

2007 7.013 Problem Set 3

Due before 5 PM on FRIDAY, March 16, 2007.

Turn answers in to the box outside of 68-120.

PLEASE WRITE YOUR ANSWERS ON THIS PRINTOUT.

Topic 1.

You have been studying transcription in a type of yeast that is frequently used in wine-making. TATA binding factor (also known as TFIID) in these yeast cells strictly recognizes the sequence 5'-TATAAT-3'. Upon binding, TFIID recruits other components of the transcription apparatus to assemble. Once RNA polymerase has bound to this region, it begins transcribing after the 25th nucleotide downstream of the TATAAT sequence.

Examine the following segments of chromosomal DNA from this species. In each case, write the first 10 nucleotides of the nascent messenger RNA that will be produced. Label the 5' and 3' ends of that molecule.

1a.

5' - GGCTATAATGCTCTACAGCAGACCCGGAAGCCAGAGCAAACGCCAAGG - 3'
 3' - CCGATATTACGAGATGTCGTCTGGGCCTTCGGGTCTCGTTTGCGGTTCC - 5' *

5' GAGCAAACGC 3'

1b.

5' - TAATGCTCTATAATCAGCAGACCTCTATTTCGCTTAGCCAGAACGACGA - 3'
 3' - ATTACGAGATATTAGTCGTCTGGAGATAAGCGAATCGGGTCTTCGTGCT - 5' *

5' AGAAGCACGA 3'

1c.

5' - GGCTATGATGCTCTACAGCAGACCCGGAAGCCAGATTATACCGGATCG - 3' *
 3' - CCGATATTACGAGATGTCGTCTGGGCCTTCGGGTCTAATATGGCCTAGC - 5'

5' CAUUAUAGCC 3'

For each of the above sequences, circle the template strand of the DNA.

* *template*

Having studied this yeast for some time, you've learned that it contains many genes that encode very small proteins, and the genes often contain tiny introns. The intron splice sites in this species always have the composition 5'-GUG-3' (for the 5' splice site) and 5'-UUG-3' (for the 3' splice site), meaning that these sequences and everything between them will be removed from the mature message. Following are three genes from this species. Only the non-template strands are shown. In each case, identify the coding portions and the introns of each gene. Assume that translation always begins at the first methionine codon encountered, and that the introns get spliced out. **In the space beneath each sequence, write out the primary sequence of the polypeptide that would be made.** Use the translation table on the last page of this problem set. Please use the single letter abbreviations for each amino acid.

Gene 1

5' -

GACTATG, ATA, ACT, ATC, GTGCAGCAGACCCGGAAGCCAGAAAACCACTGGATTGTCT, GGC, AGA, GAG, GCC, ACA, TGA
TC-3'

1d. **MITISGREAT****Gene 2**

5' -

ACAATG, TAC, TGC, GCG, ACG, CAC, GGTGACCACTTTCCCAGTTTTCTTTCTTGCA, TCG, TTC, TTA, GAA, GCG, TCG, T
AGAA-3'

1e. **MYCATHASFLEAS****Gene 3**

5' -

TAAATGTAT, ACC, CAT, ATC, AGC, ATT, TCG, GTG, GAA, CGA, TAT, CAC, GCT, AGA, GAT, TAA, TTGGAA, GCT, AGT,
TAC, TGACCCA-3'

1f. **MYTHISISEASY**

You've isolated several mutants from this species and found they each have subtle changes in the DNA sequences of the three genes listed above. Indicate what kind of mutation was involved in each case (Your choices are "missense," "nonsense," or "silent.")

- 1g. Nucleotide 10 in gene 1 was changed from "A" to "G": missense
- 1h. Nucleotide 16 in Gene 1 was changed from "C" to "A": silent
- 1i. Nucleotide 9 of Gene 2 was changed from "C" to "G": nonsense
- 1j. Nucleotide 3 in Gene 3 was changed from "A" to "G": silent

1k. Examining the rest of the sequence of the mutated Gene 3 reveals that two more changes lie further downstream within this same gene, and that these mutations are not even in the exons. One of these converts the sequence 5'-GGAAC-3' to 5'-CGAAC-3'. At the same time, the other mutation converts 5'-TTAATT-3' to 5'-TTAAAT-3'. Do you expect these changes to change the translation product. If not, *briefly* explain why not in the space below (10 words or less). If it does change the translation product, then show the modified product in the space below.

