# Lab 1 - Gr. 14 - Bioinformatics (732A93)

Andreas Stasinakis (andst745), Hector Plata (hecpl268), Julius Kittler (julki092), Mim Kemal Tekin (mimte666), Stefano Toffol (steto820)

# Contents

Assignment 1	2
Question 1: Hardy-Weinberg equilibrium	2
Question 1.1	2
Question 1.2	3
Assignment 2 : Exploring a genomic sequence	3
Question 2.1	3
Question 2.2	4
Question 2.3	4
Question 2.4	
Question 2.5	
Assignment 3: Exploring a genomic sequence	7
Question 3.1	7
Question 3.2	7
Question 3.3	
Question 3.4	8
Question 3.5	8
3.6	11
Appendix	11

# Assignment 1

# Question 1: Hardy-Weinberg equilibrium

# Question 1.1

We define the following probability space

$$(\Omega, \mathcal{F}, P)$$

Where,

$$\Omega = \{(A, A), (A, a), (a, A), (a, a)\}$$

We also define a probability measure P such as

$$P(X,Y)$$
$$X,Y \in \{A,a\}$$

So P(X,Y) is the probability that an allele is (X,Y).

By definition, p is the proportion of A's in the allele population and q is the proportion of a's in the allele population, so:

$$P(A) = P(A, A) + P(A, a) + P(a, A) = p$$

The same applies to P(a).

$$P(a) = P(a, a) + P(a, A) + P(A, a) = q$$

The fact that we assume random mating means that X and Y are IID, which entails the following:

$$P(A, A) = P(A) * P(A) = p^{2}$$
  
 $P(a, a) = P(a) * P(a) = q^{2}$   
 $P(A, a) = P(a, A) = P(A) * P(a) = pq$ 

The probability of an allele of it being a heterozygote is:

$$P(A, a) + P(a, A) = 2pq$$

$$P(\Omega) = P(A, A) + P(a, a) + 2P(A, a) = p^2 + q^2 + 2pq = 1$$

Thus, we show that by random mating the proportion of A's and a's is the same in the offsprings and the Hardy-Weinberg equilibrium is obtained and can't deviate from it.

### Question 1.2

We now look at the MN blood group, that has two possible co-dominating alleles,  $L^M$  (denoted M) and  $L^N$  (denoted N). In a population of n = 1000 Americans of Caucasian descent, the following genotype counts were observed:

- $n_{MM} = 357$  individuals were MM homozygotes;
- $n_{MN} = 485$  individuals were MN heterozygotes;
- $n_{NN} = 158$  individuals were NN homozygotes;

The relatives proportion, obtained by dividing the genotype counts by the total population, are 0.357, 0.485 and 0.158 respectively. According to the Hardy–Weinberg principle, we expect these quantities to follow the proportions  $p^2$ , 2pq and  $q^2$ , where p and q are the proportions of N and M in the alleles. In formulas:

$$p = \frac{n_{MM}}{n} + \frac{1}{2} \cdot \frac{n_{MN}}{n} = 0.357 + \frac{1}{2} \cdot 0.485 = 0.5995$$

$$q = \frac{n_{NN}}{n} + \frac{1}{2} \cdot \frac{n_{MN}}{n} = 0.158 + \frac{1}{2} \cdot 0.485 = 0.4005$$
(1)

With these empirical quantities we formulate what would the population look if it was on equilibrium and compare them with the real proportions ( $p_{MM} = 0.375, p_{NN} = 0.158, p_{MN} = 0.485$ ):

$$p_{MM}^0 = p^2 = 0.3594002$$
  

$$p_{NN}^0 = q^2 = 0.1604002$$
  

$$p_{MN}^0 = 2pq = 0.4801995$$

Performing a chi–square goodness of fit test (chisq.test() function in R) results in p-value = 1. This result shows that there is not enough statistical evidence to reject the hypothesis that the population is in a Hardy-Weinberg equilibrium.

```
## Warning in chisq.test(c(0.357, 0.485, 0.158), p = c(p_2, pq, q_2)): Chi-
## squared approximation may be incorrect
##
## Chi-squared test for given probabilities
##
## data: c(0.357, 0.485, 0.158)
## X-squared = 9.9938e-05, df = 2, p-value = 1
```

# Assignment 2: Exploring a genomic sequence

Note that, by convention, a *coding strand* is used when displaying a DNA sequence. "A coding strand, is the segment within double-stranded DNA that runs from 5' to 3', and which is complementary to the antisense strand of DNA, or template strand, which runs from 3' to 5" (https://en.wikipedia.org/wiki/Sense strand).

