Genomics Assignment 1

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Part 1: Theory

1. Central dogma: $DNA \rightarrow RNA \rightarrow protein$ DNA makes RNA makes protein.

2. a Escherichia Coli

Length: 4.6 Mb, Chromosomes: 1

b Yeast (Saccharomyces cerevisiae)

Length: 12.5 Mb, Chromosomes: 16

3. Since the probabilities of A, C, T, G are independent and identically distributed, we can apply multinomial model.

$$P(s) = \prod_{i=1}^{n} p(s(i))$$

Given $p(A) = \frac{1}{2}$, $p(C) = \frac{1}{6}$, $p(G) = \frac{1}{6}$ and $p(T) = \frac{1}{6}$

Probability of sequence 'ACGTACGTACGT'

$$= p(A) * p(C) * p(G) * p(T) * p(A) * p(C) * p(G) * p(T) * p(A) * p(C) * p(G) * p(T)$$

$$= 1/2 * 1/6 * 1/6 * 1/6 * 1/2 * 1/6 *$$

4. Probability of an ORF of length 150 codons in a random sequence, requires that we find a start codon (ATG) followed by none of the 3 stop codons for next 149 codons. We'll assume **the distribution of codons to be uniform** in which case it is 3/64 probability to pick a stop codon and 61/64 to pick a non-stop codon (total codons = 64).

This can be calculated as P(run of 148 codons) = $(\frac{61}{64})^{148} = 0.00082$

There is a 0.00082 probability of a codon of length 150 which means we can conclude with 99.92% confidence that it is highly significant.

- 5. Homology refers to similarity between DNA sequences which is a result of common ancestors and not by chance. To measure similarity:
 - first perform sequence alignment using global alignment or local alignment
 - use the Needleman-Wunch algorithm (for global alignment) or Smith-Waterman algorithm (local alignment) to find the best alignment score (uses scoring function for getting alignment score).
 - compare with score obtained from alignment of random sequences of similar type to determine statistical significance. Probability of the two sequences alignment score is calculated (scores equal and above / number of permutations) which is the p-value. If the p-value is less than 0.05 (or the confidence we decide), the two sequences are considered homologous with 95% confidence.

1

Part 2: Practical

Note: all code in ass1_1919523.m

6. Mitochondrial genome of Pan troglodytes (chimpanzee)

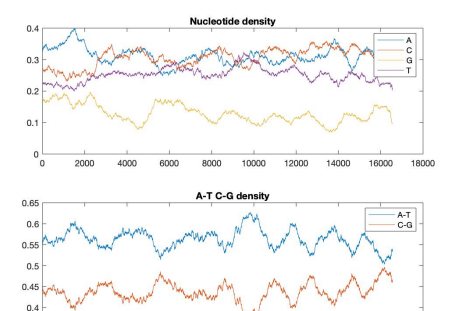
Scientific name: Pan troglodytes Accession number: NC_001643 Sequence length: 16554

Base count: A: 5154 C: 5099 G: 2133 T: 4168

GC %: 43.7, Size: 0.02 Mb Nucleotide density shown below:

0.35

2000



```
Code:
mito_gbk = getgenbank('NC_001643'); %chimp
chimp = mito_gbk.Sequence;
mitochondria_length = length(chimp)
first_300_bases = seqdisp(chimp(1:300))
figure
ntdensity(chimp)
```

4000

6000

8000

10000

12000

14000

16000

18000

7. Potential protein coding genes in a Chimpanzee mitochondrial genome can be found using:

```
orf = seqshoworfs(chimp, 'MINIMUMLENGTH', 3,
    'geneticcode', 2, 'frames', 'all');
```

Genetic code for Vertebrate Mitochondrial is 2. All frames are considered.

Threshold for the ORFs: random sequence is generated from the chimp genome by 16554 random permutations and the length of the longest ORF is selected as threshold. The possibility of seeing an ORF of length longer than the random ones makes it a good candidate to be a potential protein encoding gene, since it will be unlikely to be by chance.

```
orf1 = seqshoworfs(chimp(randperm(length(chimp))), 'MINIMUMLENGTH', 3, 'geneticcode', 2, 'frames', 'all');
```

```
ORFLength1 = [];
for i = 1:6
    for j = 1:length(orf1(i).Stop)
        ORFLength1 = [ORFLength1; orf1(i).Stop(j)+2 - orf1(i).Start(j)];
    end
end

max_threshold=max(ORFLength1)
n_max=length(find(ORFLength>=max_threshold))
```

Since a codon is made up of 3 nucleotide, by specifying MINIMUMLENGTH as max_threshold/3, we get 12 potential genes through this process. In real there are 13 genes. We could reduce the threshold in order to detect potential genes of shorter length. (Note: running the code multiple times may result in answers between 10-13. This is because we are using random permutations which change every time, changing the threshold.)