Intron is not spliced out, encodes AAs and stop codon:

MYTHISISVERYHARD

Topic 2

Towards the end of this problem set is the sequence of the alpha-tubulin gene from *Zea mays*. The sequence of the mature messenger RNA is also presented. Examine the sequence to answer the following questions.

2a. Where in the DNA sequence would you expect to find promoter sequences, including a TATA box? (Your choices are: "to the 5' side of the transcription start site," "to the 3' side of the transcription start site," or "within the first exon.")

to the 5' side of the transcription start site

2b. How many introns are there in this gene? 4

2c. How long is the 5' UTR? 76

2d. How long is the 3' UTR? 164 (including stop codon)

2e. How long is the mRNA (excluding the polyA tail and the cap)? **1590 bases**

2f. How long was the pre-mRNA primary transcript (assuming no processing occurred until after transcription had been completed; assume also that the primary transcript ended where the current polyA tail begins)?

3218 bases

2g. How many amino acids make up the protein that will be translated directly from this transcript (before any post-translational modifications)?

450 AAs

Topic 3

Having sequenced several alpha-tubulin genes in your laboratory, you notice that the alpha-tubulin gene from *Zea mays* differs from most other alpha-tubulins in one specific portion of the protein. The portion that differs most strongly is encoded by exon 2. You decide that to test the importance of these differences in the function of alpha-tubulin, you will transfer exon 2 from *Zea mays* into the alpha-tubulin gene from another species. The first step in doing this will require

Name: _____

4

you to obtain a copy of exon 2 by PCR. Using the sequence information of the alpha-tubulin gene on subsequent pages, describe two primers, each 10 bases in length, that could be used to precisely amplify exon 2 of alpha-tubulin. Write out the sequences of those primers in the space below. Be sure to label their 5' and 3' ends.

3a. Primer 1:

5' TGATTCCTCG 3'

3b. Primer 2:

5' CGTGTAGTGG 3'

Topic 4

You have discovered a new plasmid in species of foul-smelling, soil dwelling bacteria. As a first step toward understanding the composition of this plasmid, you digest the plasmid with a series of restriction enzymes, either alone or in combination. Using agarose gel electrophoresis, you determine the size of each fragment produced in these reactions and list the results in the Table below:

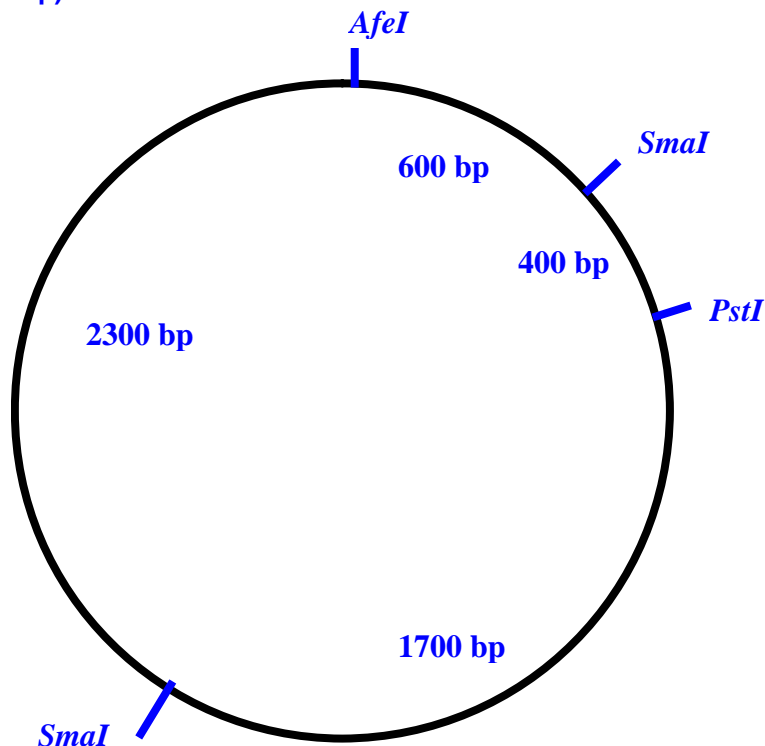
Enzyme(s) used	<i>AfeI</i>	<i>PstI</i>	<i>SmaI</i>	<i>AfeI</i> + <i>PstI</i>	<i>AfeI</i> + <i>SmaI</i>	<i>PstI</i> + <i>SmaI</i>	<i>AfeI</i> + <i>PstI</i> + <i>SmaI</i>
DNA fragments produced	5000 bp	5000 bp	2900 bp 2100 bp	4000 bp 1000 bp	2300 bp 2100 bp 600 bp	2900 bp 1700 bp 400 bp	2300 bp 1700 bp 600 bp 400 bp

4a. How big is this plasmid (in basepairs)?