### Question 2.1

We go to the following link https://www.ncbi.nlm.nih.gov/nuccore/CU329670, scroll down to "Features", click on "CDS" to see the first 5662 nucleotides of the sequence.

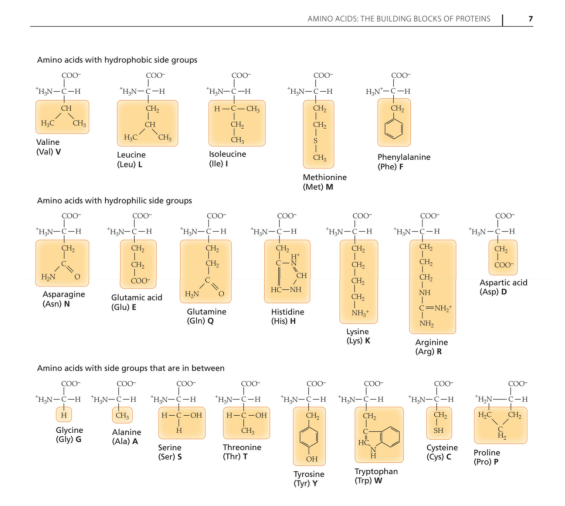
The protein product is: RecQ type DNA helicase

# Question 2.2

Note that proteins are made up of 20 different amino acids (linked together). Amino acids are molecules composed of an alpha carbon, a carboxyl group, an amino group and a side chain. The side chain is what makes an amino acid unique (see p. 5, Concepts in Bioinformatics and Genomics). The 20 different amino acids can be found in the picture below.

In our case, the first four amino acids are:

- 1. M (= **Methionine**, coded by ATG, representing the starting sequence)
- 2. V (= Valine)
- 3. V = Valine
- 4. A (= Alanine)



### Question 2.3

Note that nucleotides code for amino acids. They are molecules composed of a sugar, a phosphate and a base. The part of nucleotides that distinguishes them is the base.

There are four possible bases: A (Adenine), C (Cytosine), G (Guanine), T (Thymine)

TABLE 1-1. SINGLE-LETTER ABBREVIATIONS USED FOR DNA NUCLEOTIDE SEQUENCES

ONE-LETTER ABBREVIATION	NUCLEOTIDE NAME	BASENAME	CATEGORY
А	Adenosine monophosphate	Adenine	Purine
С	Cytidine monophosphate	Cytosine	Pyrimidine
G	Guanosine monophosphate	Guanine	Purine
Т	Thymidine monophosphate	Thymine	Pyrimidine
N	Any nucleotide	Any base	NA
R	A or G	A or G	Purine
Υ	CorT	C or T	Pyrimidine
- or *	_	_	Gap

Specific combinations of these nucleotides code for specific amino acids. In genetic code, three nucleotides (called codon) always code for one amino acid. E.g. ATG codes for met (Methionine) which is a starting point of a protein. Which codos code for which amino acids is shown in the overview below:

1st		2nd base							
base TTT TTC		Т		С		Α	G		base
	TTT	(D) (E) D)	TCT		TAT	/T 00 T	TGT	(0. (0) 0	Т
	TTC	(Phe/F) Phenylalanine	TCC	(Ser/S) Serine	TAC (Tyr/Y) Tyrosine	TGC	(Cys/C) Cysteine	С	
Т	TTA		TCA		TAA[B]	Stop (Ochre)	TGA[B]	Stop (Opal)	Α
	TTG		TCG		TAG[B]	Stop (Amber)	TGG	(Trp/W) Tryptophan	G
	CTT	TC (Leu/L) Leucine	CCT		CAT	(His/H) Histidine	CGT	(Arg/R) Arginine	Т
	СТС		ccc	(Pro/P) Proline	CAC		CGC		С
С	CTA		CCA		CAA	(Gln/Q) Glutamine	CGA		Α
	CTG		CCG		CAG		CGG		G
	ATT		ACT	(Thr/T) Threonine	AAT	(Asn/N) Asparagine	AGT	(Ser/S) Serine	Т
	ATC (Ile/I) Isoleucine	(Ile/I) Isoleucine	ACC		AAC		AGC		С
Α	ATA		ACA		AAA	// // // // // // // // // // // // //	AGA	(Arg/R) Arginine	Α
	ATG[A]	(Met/M) Methionine	ACG		AAG	(Lys/K) Lysine	AGG		G
	GTT		GCT		GAT	/4 /D) 4 - 10 - 11	GGT		Т
GTC		GCC		GAC	(Asp/D) Aspartic acid	GGC	(Ch.(C) Chreins	С	
G	G GTA	(Val/V) Valine GC	GCA		GAA	(Glu/E) Glutamic acid	GGA	(Gly/G) Glycine	Α
			GCG		GAG		GGG		G

