## Homework #2

## Please present 4 significant figures in your final answers for probabilities. Also, make sure to explain your thought process as if the reader is one of your classmates.

**1. (7 points)** Mutations to DNA occurs when the cell makes mistakes as it makes copies of its DNA. Suppose the rate of mutation per base, per generation is about  $2.5 \times 10^{-8}$  and there are about  $3.2 \times 10^{9}$  sites in the human genome. Assume that mutations at different sites are independent of each other.

(a) Calculate the rate  $\lambda$  at which mutations occur in a cell with one copy of the human genome.

Answer: λ =(3.2 x 10<sup>9</sup>)(2.5 x 10<sup>-8</sup>) = 80

(b) If one observes the cell for 1 generation what is the probability that at least 80 and at most 85 mutations will occur?

Answer: Let X~Pois(80) P(85<= X <=80) = P(X=80) + P(X=81) + P(X=82)+ P(X=83)+ P(X=84)+ P(X=85) = 0.04456 + 0.04401 + 0.04293 + 0.04138 + 0.03941 + 0.03709 = 0.24938

**2. (7 points)** For some reason a lot of people hate raisins in cookies, but you love them. So, when you were baking 100 cookies for a party, you only put in 100 raisins in the batter. Your best friend who absolutely hates raisins picks 3 cookies at random. What is the probability that at least one of the cookies is raisin-free? (Hint: Use both Poisson and Binomial)

Answer: Expected number of raisins per cookie  $\lambda$ =1. Probability that a randomly picked cookie has no raisins is P(X=0) = e<sup>- $\lambda$ </sup> = e<sup>-1</sup> = 0.3679

Probability that at least one cookie out of 3 has no raisins is 1 – P(Y=0) = 1 – <sup>3</sup>C<sub>0</sub> (0.3679)<sup>0</sup> (0.6321)<sup>3</sup> = 1- 0.2526 = 0.7474

**3. (11 points)** Sequencing technologies can only "read" short fragments from a genome. Given that the process through which the sequences are generated is random, it is possible that certain parts of the genome will remain uncovered unless an impractical amount of sequences are generated.

We know that the size of the human genome is  $3x10^9$  bp. Now a new human genome has been sequenced and it's randomly covered by 30 million reads (read length is 300 bp). We assume that the number of times a base in the human genome is covered follows a Poisson distribution.

(a) What is the probability that a particular base is covered by at least one read?

Answer: the average time a base is covered is  $30 \times 10^6 \times \frac{300}{3 \times 10^9} = 3$ . The probability that a particular base is not covered by any read is  $P(X = 0) = e^{-3} = 0.04979$  So, P(X>=1) = 1 - 0.04979 = 0.95021

(b) Calculate the number of contigs Answer:  $N_{contigs} = N * e^{-\lambda} = 30 \times 10^6 * e^{-3} = 1.494 \times 10^6$ 

(c) What is the average le	ngth of a contig?
Answer: G <sub>covered</sub> / N <sub>contigs</sub>	= G * P(X>0) /1.494 x 10 <sup>6</sup>
	= 3x10 <sup>9</sup> * 0.95021 / 1.494 x 10 <sup>6</sup>
	= 1908

**4. (11 points)** Assume X is normally distributed with a mean of 4 and a standard deviation of 2.

(a) Determine P(X > 2)

Answer:  $P(X > 2) = 1 - P(X \le 2) = 1 - 0.15866 = 0.84134$ 

(b) Determine P(0 < X < 7) Answer: P(0 < X < 7) = P(X < 7) - P(X < 0) = 0.93319 - (1 - 0.9772) = 0.9104

(c) If P(x < X < 7) = 0.2, what is x? Answer: If P(x < X < 7) = 0.2, then P(X < x) = 0.73319. By looking at the table, we know that x = 5.24498.

**5.** (7 points) The height of people is often assumed to be normally distributed. Let the mean height of men in the US is  $\mu$ = 1.77m, and standard deviation is  $\sigma$  = 0.08.

- (a) What is the probability that a randomly selected man is taller than 1.83m? Answer: The probability that a randomly selected man is taller than 1.83m is  $P(X > 1.83) = 1 - P(X \le 1.83) = 1 - 0.77337 = 0.22663$ .
- (b) What is the probability that if 10 men are randomly selected, at least 3 of them are taller than 1.83m? Let Y~Binom(10, 0.22663)

P(Y>=3)	= 1 - (P(Y=0) + P(Y=1) + P(Y=2))
	= 1 – (0.59659)
	= 0.40341

**6. (7 points)** Measurement error that is normally distributed with a mean of zero and a standard deviation of 0.5 grams is added to the true weight of a sample. Then the measurement is rounded to the nearest gram. Suppose that the true weight of a sample is 156.5 grams.

(a) What is the probability that the rounded result is exactly 157 grams?

Answer: Let us denote X as measurement and  $\epsilon$  as error. Then, we have  $X = 156.5 + \epsilon$ . The probability that the rounded result is exactly 157 grams is  $P(156.5 \le X < 157.5) = P(X < 157.5) - P(X < 156.5) = P(\epsilon < 1) - P(\epsilon < 0) = 0.97725 - 0.5 = 0.47725$ 

(b) What is the probability that the rounded result is 155 grams or lesser? Answer:  $P(X < 155.5) = P(\epsilon < -1) = 1 - P(\epsilon < 1) = 0.02275$