5 kbp or 5000 bp

4b. Using the circle below, draw a restriction map of the plasmid. Be sure to indicate the distances between adjacent restriction sites (in basepairs).

***AfeI* (0 bp)
SmaI (600, 2700bp)
PstI (1000 bp)**



Topic 5

Gaucher's disease is genetic disorder with a higher incidence among the Ashkenazi Jewish population. Affected individuals cannot metabolize certain lipids, primarily because of a deficiency in the enzyme glucocerebrosidase (GCA), causing glucocerebrosides to accumulate in the blood and other tissues. The deficiency is caused by a mutation in the gene encoding GCA.

In an effort to develop a gene therapy for this disorder, you have set out upon an ambitious strategy:

- you intend to produce a copy of the GCA gene that is specifically expressed in liver cells;
- You will introduce this gene into human liver cells by using the specialized, non-disease causing virus called HEP-RC, which can infect liver cells;
- you will then transplant the resulting cells into the liver of a patient, hopefully establishing a cluster of cells in the liver that allows the patient to metabolize glucocerebrosides.

5a. You have a cloned copy of the wild-type (functional) GCA coding sequence. As a first task, select a promoter that, when joined to the GCA gene, is most likely to ensure proper expression of the GCA gene in liver cells:

_____ The promoter from the gene that encodes β -globin, an important blood-specific protein

 X The promoter from the gene that encodes the CK-19 protein, which is normally transcribed only in the liver

_____ The promoter from the gene that encodes GFAP, a brain-specific protein

_____ The promoter from the gene that encodes α -tubulin, which is expressed in almost all cells

5b. In the space below, justify your choice in 15 words or less.

The promoter will interact with transcriptional activation pathways in the host cell- a liver cell.

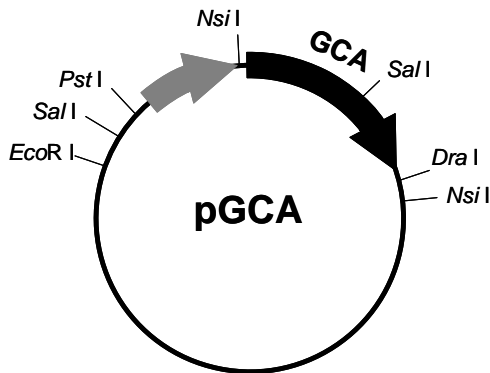
Having selected the best promoter, your assistant creates a plasmid in which the promoter and the GCA are fused together. This plasmid is called pGCA. You also have the HEP-RC genomic DNA (see below). You now need to excise the promoter-GCA region from pGCA, and you need to cut the HEP-RC DNA with restriction enzymes in a manner that allows you to ligate them together, to create a recombinant virus for use in your gene therapy strategy.

5c. Based on these diagrams, which two restriction enzyme(s) should you use to excise the promoter-GCA region from pGCA so that it can later be ligated into the viral genome. (Think about how you'll be cutting the HEP-RC DNA before you finalize your decision.)

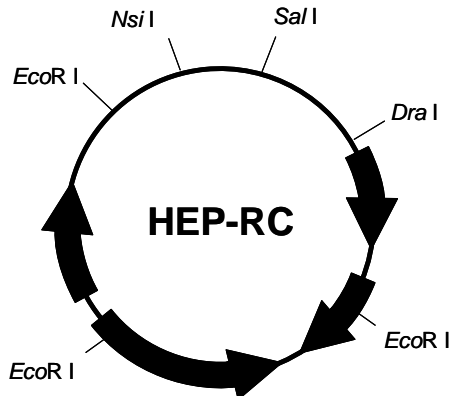
Pst I, Dra I

5d. Which restriction enzymes would be suitable for preparing the HEP-RC genome to accept the resulting promoter-GCA fragment?

Nsi I, Dra I



pGCA contains the promoter you have selected (grey arrow) and the coding sequence from the GCA gene, (heavy black arrow). Also indicated are the relative positions of several restriction sites.