Saving the nucleotide sequence from GenBank:

The complete nucleotide sequence of the coding strand from GenBank (https://www.ncbi.nlm.nih.gov/nuccore/CU329670.1?from=1&to=5662) was saved in FASTA format as **2.3\_nucleotid-sequence.FASTA**. Note that only the last 12 characters AGCGACGACCAT actually correspond to the amino acids MVVA if the reverse compliment is taken (see 2.4).

Using backtrack to obtain the amino acid sequence from the protein sequence:

We used backtrack to obtain the amino acid sequence corresponding to the protein sequence MVVA (https://www.ebi.ac.uk/Tools/st/emboss\_backtranseq/). After pasting MVVA and selecting "Schizosaccharomyces pombe (CAI equivalent)", the following sequence is returned: **ATGGTCGTCGCT** 

# Question 2.4

The above mentioned obtained coding strand sequence ATGGTCGCT does not exist in the provided nucleotide sequence (that was saved in FASTA format as 2.3\_nucleotid-sequence.FASTA) since this sequence is not found when searching the file.

#### Option 1

However, we can modify the displayed sequence to get what we are looking for. On GenBank, under "Display options", one has to select both "Show sequence" and "Show reverse complement" (https://www.ncbi.nlm.nih.gov/nuccore/CU329670.1?from=1&to=5662). Afterwards, the displayed nucleotid sequence under "ORIGIN" starts with ATGGTCGTCGCT which codes for our amino acids MVVA.

#### Option 2

Alternatively, one can take the last 12 characters from the original nucleotid sequence (the one in the FASTA file, i.e. the sequence before selecting "Show reverse complement" on GenBank). These characters are: AGCGACGACCAT. Copy this string and paste it here: http://arep.med.harvard.edu/labgc/adnan/projects/Utilities/revcomp.html Then click "reverse complement". The result is again the sequence that we are looking for: ATGGTCGTCGCT which codes for our amino acids MVVA.

#### Conclusion

ATGGTCGTCG is exactly what we got in 2.3 using backtranseq to obtain the amino acid sequence from the protein sequence MVVA. The only tricky thing here was that we needed to take the reverse complement of the correct characters from the nucleotid sequence (namely the last 12 characters).

### Question 2.5

Number range that corresponds to these amino acids (protein sequence):

In the saved FASTA file, the *last 12 characters* AGCGACGACCAT correspond to the amino acids MVVA (after taking the reverse complement). The number range corresponding to them is 5651 to 5662.

If we had selected "Show reverse complement" on GenBank before, then the first 12 characters ATG-GTCGTCGCT would correspond to the amino acids MVVA. The number range would then be 1 to 12.

Stop codon in the nucleotide sequence:

Note that there are three possible stop codons: TAA, TAG, TGA. Still, it is not easy to identify the stop codon manually. However, one can use: https://www.ncbi.nlm.nih.gov/orffinder and paste the complete (reverse complement) nucleotide sequence in FASTA format. With the default parameters, this is automatically translated to the correct amino acid sequence. Also, after scolling down a bit, one can see that the last stop happens at 5661. We find TC from 5661 to 5662 (and at 5662, we have the last nucleotid). From 5658 to 5660, we have TGA which is one of the above mentioned stop codons. We can conclude that the stop codon in the nucleotid sequence is hence TGA from 5658 to 5660 (looking at the reverse complement nucleotid sequence).

