## Math 5233 practice exam.

- (1) (a) What are mitochondria?
  - (b) Why are they important for evolutionary studies?
  - (c) Why might a scoring matrix derived from mitochondrial sequences differ from one derived from nuclear sequences?
- (2) Decide whether each of the alignments below are from protein-coding regions of DNA or not. Explain your reasoning.

Tapir	TCACCCACCATTAAAAGTATAGGAGATAGAAATTTT
Sable	TTAACCACACATTACA-GTATAGGAGATAGAAATTCT
RiverDolphin	TCACACCCCTTCTAAA-GTATAGGAGATAGAAATTTAAACACC
Human	TTACCCAAATAAA-GTATAGGCGATAGAAATTGAA
Siamang	TCACCCACATAAA-GTATAGGCGATAGAAATTATTA
Baboon	TAACCCATATTAA-GTATAGGCGATAGAAATCTTA

Figure 1. A

Sheep	TCCCGGAGTTCAAGAAATCATCTCACACAGCGGTCTCGGAC
Dog	TCC <mark>C</mark> GGAGTTCAAGAAATCATCTCACAC <mark>A</mark> GC <mark>A</mark> GT <mark>C</mark> TC <mark>A</mark> GAC
Human	TCTCGGAGTTCAAGAAATCATCTCACACGGCCGTCTCAGAC
Mole	TCCAGAAGTTCAAGAAATCATCTCACACAGAGTCTCAGAC
Chicken	TCC <mark>AGG</mark> AGTTCAAGAAATC <mark>T</mark> TC <mark>A</mark> CA <b>T</b> AC <b>T</b> GC <b>A</b> GT <mark>A</mark> TC <mark>A</mark> GAT
AfricanFrog	TCC <mark>AAA</mark> AG <mark>C</mark> TCAA <mark>TT</mark> AAATCATC <mark>CTC</mark> TCACAC <mark>A</mark> GCAGT <b>T</b> TC <mark>C</mark> GAC

Figure 2. B

- (3) Describe how a progressive pairwise alignment of the sequences ACCCGC, ACCGC, ACTCTC, and ACGCGC (in that order) could generate a sub-optimal multiple alignment.
- (4) Using a scoring system of +5 for a match, -3 for a mismatch, and -4 for a gap, use the Smith-Waterman algorithm to fill in a dynamic programming table and find the best local alignment between the sequences TAGCG and TAAGCG. If there are ties, list all of the best alignments.
- (5) Suppose two different types of nucleotide sequences are used to study the evolutionary distances between humans, chimps (Pan troglodytes), and mice (Mus musculus). The first sequence type is mRNA of the H4 histone protein, and the fraction of sites that differ between the species are: human-mouse: 40/356, human-chimp: 5/356, and chimp-mouse: 43/364. The second sequence type is the D-loop region of the

mitochondrion, and for that the fractional differences are: human-mouse: 527/1092, human-chimp: 138/1107, and chimp-mouse: 531/1088.

- (a) What would you conclude from each type of sequence if you used the Jukes-Cantor model?
- (b) What are some possible problems with using each type of sequence with the Jukes-Cantor model? What sort of third sequence type would be good to use to check your results?
- (6) Suppose that P and Q are probability distributions on N items that is,  $P = \{p_1, \ldots, p_N\}$  and  $Q = \{q_1, \ldots, q_N\}$  are both lists of N numbers between 0 and 1, which sum to 1.
  - (a) Show that the difference in entropy

$$H_Q - H_P = -\sum_{i=1}^{N} q_i \log q_i + \sum_{i=1}^{N} p_i \log p_i$$

is equal to the relative entropy of P relative to Q

$$H(P||Q) = \sum_{i=1}^{N} p_i \log (p_i/q_i)$$

if Q is the uniform distribution (each item has a 1/N probability).

- (b) If N = 2, for a fixed distribution P for which distributions Q will  $H_Q H_P = H(P||Q)$ ?
- (c) Extra Credit: same as the previous question, for N=3. Describe the situation as completely as possible.
- (7) Suppose we view each of the approximately 30,000 proteins in the human genome as a message from the genome to the cell, and suppose that their frequency of expression follows Zipf's law: the most frequent protein is twice as likely as the second-most frequent protein, three times as likely as the third-most frequent protein, and so on.
  - (a) What is the entropy of this distribution?
  - (b) How does it compare with the entropy of the uniform distribution for 30,000 items?
- (8) For the DNA scoring matrix which has  $S_{ii} = 3$  and  $S_{ij} = -5$  for  $i \neq j$ , if you assume a uniform background distribution of nucleotides  $P_A = P_C = P_G = P_T$  calculate the implied transition matrix  $Q_{ij}$  and the (relative) entropy  $\sum_{i,j} Q_{ij} \log_2(\frac{Q_{ij}}{P_i P_j})$ .