The HEP-RC genome is a circular double stranded DNA molecule and contains several genes (heavy arrows) that must not be disrupted if the viral DNA is to persist in liver cells.

Restriction Enzyme Recognition Sequences
Arrowheads indicate where the enzyme cleaves the phosphodiester bonds between DNA bases

DraI
5'... TTT[▼]AAA... 3'
3'... AAA[▲]TTT... 5'

EcoRI
5'... GAATTC[▼]... 3'
3'... CTTAAG[▲]... 5'

NsiI
5'... ATGCAT[▼]... 3'
3'... TACGTA[▲]... 5'

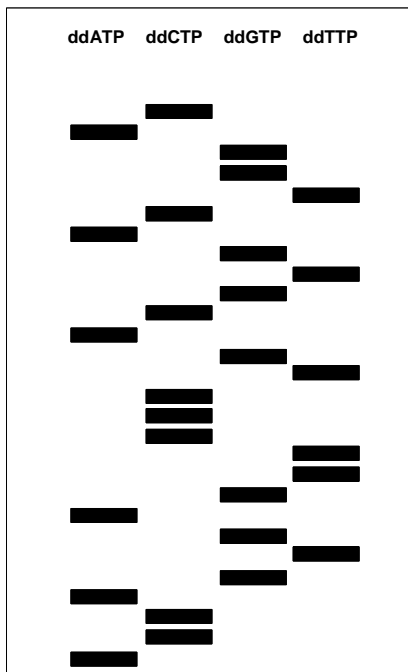
PstI
5'... CTGCAG[▼]... 3'
3'... GACGTC[▲]... 5'

SalI
5'... GTCGAC[▼]... 3'
3'... CAGCTG[▲]... 5'

Topic 6

6a. In your laboratory, you have discovered a gene that seems to be involved in cancer progression. As a first step toward understanding its function, you clone the gene and sequence it. Below is an image of the sequencing gel that you prepared using the Sanger dideoxy nucleotide sequencing technique. From this image determine the sequence of the DNA that was examined. Label the 5' and 3' ends of that DNA sequence. Remember that smaller DNA molecules migrate more quickly through a gel and will reach the bottom of the gel sooner than larger molecules. At the top of each lane is indicated the type of dideoxynucleotide that had been included in each polymerization reaction.

5' ACCAGTGAGTTCCTGACGTGACTGGAC 3'



(You do not need to turn in this page)

One strand of the DNA sequence from the alpha-tubulin locus is shown below in **BOLD**. Where appropriate, the sequence of the mature mRNA is shown directly beneath the corresponding DNA sequence (NOT BOLD). Relative numbering of each is indicated to the left. "@" indicates the position of the 7-methyl-G cap. Exclamation points (!!!) are positioned beneath the nucleotides that constitute the start codon; Asterisks (***) are positioned beneath the nucleotides that constitute the stop codon.

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1  TACGGACGGG AGAAGCGCGC GGGCGGGCCG GCCGGCCTCA CGTGAACAAC GCCCGCCCAC
61 GAAACCGCGT GGCTGTGCAA AAGCAACCGT TTCCAGTTTC CTCCCCAGTG CACCCCCTCC
121 CCTCTCCCCT CCCTCTCTCT CCTCCGGCCC CCTTTTTTTA TGCCCCGATCT CACACATTTT
181 GGAAGGAGAG AGAGAGAGGC AGAGGGAGAG ATTGGAGGGA GGCCCCTGCC GTAGGCAAGA
1                                     @CCCUGCC GUAGGCAAGA

241 GAAACCGCGG CGCGCGGAGA GGGAGGAAGG GCAGACGCAG ACGCAGACGG GCGAACAAGA
18 GAAACCGCGG CGCGCGGAGA GGGAGGAAGG GCAGACGCAG ACGCAGACGG GCGAACAAGA
!

301 TGAGGGAGAT CATCAGCATC CACATCGGCC AGGCCGGGAT CCAGGTCCGG AACGCCTGCT
78 UGAGGGAGAU CAUCAGCAUC CACAUCGGCC AGGCCGGGAU CCAGGUCCGG AACGCCUGCU
!!