Chromosome on which the genomic sequence lies:

We can see on GenBank (in the header under the definition as well as in the features under source), that this genomic sequence lies on the chromosome 1 of schizosaccharomyces pombe.

# Assignment 3: Exploring a genomic sequence

# Question 3.1

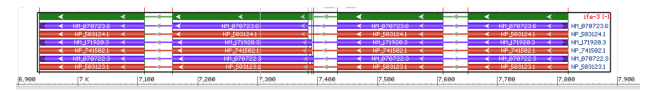
Caenorhabditis elegans is a free-living, nematode worm, which lives in soil environments all over the world. it is divided in two sexes:

- i) male
- ii) self-fertilizing hermaphrodite

Finally, it has all human sensations, such as taste and smell, despite the fact that is has no eyes. (https://en.wikipedia.org/wiki/Caenorhabditis elegans)

In general C.elegans has a similar construction and characteristics as humans. It is also important that the genes, which are responsible for human's evolution, had similar ancestor with C.elegans. Therefore, instead of making experiments on humans, which most of the time is expensive and difficult, scientist can work with C.elegans. That is the reason why studying C.elegans biology is crucial for scientific field. (http://www.people.ku.edu/~erikl/Lundquist\_Lab/Why\_study\_C.\_elegans.html)

# Question 3.2



This diagram shows exons and introns. We have 4 exons in our searching query. Our searching query found between 6529 and 8028. Our exons and introns:

• introns1: 6529 - 6935

• exon1: 6936 - 7110

• introns2: 7111 - 7157

• exon2: 7158 - 7393

• introns3: 7394 - 7432

• exon3: 7433 - 7609

• introns4: 7610 - 7650

• exon4: 7651 - 7818

• introns5: 7819 - 8028

# Question 3.3

We can find all the information needed from the summary:

### Caenorhabditis elegans chromosome V

Sequence ID: NC 003283.11 Length: 20924180 Number of Matches: 1

Range 1: 6529 to 802	8 GenBank G	▼ Next	Match 🛕 Previous Match	
Score	Expect	Identities	Gaps	Strand
2771 bits(1500)	0.0	1500/1500(100%)	0/1500(0%)	Plus/Plus

The numbering of the sequences in the alignment is 1500. The identities section in the summary shows this result. In general, identities shows the number of identical bases between the query and the subject sequence.

The summary also conteins information about the orientation both of the query and the subject sequence, which can be found in "Strand" section. In this case the database genomic sequence progress has the same direction as the query sequence because both of them are plus and they are increasing.

# Question 3.4

Our query sequence is found at "Caenorhabditis elegans chromosome V" and its range is between 6529 and 8028. We can see chromosome name in the figure of question 3.3.

Using http://www.bioinformatics.nl/cgi-bin/emboss/revseq , we reverse the query sequence, we run again the BLAST tool and we get the below summary :

Using http://www.bioinformatics.nl/cgi-bin/emboss/revseq , we reverse the query sequence, we run again the BLAST tool and we get the below summary :

# Caenorhabditis elegans chromosome V

Sequence ID: NC\_003283.11 Length: 20924180 Number of Matches: 1

Range 1: 6529 to 802	8 GenBank G	<u>Graphics</u>	▼ Ne	ext Match 🛕 Previous Match
Score	Expect	Identities	Gaps	Strand
2771 bits(1500)	0.0	1500/1500(100%)	0/1500(0%)	Plus/Minus

It is clear from this summary that the reverse sequence has opposite direction from the subject sequence ( Plus / Minus in the "Strand" section ), while all the other information, such as Identities and Score, is the same.

