Quiz Section Week 8 May 16, 2017

String handling and regular expressions

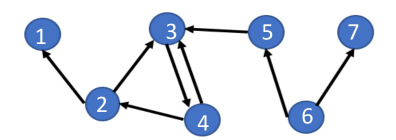
A bit more on generating random numbers

Machine Learning things to think about

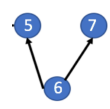
HW 6 problem 2

Subgraphs vs motifs

Motif = a pattern of connections between nodes Subgraph = an actual set of nodes and edges







How many subgraphs? Why n-2?

What would the graph have to look like to have that many subgraphs? Try for a 4 node graph

Homework programming: many of you assumed you know how many lines the file is

```
seq = lines[1]

seq2 = lines[3]
```

What if you wanted to run your code on a file with 1000 sequences?

```
for line in fin:
    if line[0] == '>':
        headers.append(line.rstrip())
    else:
        seqs.append(line.rstrip())
        seqs.append(line.rstrip())
        count = 0
    while count < len(lines):
        headers.append(lines[count])
        seqs.append(lines[count+1])
        count = count + 2</pre>
```

More on manipulating strings

Fasta sequence files are usually formatted as follows:

```
>NC_018723.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A1, Felis_catus_8.0, whole genome shotomagactic care and ca
```

>NC_018725.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A3, Felis_catus_8.0, whole genome shoto

How to read a Fasta file like this one into two lists of strings, one of headers and one of sequences?

Reading in a file with different procedure depending on content of each line

```
fin = open('cat_genome.fasta', 'r')
seqs = []
headers = []
current_seq = ""
for line in fin:
```

Reading in a file with different procedure depending on content of each line

```
fin = open('cat genome.fasta', 'r')
seqs = []
headers = []
current seq = ""
for line in fin:
     if line[0] == '>':
          headers.append(line.rstrip())
          seqs.append(current seq)
          current seq = ""
     else:
          current seq = current seq + line.rstrip()
```

Maybe we want to pull different pieces of information out of the sequence header

. . .

>NC_018725.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A3, Felis_catus_8.0, whole genor

- NCBI ID
- Species
- Isolate
- Breed
- Chromosome number

We can use *regular expressions* to analyze patterns in strings

```
import re
line = '>NC 018723.2 Felis catus isolate Cinnamon
breed Abyssinian chromosome A1, Felis catus 8.0,
whole genome shotgun sequence'
                                    But what if we don't always have
re.match('>NC', line)
                                    these characters exactly?
re.findall('chromosome', line)
re.sub('whole genome shotgun', 'WGS', line)
```

https://docs.python.org/2/library/re.html

Regular expression glossary (incomplete!)

- any character
- * repeated 0 or more times
- + repeated 1 or more times
- {n} repeated n times
- [A|B] either A or B
- [A-Z] any uppercase letter, [0-9] any numeric character
- beginning of line
- \$ end of line
- \ escape (actually search for one of the characters above)

We can use *regular expressions* to analyze patterns in strings

```
import re
line = '>NC 018723.2 Felis catus isolate Cinnamon breed
Abyssinian chromosome A1, Felis catus 8.0, whole genome
shotgun sequence'
re.match('chromosome.{3}', line)
'chromosome A1'
re.sub('[W|w]hole [G|q]enome [S|s]hotqun', 'WGS', line)
'>NC_018723.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A1, Felis_catus_8.0, WGS'
re.findall('^>[A-Z]+ [0-9]+\.[0-9]', line)
'>NC 018723.2'
                                             https://docs.python.org/2/library/re.html
```

We can use *regular expressions* to analyze patterns in strings

```
import re
line = 'ATGGCTATC'
re.match('AT[G|C]', line)
re.findall('AT[G|C]', line)
re.sub('AT[G|C]', 'QQQ', line)
```