361 GGGAGCTCTA CTGCCTCGAG CACGGCATCG AGCACGATGG CACCATGCCC AGGTGAGTGC
138 GGGAGCUCUA CUGCCUCGAG CACGGCAUCG AGCACGAUGG CACCAUGCCC AG

421 TGCATGCCCT GTCGATCCGC CGGGACGCGC TCCTCCTGTT TACCCATTTT TTGCGACTTA
481 AAGCTGTTGC GCGTTCTATG GAGCCGGCGC ATCAAAATTC TGGCGTGCTC ATTCGTCGCA
541 TTTCTAGAT TCGCTGCGTT GTTTTCTTCC AATTCGTCGG AGCAAATGAC GGGGCGTGTA
601 GGTGTAGTGA TGGTACTTCC ATATCCGTTT CGTATTTTTT GGTTTCGTCG TCCCTTTCCG
661 TTGCACTCGC AGTTTTGGTT CCTTTGCGAC TGGATTTGTT TGTTGGGTGG GATGTGTTGT
721 TGTTGTTGTC TGTTTGGGTT TTGGTGCATT TTGCCGGTTG TTCGAGAGAT TGCTATGAGC
781 CGGGACCAGA TGCCACGTTT CCTAATTAAG AGTTTGCCT AGCGTTTTTT TTTTGGACTA
841 TTAATACTAG GCCTGTAAAT TAGGCAGGCG CCACAGAAGG ATGCGTTGAG CTCTGAAGAA
901 TTGAATCTGC AGGGATGTCA TCTAAATAAA TAATATAACT GGGATGTTCT GATGAGTGTG
961 AGACCATGGA TCTCCCTGGA TTCACGAGTT TCATCTGACC CTTTGTAATA AGTAGCACCC
1021 TTCTCTCTGT TTGCCTGATC TTGTAGTACC ATAGAAAATT GCCCTAATGC TGCTCCGCCCT
1081 GATAATAGAA AAAATCCTGT TTTAGGACCC CAAGGAGGCA GGTTCGTTAG GCTATAAATT
1141 CTCTTTAAGC GCATATGCCA ATGGTTCGTG CCTTATTGTT ATTGGAGTTC CATGCATATC
1201 ATTTTTTATA ACACTGGTTT GATTGTGATG CAGTGATTCC TCGGTTGGCG TCGCACAATGA
190                                     UGAUUC UCGGUUGGCG UCGCACAUGA

1261 TGCCTTCAAC ACGTTCTTCA GCGAGACTGG TTCCGGCAAG CATGTGCCCA GGGCCATCTT
217 UGCCUUAAC ACGUUCUUA GCGAGACUGG UUCGCGCAAG CAUGUGCCCA GGGCAUCUU

1321 CGTCGACCTT GAGCCCACTG TCATCGACGA GGTTCGCACT GGCTCGTACC GCCAGCTCTT
277 CGUCGACCUU GAGCCACUG UCAUCGACGA GGUUCGCACU GGCUCGUACC GCCAGUCUU

1381 CCACCCAGAG CAGCTCATCT CGGGAAGGA GGATGCAGCT AACAACTTTG CCCGTGGCCA
337 CCACCCAGAG CAGCUCAUCU CGGGAAGGA GGAUGCAGCU AACAAUUUG CCCGUGGCCA

1441 CTACACGGTA ATTCACCTTG TGTCTGAATA GATAATAGGA GTGGGTCCTA TGCTTCTGCA
397 CUACACG

1501 ACTGTGAAA TGTTC AATTC TAATTATGGT GCAATGCAGT TGGGAAGGAG ATTGTCGATC
404                                     GU UGGGAAGGAG AUUGUCGAUC

1561 TATGCCTGGA CCGTGTGCGC AAGCTTGCAG ACAACTGCAC TGGGCTGCAG GGATTCTTGG
426 UAUGCCUGGA CCGUGUGCGC AAGCUUGCAG ACAACUGCAC UGGGCUGCAG GGAUUCUUGG

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1621 TGTTC AATGC TGTGG TGGT GGTACTGGCT CTGGACTTGG TTCACTGCTG TTGGAGCGCC
 486 UGUUCA AUGC UGUUGGUGGU GGUACUGGCU CUGGACUUGG UUCACUGCUG UUGGAGCGCC

1681 TTTCAGTTGA TTATGGCAAG AAGTCTAAGC TTGGTTTCAC CATTATCCT TCCCCACAGG
 546 UUUCA GUUGA UUAUGGCAAG AAGUCUAAGC UUGGUUUCAC CAUUUAUCCU UCCCCACAG