### Question 3.5

We will find sequences of exons by this code:

```
## gets sequence, starting index and ending index as a parameter
## split and return the sequence between start and index
get_exon = function(seq, start, end){
  org_seq_start = 6529
  fst_index = start - org_seq_start + 1
   return(substr(seq, fst_index, fst_index + (end-start)))
}

## read the query sequence
fasta_file = readLines("732A51_BioinformaticsHT2018_Lab01Ex03.fasta")

## Warning in readLines("732A51_BioinformaticsHT2018_Lab01Ex03.fasta"):
## incomplete final line found on
## '732A51_BioinformaticsHT2018_Lab01Ex03.fasta'
```

```
query_sequence = paste(fasta_file[-1], collapse = "")
## Starting and end indexes of exons.
exon1_start = 6936
exon1_end = 7110
exon2\_start = 7158
exon2 end = 7393
exon3 start = 7433
exon3_end = 7609
exon4 start = 7651
exon4 end = 7818
## get exons with our split function
exon1 = get_exon(query_sequence, exon1_start, exon1_end)
exon2 = get_exon(query_sequence, exon2_start, exon2_end)
exon3 = get_exon(query_sequence, exon3_start, exon3_end)
exon4 = get_exon(query_sequence, exon4_start, exon4_end)
## save exons for protein analysis
writeLines(exon1, file("q3.5_exon1.txt"))
writeLines(exon2, file("q3.5_exon2.txt"))
writeLines(exon3, file("q3.5_exon3.txt"))
writeLines(exon4, file("q3.5_exon4.txt"))
```

#### exon 1

- get exon function output:
- ## TTAAGGAGTTGGGGTGGCTGGAGAAGTTCCTGTAGCCTCCGTGCCGGGAT ...
  - Sequence Text View Tool in blast nucleotide search

### exon 2

- **get\_exon** function output:
- ## CTCAAAATCTCAGTATCCGGAATGCTCAATTTCTGCTTCAAAACCTGTCC ...
  - Sequence Text View Tool in blast nucleotide search

TAAAATTGTAAAATTATCTCAAAATCTCAGTATCCGGAATGCTCAATTTCTGCTTCAAAACCTGTCCGATGCGAAGATTGACATCATCGCGAGTAGCATC
R L I E T D P I S L K Q K L V Q G I R L N V D D R T A D

ACGAGTCCACAAGGAAACCTTGTCACCCTTTTGACGAACATTCACGACAGCTCCGCAGATGTAGTCTCCGTACTCGTCGAATTGCTCTCCAACAATAGCC
R T W L S V K D G K Q R V N V V A G C I Y D G Y E D F Q E G V I A

ATCAACAGCTCCAACCAGTAGTGATCGAGCAATTGCGTTCTTCTCTTCTATGATTCATTGATTAAAAATATATTTCTCAAAAACGTACTTGCTTAT
M L L E L W Y H D L L Q T R R

#### exon 3

- get\_exon function output:
- ## TTGCTTATCGACAACCAACCAACGTCCACCTTGAACGTTGTTGACGT ...
  - Sequence Text View Tool in blast nucleotide search

#### exon 4

- **get\_exon** function output:
- ## CTTCAGACAATCCTCCCATTCCTTGTTACGGTCAGCTTTCAAGTACCAGA ...
  - Sequence Text View Tool in blast nucleotide search

7641 AATCTTCTA CTTCAGACAATCCTCCATTCCTTGTTACGGTCAGCTTTCAAGTACCAGAGAGCCCAGCGATTCTGGAGGGGGGTGTCTGGTGAGAAGCTC

K L C D E W E K N R D A K L Y W L A W R N Q L P H R T L L E

7741 TGGAGGACTGAAGCATCGGACGCATTCACATCGCCGGAAGCTGACAATGCTTTGTTTTCCGCTACGGATGTCTCAT

P P V S A D S A N V D G S A S L A K N E A V S T S M

After getting whole DNA code from all exons, we can search it on blastx to find which protein it is and its protein sequence. We can also see the protein sequence in the screenshots of Sequence Text View Tool in blast nucleotide search for each exons.

• blastx

# Eukaryotic translation initiation factor 4E-3 [Caenorhabditis elegans]

Sequence ID: NP\_503123.1 Length: 251 Number of Matches: 1

► See 2 more title(s)