8. The first potential protein encoding gene can be extracted using

```
protein = chimp(orf(1).Start(1):orf(1).Stop(1)+2)
```

Test statistic: ORF length calculated as length(protein)/3 = 317 codons

Null hypothesis: the chosen ORF is generated by a random process.

Significance level = 0.005

Using uniform distribution assumption, the probability of seeing a codon of length 317 is $\frac{61}{64}^{315} = 0.00000027$ (p-value).

The p-value is much smaller than the significance level. Therefore, there's a 99.999973% confidence of this being a significant gene.

9. To convert sequence to aminoacid we use

```
aminoacid = nt2aa(sequence, 'geneticcode',2)
```

Amino acid (first 50) = 'MTNLLLLIVPILIAIAFLMLTERKILGYIQLRKGPNIVGPYGLLQPFADA'

Common chimpanzee and 5 closest organisms

Name	Taxonomic Name	Protein	Identity score
Common chimpanzee	Pan troglodytes	NADH-ubiquinone oxi-	618
		doreductase chain 1	
Nigeria-Cameroon chimpanzee	Pan troglodytes ellioti	NADH dehydrogenase	616
		subunit 1	
Western chimpanzee	Pan troglodytes verus	NADH dehydrogenase	615
		subunit 1	
Eastern chimpanzee	Pan troglodytes schweinfurthii	NADH dehydrogenase	615
		subunit 1	
Bonobo	Pan paniscus	NADH dehydrogenase	613
		subunit 1	
Central chimpanzee	Pan troglodytes troglodytes	NADH dehydrogenase	613
		subunit 1	

Multialignment can be generated using

```
seq1 = getgenpept('Q9T9W3', 'SequenceOnly', true);
seq2 = getgenpept('AIV00479', 'SequenceOnly', true);
seq3 = getgenpept('AEQ36262', 'SequenceOnly', true);
seq4 = getgenpept('ANQ92411', 'SequenceOnly', true);
seq5 = getgenpept('AMB65312', 'SequenceOnly', true);
seq6 = getgenpept('AJO25286', 'SequenceOnly', true);
seqs = {seq1, seq2, seq3, seq4, seq5, seq6};
showalignment(multialign(seqs))
```

```
MPMTNLLLLIVPILIAMAFLMLTERKILGYMQLRKGPNIVGPYGLLQPFADAMKLFTKEPLKPSTSTITLYITAPTLALTIALLLWTPLPMF
--MTNLLLLIVPILIAMAFLMLTERKILGYMQLRKGPNIVGPYGLLQPFADAMKLFTKEPLKPSTSTITLYITAPTLALTIALLLWTPLPMF
TPMTNLLLLIVPILIAMAFLMLTERKILGYMQLRKGPNIVGPYGLLQPFADAMKLFTKEPLKPSTSTITLYITAPTLALTIALLLWTPLPMF
--MTNLLLLIVPILIAMAFLMLTERKILGYMQLRKGPNIVGPYGLLQPFADAMKLFTKEPLKPSTSTITLYITAPTLALTIALLLWTPLPMF
--MTNLLLLIVPVLIAMAFLMLTERKILGYMQLRKGPNIVGPYGLLQPFADAMKLFTKEPLKPSTSTITLYITAPTLALTIALLLWTPLPMF
--MTNLLLLIVPILIAMAFLMLTERKILGYMQLRKGPNIVGPYGLLQPFADAMKLFTKEPLKPSTSTITLYITAPTLALTIALLLWTPLPMF
```

We can see that the sequence discovered using p-value analysis is an actual protein found in many different types of chimpanzees (even human). This shows that we can detect genes accurately with probability and statistical analysis.

10. Human mitochondrion genome

Accession no: NC_012920

Name: Homo sapiens mitochondrion

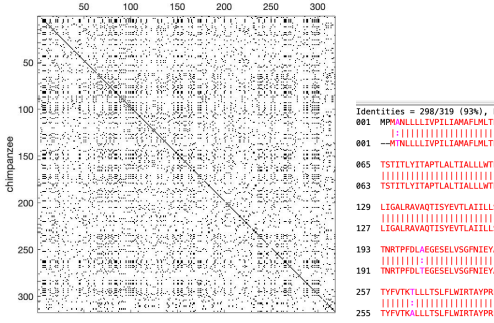
GC%: 44.4

Can be seen from BLASTing the discovered protein that it is also found in human mitochondrion. The protein is NADH dehydrogenase subunit 1 (mitochondrion) AFF91323. We can get the sequence using

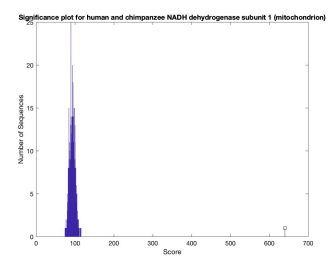
humanProtein = getgenpept('AFF91323', 'SequenceOnly', true);

human

A dotplot of sequence matches shows an almost perfect match. The alignment (BLOSUM50) is shown below



Global alignment gives a score of 640 for scoring matrix BLOSUM50 and 393.6 for BLOSUM30. A significance test gives a p-value of 0.