1741 TATGACCTTA TGACAGAACC TTACATTAGC AACCTAGTTT CTGTGGATTT CACACAAAAA
 1801 TATATTCATA TAATTATTCT TTCCAAAAGG AGTTTGGCTT AAGTATATGA GAATTGTAAA
 1861 ACATACATGA AGTATTAGGT TGACAATTTA TATTTATACA CCAACATGAA ATTTTTTTTAT
 1921 ATGTAAATTA ATCTACGGAT TATGCAATTT GGTGGCACTG TCCTTGCCCT CAAAATTACA
 1981 GACAGAGCAC TGATTATTGG ATCAGCCTAG CATTATATAA ATCATGTGTG GAACAAAACA
 2041 AGCAACTGTT GTGAACTTAT GGTAGTGGAG ACCTTCTTGT ATATTGTTTA GACCATGGTT
 2101 AGTACAGCTG TCACAGTCTT CAGGTCTTCT TTATGAGTGT CTTTAAATCA TGCATTGCAT
 2161 ATGTGCCAGC ATTATCCTAT AGTGCTGCTC CTTCTGTCC ATGTGATGTG GTCCATTTCAT
 2221 CTTGTTTCATG TGAGTAGGTT TTCTGCATCT TAGCTGTTGT GGGGAAAAGT GGGTAGAGAT
 2281 GTTCGATGAA TTTGGATCTG ATCCATCATT TAACTTTTCA TGTAACGCCA CTCTGGAATG
 2341 CAGGTGTCAA CAGCTGTTGT AGAGCCATAC AACAGTGTCC TCTCGACCCA CTCCTTGCTC
 605 GUGUCA CAGCUGUUGU AGAGCCAUAC AACAGUGUCC UCUCGACCCA CUCCUUGCUC

2401 GAGCACACTG ATGTTGCAGT CCTCCTGGAC AACGAGGCTA TCTATGACAT ATGCAGGAGG
 662 GAGCACACUG AUGUUGCAGU CCUCCUGGAC AACGAGGCUA UCUAUGACAU AUGCAGGAGG

2461 TCCCTTGACA TCGAAAGGCC AACCTATAACC AACTTGAACA GGCTGATCTC ACAGATCATA
 722 UCCCUUGACA UCGAAAGGCC AACCUAUACC AACUUGAACA GGCUGAUCUC ACAGAUCAUA

2521 TCATCACTTA CCACCTCCCT GAGGTTTGTAT GGGGCTATCA ATGTGGATGT TACTGAGTTC
 782 UCAUCACUUA CCACCUCCU GAGGUUUGAU GGGGCUAUCA AUGUGGAUGU UACUGAGUUC

2581 CAGACCAACC TTGTTCCATA CCCACGTATT CATTTCATGC TTTCTCATA TGCCCCCTGTA
 842 CAGACCAACC UUGUUCCAUA CCCACGUAAU CAUUUCAUGC UUUCCUCAUA UGCCCCUGUA

2641 ATATCTGCTG AGAAGGCCTA CCATGAGCAG CTCTCTGTGC CTGAAATCAC CAATGCTGTA
 902 AUAUCUGCUG AGAAGGCCUA CCAUGAGCAG CUCUCUGUGC CUGAAAUCAC CAAUGCUGUA

2701 TTTGAGCCCT CAAGCATGAT GGCCAAGTGT GACCCAAGGC ACGGAAAGTA CATGGCTTGC
 962 UUUGAGCCCU CAAGCAUGAU GGCCAAGUGU GACCCAAGGC ACGGAAAGUA CAUGGCUUGC

2761 TGCTTGATGT ACCGCGGTGA TGTGTTTCTT AAGGATGTCA ACGCTGCAGT TGCAACCATC
 1022 UGCUUGAUGU ACCGCGGUGA UGUUGUUCU AAGGAUGUCA ACGCUGCAGU UGCAACCAUC

2821 AAGACCAAGA GAACTGTCCA GTTTGTGGAC TGGTAAGATG TCTTTGTGCG ATTTGAGTTG
 1082 AAGACCAAGA GAACUGUCCA GUUUGUGGAC UG

2881 TGCTTATTCT ATGATATCAC ATGCTTGTCA CTTGATTGCG GATGGTAAGT GGGTAACTAA
 2941 TTCAATTATG CTGTCGCCAC ACAGGTGCCC CACTGGATTG AAGTGTGGCA TCAACTATCA
 1114 GUGCCC CACUGGAUUC AAGUGUGGCA UCAACUAUCA

