
AGTCCTAGGATGTTC
                                             print(d)
> Species D
                                                     read fasta biopython.py
AGTTCTGGCATGTTC
    example.fasta
      >>> print(d)
      SingleLetterAlphabet() alignment with 4 rows and 15 columns
      AGTCCTAGCATGTTC Species A
      AGTCATAGCATGTTC Species B
      AGTCCTAGGATGTTC Species C
      AGTTCTGGCATGTTC Species D
```

Biopython alignment data structure

# Biopython *sequence objects* behave like strings, while having expanded features for bioinformatics tasks

```
>>> from Bio.Seq import Seq
>>> my_seq = Seq('GATTACA')  # create a sequence object
>>> my_seq  # return value
Seq('GATTACA')
>>> my_seq[0:2]  # extact subsequence
Seq('GA')
>>> len(my_seq)  # get length
7
>>> Seq('r u rly a string?')  # no alphabet imposed
Seq('r u rly a string?')
```

# Sequence objects are easily cast as strings; sequences support *upper*, *find*, *count*, *etc*.

```
>>> from Bio.Seq import Seq
>>> my seq = Seq('GATTACA') # make sequence
>>> str(my seq)
                  # typecast as string
'GATTACA'
                  # change case (.upper supported)
>>> my seq.lower()
Seq('gattaca')
>>> my seq.find('TAC')
                     # find start index for subseq
>>> my seq.count('TAC') # count occurrences
>>> Seq('AAAA').count('A')  # non-overlapping count
>>> Seq('AAAA').count overlap('AA') # overlapping count
>>> (my seq.count('C')+my seq.count('G')) / len(my seq) * 100
28.57142857142857
                            # compute GC content
```

# Sequence objects are indexed and concatenated using the same syntax as is used for strings

```
>>> from Bio.Seq import Seq
>>> my_seq = Seq('GATCGATGGGCCTATATAGGA')
>>> my_seq[4:12]  # extract subsequence
Seq('GATGGGCC')
>>> my_seq[0::3]  # first codon position
Seq('GCTGTAG')
>>> my_seq[1::3]  # second codon position
Seq('AGGCATG')
>>> my_seq[2::3]  # third codon position
Seq('TAGCTAA')
>>> my_seq[:7]+Seq('NNNNNNN')+ my_seq[14:]
Seq('GATCGATNNNNNNNNTATAGGA')
```

Sequence objects provide special methods for computing properties of DNA sequences, e.g. complement, reverse-complement, and GC %

```
>>> from Bio.Seq import Seq
>>> my_seq = Seq('GATCGATGGGCCTATATAGGATCGAAAATCGC')
>>> my_seq
Seq('GATCGATGGGCCTATATAGGATCGAAAATCGC')
>>> my_seq.complement()  # return complement
Seq('CTAGCTACCCGGATATATCCTAGCTTTTAGCG')
>>> my_seq.reverse_complement() # return reverse-complement
Seq('GCGATTTTCGATCCTATATAGGCCCATCGATC')
>>> from Bio.SeqUtils import GC
>>> GC(my_seq)  # what is the GC content?
46.875
```

### Biopython sequence objects support methods to *transcribe* DNA into mRNA and to *translate* mRNA (or DNA) into AA

```
>>> from Bio.Seq import Seq
>>> cDNA = Seg('ATGGCCATTGTAATGGGCCGCTGAAAGGGTGCCCGATAG')
                                  # coding DNA sequence
>>> cDNA
Seg('ATGGCCATTGTAATGGGCCGCTGAAAGGGTGCCCGATAG')
>>> mRNA = cDNA.transcribe() # transcribe DNA into mRNA
>>> mRNA
Seq('AUGGCCAUUGUAAUGGGCCGCUGAAAGGGUGCCCGAUAG', RNAAlphabet())
>>> mRNA.translate()
                                  # translate mRNA into AA
Seq('MAIVMGR*KGAR*', HasStopCodon(ExtendedIUPACProtein(), '*'))
>>> cDNA.translate()
                                 # translate DNA into AA
Seg('MAIVMGR*KGAR*', HasStopCodon(ExtendedIUPACProtein(), '*'))
>>> cDNA.translate(to stop=True) # terminate AA at stop codon
Seg('MAIVMGR', ExtendedIUPACProtein())
>>> # consider an alternate table (default: 'Standard')
>>> cDNA.translate(table='Vertebrate Mitochondrial')
Seq('MAIVMGRWKGAR*', HasStopCodon(ExtendedIUPACProtein(), '*')
>>> cDNA.translate(table='"'Vertebrate Mitochondrial'"', to stop=True)
Seq('MAIVMGRWKGAR', ExtendedIUPACProtein())
```

# Sequence objects are translated according to a CodonTable object; default is the Standard translation table