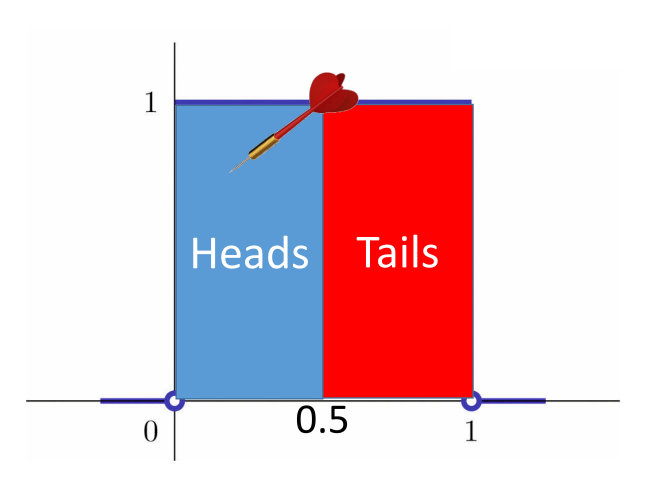
Exercise: use regular expressions to extract the annotation IDs: Felis_catus_8.0

- >NC_018723.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A1, Felis_catus_8.0, whole genome shotgun sequence
- >NC_018724.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A2, Felis_catus_7.0, whole genome shotgun sequence
- >NC_018725.2 Felis silvestrus isolate Cinnamon breed Abyssinian chromosome A3, Felis_silvestris_1.0, whole genome shotgun sequence

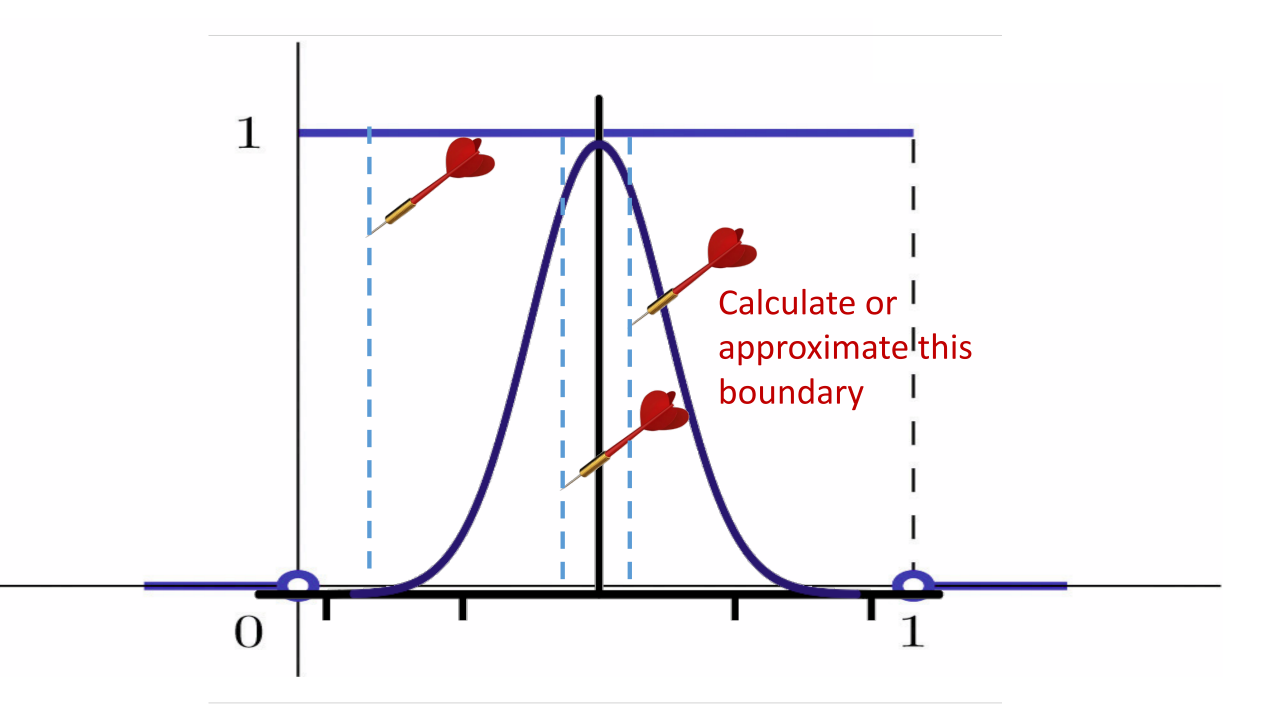
Exercise: use regular expressions to extract the breed from these headers (e.g. the word "breed" and the word after it)