3001 GCCACCCTCT GTTGTCCCTG GAGGTGACCT GGCAAAGGTC CAGCGTGCCG TGTGCATGAT
 1150 GCCACCUCU GUUGUCCUG GAGGUGACCU GGCAAAGGUC CAGCGUGCCG UGUGCAUGAU

3061 CAGCAACAAC ACCGCCGTTG CCGAGGTGTT TTCACGCATC GACCACAAGT TTGACCTTAT
 1210 CAGCAACAAC ACCGCCGUUG CCGAGGUGUU UUCACGCAUC GACCACAAGU UUGACCUAU

3121 GTACGCCAAG CGCGCATTG TTTACTGGTA TGTCGGTGAG GGTATGGAAG AGGGTGAGTT
 1270 GUACGCCAAG CGCGCAUUCG UUCACUGGUA UGUCGGUGAG GGUUAUGGAAG AGGGUGAGUU

3181 CTCAGAAGCC CGTGAGGACT TGGCTGCTCT TGAGAAGGAC TATGAGGAAG TCGGTGCAGA
 1330 CUCAGAAGCC CGUGAGGACU UGGCUGCUCU UGAGAAGGAC UAUGAGGAAG UCGGUGCAGA

3241 GGGTGCCGAT GACGAGGGTG ACGAGGGAGA CGACTATTGA ATAGCTGGCT AATAAGTAGT

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1390 GGGUGCCGAU GACGAGGGUG ACGAGGGAGA CGACUAUUGA AUAGCUGGCU AAUAAGUAGU

3301 **TCTCTGGTGG TTAATGGTTG GACTATTTTG GAGCTATATA CTCTATGGCT CCACTCCATT**
1450 UCUCUGGUGG UAAAUGGUUG GACUAUUUUG GAGCUAUUAU CUCUAUGGCU CCACUCCAUI

3361 **GGATACTGCT GCTGGTTGTG TGTTCATT TGTACTATG TAGTAAATTG TTCGTAACCC**
1510 GGAUACUGCU GCUGGUUGUG UGUUUCCAUI UUGUACUAUG UAGUAAAUUG UUCGUAACCC

3421 **CCCTATTGGC CATGATTGTT CCATATCATC CTTCTTTGCG TTTGCAACGC TATTCGTCCA**
1570 CCCUAUUGGC CAUGAUUGUU CAAAAAAAAA AAAAAAAAAA AAAAAAAAAA AAAAAAAAAA

3481 **ATTTCCGGTG TATATGCTAT AATGCTATTA TGTGCAAGT GTGCTGTGCT CAACCATTGG**
3541 **GTTTTATTTA TATTTAGAAA CAGAAGGCAT TATTACTCTT CGCTGAAGCT TGTTTCTGAA**
3601 **TCAGGGGAAC ATTGAGATT TGAAGATTT TGACAGCAGT CTACCACTGT GAGGGCCTAA**
3661 **GGATCCCACG GGAG**

Codon Table

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	U	C	A	G
U	UUU Phe (F)	UCU Ser (S)	UAU Tyr (Y)	UGU Cys (C)
	UUC “	UCC “	UAC “	UGC “
	UUA Leu (L)	UCA “	UAA <i>Stop</i>	UGA <i>Stop</i>
	UUG “	UCG “	UAG <i>Stop</i>	UGG Trp (W)
C	CUU Leu (L)	CCU Pro (P)	CAU His (H)	CGU Arg (R)
	CUC “	CCC “	CAC “	CGC “
	CUA “	CCA “	CAA Gln (Q)	CGA “
	CUG “	CCG “	CAG “	CGG “
A	AUU Ile (I)	ACU Thr (T)	AAU Asn (N)	AGU Ser (S)
	AUC “	ACC “	AAC “	AGC “
	AUA “	ACA “	AAA Lys (K)	AGA Arg (R)
	AUG Met (M)	ACG “	AAG “	AGG “
G	GUU Val (V)	GCU Ala (A)	GAU Asp (D)	GGU Gly (G)
	GUC “	GCC “	GAC “	GGC “
	GUA “	GCA “	GAA Glu (E)	GGA “
	GUG “	GCG “	GAG “	GGG “

The Genetic Code. Three-letter and one-letter abbreviations for each amino acid are listed alongside their corresponding codons. Ditto marks (“) indicate the same amino acid as above.