```
>>> from Bio.Data import CodonTable
>>> standard table = CodonTable.unambiquous dna by name["Standard"]
>>> standard table.stop codons
['TAA', 'TAG', 'TGA']
>>> standard table.start codons
['TTG', 'CTG', 'ATG']
>>> standard table.forward table['GTG'] # returns AA for codon
' 77 '
>>> standard table.back table['V']
' GTT '
>>> print(standard table)
                                                          Table 1 Standard, SGC0
                                                                              TAT Y
                                                                                      TGT C
                                                              TTT F
                                                              TTC F
                                                                      TCC S
                                                                              TAC Y
                                                                                      TGC C
                                                              TTA L
                                                                      TCA S
                                                                              TAA Stop | TGA Stop |
                                                                      TCG S
                                                                              TAG Stop
                                                                                      TGG W
                                                              CTT L
                                                                      CCT P
                                                                              CAT H
                                                                                      CGT R
                                                              CTC L
                                                                      CCC P
                                                                              CAC H
                                                                                      CGC R
                                                              CTA L
                                                                      CCA P
                                                                              CAA O
                                                                                      CGA R
                                                              CTG L(s)
                                                                      CCG P
                                                                              CAG O
                                                                                      CGG R
                                                                              AAT N
                                                                                      AGT S
                                                              ATT I
                                                                      ACT T
                                                              ATC I
                                                                      ACC T
                                                                              AAC N
                                                                                      AGC S
                                                                                      AGA R
                                                              ATA I
                                                                      ACA T
                                                                              AAA K
                                                                      ACG T
                                                                              AAG K
                                                                                      AGG R
                                                              ATG M(s)
                                                                                      GGT G
                                                                      GCT A
                                                                              GAT D
                                                              GTC V
                                                                      GCC A
                                                                                      GGC G
                                                                              GAC D
                                                              GTA V
                                                                      GCA A
                                                                              GAA E
                                                                                      GGA G
                                                              GTG V
                                                                      GCG A
                                                                              GAG E
                                                                                      GGG G
```

# Biopython supports over translation using over 40 genetic code tables; choose carefully!

```
>>> from Bio.Data import CodonTable
>>> CodonTable.unambiguous dna by name.keys() # list names of codon tables
dict keys(['Standard', 'SGC0', 'Vertebrate Mitochondrial', 'SGC1',
           'Yeast Mitochondrial', 'SGC2', 'Mold Mitochondrial',
           'Protozoan Mitochondrial', 'Coelenterate Mitochondrial',
           'Mycoplasma', 'Spiroplasma', 'SGC3', 'Invertebrate Mitochondrial',
           'SGC4', 'Ciliate Nuclear', 'Dasycladacean Nuclear',
           'Hexamita Nuclear', 'SGC5', 'Echinoderm Mitochondrial',
           'Flatworm Mitochondrial', 'SGC8', 'Euplotid Nuclear', 'SGC9',
           'Bacterial', 'Archaeal', 'Plant Plastid', 'Alternative Yeast Nuclear',
           'Ascidian Mitochondrial', 'Alternative Flatworm Mitochondrial',
           'Blepharisma Macronuclear', 'Chlorophycean Mitochondrial',
           'Trematode Mitochondrial', 'Scenedesmus obliquus Mitochondrial',
           'Thraustochytrium Mitochondrial', 'Pterobranchia Mitochondrial',
           'Candidate Division SR1', 'Gracilibacteria',
           'Pachysolen tannophilus Nuclear', 'Karyorelict Nuclear',
           'Condylostoma Nuclear', 'Mesodinium Nuclear', 'Peritrich Nuclear',
           'Blastocrithidia Nuclear'])
>>> len(CodonTable.unambiguous dna by name) # how many?
43
```

### Sequence objects are immutable, but can easily be converted to and from *mutable sequences*

```
>>> from Bio.Seq import Seq
>>> my seq
                                  # create sequence
Seq('GATTACA')
>>> my seq[0] = 'C'
                                  # attempt to modify seq
Traceback (most recent call last):
  File "<stdin>", line 1, in <module>
TypeError: 'Seq' object does not support item assignment
>>> mut seg = my seg.tomutable() # convert to mutable seg
>>> mut seq[0] = 'C'
                       # modify seg successfuly
>>> mut sea
MutableSeq('CATTACA')
>>> new seq = mut seq.toseq() # convert to immutable seq
>>> new seq
Seq('CATTACA')
```

## Read fasta files using *SeqIO.parse()*; this function returns a container of iterable SeqRecords