Range 1	L: 1 to :	236 GenPept	Graphics	▼	Next Match 🛕 Pres	vious Match	
Score		Expect	Method	Identities	Positives	Gaps	Frame
489 bi	ts(126	0) 3e-178	Compositional matrix adjust.	236/236(100%)	236/236(100%)	0/236(0%)	-1
Query	756		KALSASGDVNASDASVPPELLTRHPI KALSASGDVNASDASVPPELLTRHPI			577	
Sbjct	1		KALSASGDVNASDASVPPELLTRHPI			0	
Query	576		SLYNHIQSAGGLNWGSDYYLFKEGIR SLYNHIOSAGGLNWGSDYYLFKEGIR			97	
Sbjct	61		LYNHIQSAGGLNWGSDYYLFKEGI			.20	
Query	396	-	ELLMAIVGEQFDEYGDYICGAVVN\ ELLMAIVGEQFDEYGDYICGAVVN\	-		17	
Sbjct	121		ELLMAIVGEQFDEYGDYICGAVVNV			.80	
Query	216		TEILRYEVHKDSSARTSSTVKPRIC TEILRYEVHKDSSARTSSTVKPRIC				
Sbjct	181	-	TEILRYEVHKDSSARTSSTVKPRIC				

As we can see from output of blastx, all exons are used for producing Eukaryotic translation initiation factor 4E-3 [Caenorhabditis elegans] protein.

After DNA code translation, blastx compared our protein sequence with a negative direction. While our query sequence order is decreasing, out subject sequence order is increasing.

From https://www.wormbase.org/species/c\_elegans/gene/WBGene00002061#0-3gi1-3 and https://www.ncbi.nlm.nih.gov/gene/178536 we can gain the below information:

First of all, the gene's name is ife-3 and it belongs to the gene class ife. It can be found in humans, monkeys, cows and other animals and insects. It is one of the five c. elegans homologs responsible for the protein eIF4E(Eukaryotic translation initiation factor 4E-3). ??ts ancestry also related to Eukaryota, Metazoa, Ecdysozoa and other elegans. Moreover, except for c.elegans, which referred in previous parts of the exercise, BLASTP showed matches with some other species. For instance, the highest match (100%) is with c. brenneri and the lowest (73.7) with S. cerevisiae. Finally, from the gene's ontology perspective, ife - 3 participates in some molecular functions and biological functions such as RNA binding, protein metabolic process and gene expression.

# **Appendix**

```
knitr::opts_chunk$set(fig.width = 7, fig.height = 3, echo = FALSE)
library(dplyr)
library(tidyr)
library(magrittr)
library(kableExtra)
# Question 1.2
# The prob of one allele is the proportion of the homoziqote + half the proportion of the heterozyqote
p = 0.357 + 0.485 / 2
q = 0.158 + 0.485 / 2
# Expected values (Hardy-Weinberg equilibrium)
p_2 = p^2
q_2 = q^2
pq = 2*p*q
Xsq = chisq.test(c(0.357, 0.485, 0.158), p=c(p_2, pq, q_2))
# Xsq$observed
# Xsq$expected
knitr::include_graphics("images/2.2_amino-acids.png")
knitr::include_graphics("images/2.3_nucleotides.png")
knitr::include_graphics("images/2.3_codons-aminoacids")
## gets sequence, starting index and ending index as a parameter
## split and return the sequence between start and index
get_exon = function(seq, start, end){
  org_seq_start = 6529
  fst_index = start - org_seq_start + 1
  return(substr(seq, fst_index, fst_index + (end-start)))
}
```

```
## read the query sequence
fasta_file = readLines("732A51_BioinformaticsHT2018_Lab01Ex03.fasta")
query_sequence = paste(fasta_file[-1], collapse = "")
## Starting and end indexes of exons.
exon1_start = 6936
exon1_end = 7110
exon2\_start = 7158
exon2_end = 7393
exon3_start = 7433
exon3_end = 7609
exon4_start = 7651
exon4_end = 7818
## get exons with our split function
exon1 = get_exon(query_sequence, exon1_start, exon1_end)
exon2 = get_exon(query_sequence, exon2_start, exon2_end)
exon3 = get_exon(query_sequence, exon3_start, exon3_end)
exon4 = get_exon(query_sequence, exon4_start, exon4_end)
## save exons for protein analysis
writeLines(exon1, file("q3.5_exon1.txt"))
writeLines(exon2, file("q3.5_exon2.txt"))
writeLines(exon3, file("q3.5_exon3.txt"))
writeLines(exon4, file("q3.5_exon4.txt"))
cat(substr(exon1,1,50), "...")
cat(substr(exon2,1,50), "...")
cat(substr(exon3,1,50), "...")
cat(substr(exon4,1,50), "...")
```