- >NC_018723.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A1, Felis_catus_8.0, whole genome shotgun sequence
- >NC_018724.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A2, Felis_catus_8.0, whole genome shotgun sequence
- >NC_018725.2 Felis catus isolate Cinnamon breed Abyssinian chromosome A3, Felis_catus_8.0, whole genome shotgun sequence

More on random numbers: random.random() returns a uniformly distributed random value between 0 and 1



What if we want to sample random numbers from another distribution?



Python has additional useful random functions

https://docs.python.org/2/library/random.html

```
>>> random.choice(['apple','banana','pear'])
'pear'
>>> random.randint(10,100)
55
>>> random.gauss(0,1) #mean 0, std dev 1
-0.1175
```

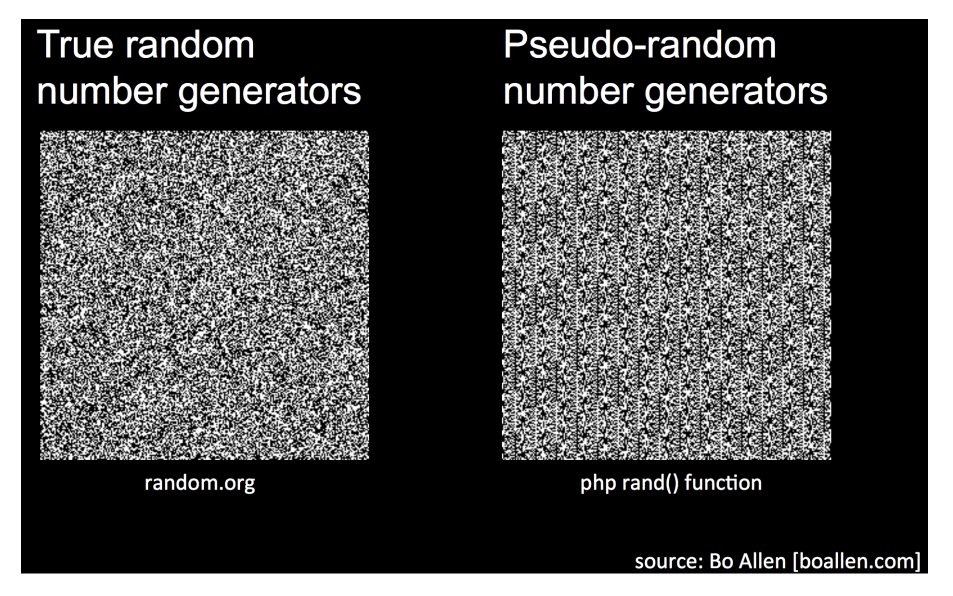
What does it mean to generate random numbers anyway?

```
int getRandomNumber()
{
    return 4; // chosen by fair dice roll.
    // guaranteed to be random.
}
```

Computation is deterministic!

Most random number generating algorithms are pseudo-random

What are some truly random processes?



Pseudo-random number generators require a "starting point" called a *seed*

- a seed lets us initialize the random number generator
 - If you know the seed, the sequence of numbers is predictable and fixed
 - If you don't know the seed, the sequence is hopefully unpredictable (but still deterministic)

```
>>> random.seed(number)
```

 You can set the seed to be something unpredictable, like a function of the time at which the code is running

Why might we want predictable "random" numbers? Why might we want unpredictable "random" numbers?