```
>>> from Bio import SeqIO
>>> f = SeqIO.parse('example.fasta', 'fasta')
>>> for row in f:
... print( row.id + ' : ' + row.seg )
Species A : ACGCTG
Species B : ACTCTG
Species C : AGTATC
Species D : AGTCTC
>>> # convert to dict of SegRecords
>>> d = SeqI0.to dict(SeqI0.parse('example.fasta','fasta'))
>>> d
{'Species A': SeqRecord(seq=Seq('ACGCTG', SingleLetterAlphabet()), id='Speci
 'Species_B': SeqRecord(seq=Seq('ACTCTG', SingleLetterAlphabet()), id='Speci
 'Species C': SegRecord(seg=Seg('AGTATC', SingleLetterAlphabet()), id='Speci
 'Species D': SegRecord(seg=Seg('AGTCTC', SingleLetterAlphabet()), id='Speci
>>> d['Species A']
SeqRecord(seq=Seq('ACGCTG', SingleLetterAlphabet()), id='Species A', name='S
```

# Write fasta files using *SeqIO.write()*; this function expects a list of *SeqRecord* objects

```
>>> from Bio.Seq import Seq
>>> from Bio.SeqRecord import SeqRecord
>>> # make sequence records to write
>>> rec1 = SeqRecord(seq=Seq('ACGTTA'),id='Species_A',description='')
>>> rec2 = SeqRecord(seq=Seq('TCGTTA'),id='Species_B',description='')
>>> rec3 = SeqRecord(seq=Seq('ACGTGT'),id='Species_C',description='')
>>> my_records = [rec1, rec2, rec3] # list of records
>>> SeqIO.write(my_records, 'new_file.fasta', 'fasta')
```

```
$ cat new_file.fasta
>Species_A <unknown description>
ACGTTA
>Species_B <unknown description>
TCGTTA
>Species_C <unknown description>
ACGTGT
```

# Create *MultSeqAlignment* objects using *Bio.AlignIO*; access subsets of alignment with slice-indexing

```
>>> from Bio import AlignIO
>>> alignment = AlignIO.read("new file.fasta", "fasta")
>>> print(alignment) # full alignment
SingleLetterAlphabet() alignment with 3 rows and 6 columns
ACGTTA Species A
TCGTTA Species B
ACGTGT Species C
GCATGT Species D
>>> print(alignment[1:3,:])
SingleLetterAlphabet() alignment with 2 rows and 6 columns
TCGTTA Species B
ACGTGT Species C
>>> print(alignment[:,3:5])
SingleLetterAlphabet() alignment with 4 rows and 2 columns
TT Species A
TT Species B
TG Species C
TG Species D
>>> print(alignment[1:3,3:5])
SingleLetterAlphabet() alignment with 2 rows and 2 columns
TT Species B
TG Species C
```

# MultSeqAlignment objects can be constructed within Python (i.e. not parsed from file)

```
>>> from Bio.Seg import Seg
>>> from Bio.SegRecord import SegRecord
>>> from Bio.Align import MultipleSegAlignment
>>> from Bio import AlignIO
>>> alignment = MultipleSeqAlignment(
            SeqRecord(Seq("ACTCCTA"), id='seq1'),
           SeqRecord(Seq("AAT-CTA"), id='seq2'),
            SegRecord(Seg("CCTACT-"), id='seg3'),
            SegRecord(Seg("TCTCCTC"), id='seg4'),
>>> print(alignment)
Alignment with 4 rows and 7 columns
ACTCCTA seq1
AAT-CTA seq2
CCTACT- seq3
TCTCCTC seq4
>>> AlignIO.write(alignment, 'new alignment.fasta', 'fasta')
```

### Combining ideas to write a Biopython script

```
from Bio import AlignIO
from Bio.Seq import Seq
from Bio.SeqRecord import SeqRecord
from Bio. Align import MultipleSeqAlignment
ifn = 'example nt.fasta'
ofn = 'example aa.fasta'
fmt = 'fasta'
align = AlignIO.read(ifn, fmt)
x = []
for row in align:
    row.id = 'tmp ' + row.id
    s = row.seq.tomutable()
    s[0:3] = 'GGG'
    row.seg = s.toseg().translate()
    x.append(row)
new align = MultipleSeqAlignment(x)
AlignIO.write(new align, ofn, fmt)
```

### Combining ideas to write a Biopython script

```
# import libraries
from Bio import AlignIO
from Bio.Seg import Seg
from Bio.SegRecord import SegRecord
from Bio.Align import MultipleSegAlignment
ifn = 'example nt.fasta'
ofn = 'example aa.fasta'
                                 # file format
fmt = 'fasta'
# create empty list
x = []
for row in align:
                                 # loop over alignment rows
   row.id = 'tmp ' + row.id # change row id
   s = row.seq.tomutable() # let row sequence be edited
   s[0:3] = 'GGG'
                                 # overwrite first codon w/ GGG
   row.seg = s.toseg().translate()
   x.append(row)
new align = MultipleSeqAlignment(x) # create new MSA from x
AlignIO.write(new align, ofn, fmt)
```

### Lab 10B

github.com/WUSTL-Biol4220/home/labs/lab\_10B.md