This is expected since humans and chimpanzees are descended from same ancestors, primates and therefore their genes can be homologous.

11. First the human mitochondrion genome is processed to find significant protein coding regions following the same process as the chimpanzee genome in answer 7. This results in 11 sequences, and we had 12 sequences from chimpanzee. They are all subjected to global alignment that yields 132 scores. The alignments with score more than 100 are further analysed.

Test statistic: global alignment score compared with scores of 1000 random permutations of similar type.

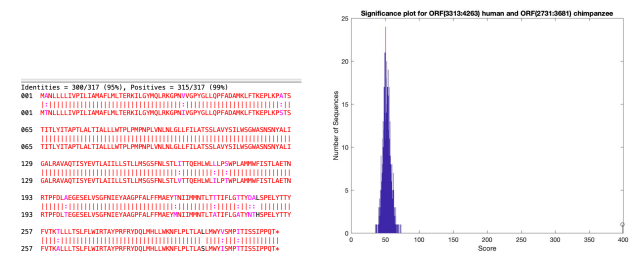
Null hypothesis: the alignment is generated by a random process.

Significance level = 0.005

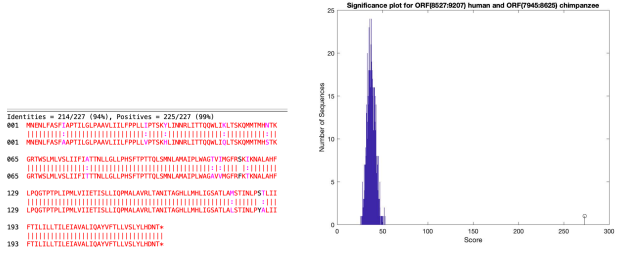
Finding: Doing a statistical test yields all the alignments to be highly significant since **the p-value is 0** for all. None of the random permutations generate a score comparable to the alignment score. Since humans and chimpanzees do share ancestors, we end up finding multiple homologous genes.

Below are all the results with alignments and significance plot for three:

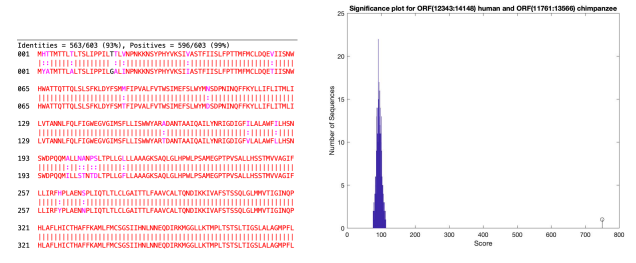
(a) ORF in position (3313:4263) in human is homologous to ORF in position (2731:3681) in chimpanzee.



(b) ORF in position (8527:9207) in human is homologous to ORF in position (7945:8625) in chimpanzee.



(c) ORF in position(12343:14148) in human is homologous to ORF in position (11761:13566) in chimpanzee.



- (d) ORF in position (7586:8269) in human is homologous to ORF in position (7003:7686) in chimpanzee.
- (e) ORF in position(10760:12184) in human is homologous to ORF in position (10178:11602) in chimpanzee.
- (f) ORF in position(14747:15907) in human is homologous to ORF in position (14165:15325) in chimpanzee.
- (g) ORF in position (4578:5513) in human is homologous to ORF in position (4191:4931) in chimpanzee.
- (h) ORF in position (5904:7445) in human is homologous to ORF in position (5321:6862) in chimpanzee.
- (i) ORF in position (9207:10055) in human is homologous to ORF in position (8625:9473) in chimpanzee.

The code in ass1_1919523.m can be used to view all the alignments and significance plots.

12. We can use BLAST to find alignments easily as done in question 10. A way to do it manually is described in question 11. First the significant potential protein encoding genes are extracted using random permutation significant test. All the sequences are compared to each other and scores are obtained using nwalign. We discard low scores (threshold 50 or 100) and remaining are tested for significance. If the p-value is less than 0.05, the sequence pair may be homologous.

The procedure will yield results shown in 11.

Although this process can be slow for large number of matching genes, it gives accurate results.