7.014 Quiz II 3/18/05

Your Name:		TA's Name:		
Write your nan	ne on this pag	e and your initials on all the other pages in the space provided		
		ding this coversheet. Check that you have pages 1-10. Genetic e amino acids can be found on page 10.		
This exam has	4 questions.	Read all questions before starting to write.		
Write your ans	wers as clearly	y and precisely as possible in the space provided.		
This is a <u>closed</u>	<u>l book</u> exam.			
Question	Value	Score		
1	29			
2	30			
3	25			
4	16			

TOTAL:

100

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Question 1 (29 points)

Barney is an alien. On his ship, hidden in the Stata building, you find alien bacteria that metabolize wood. You call this species *A. termiticus*, and call your original strain BLT (for "Barney's little termiticus").

You subject a sample of BLT to mutagens, and isolate a new strain that no longer metabolizes wood. You conclude that you have succeeded in disrupting at least one gene necessary to metabolize wood. You call the mutant strain M.

You mix a sample of M with a sample of heat-killed wild type BLT, and the resulting strain metabolizes wood. You summarize your data in the following table:

Strains	Metabolizes wood?
BLT	Yes
Heat-killed BLT	No
M	No
Heat-killed BLT +M	Yes

a) Did the content of any of the BLT or M cells change in the experiment? If yes, which cells underwent the change, and what change occured? If not, explain why there was no change.

You plan to characterize the alien genetic material. You start by breaking some *A. termiticus* cells open to determine their molecular composition. You find that they contain various small molecules, carbohydrates, lipids, and two other macromolecules, A and B.

In order to determine which macromolecule is the carrier of genetic information, you repeat your previous experiment, but include test tubes where you treat the sample of the heat-killed BLT with either an agent that destroys macromolecule A (A-ase) or macromolecule B (B-ase). You find the following results (including the repeat of your previous experiment in the first 4 lines):

Strains and agents	Metabolizes wood?
BLT	Yes
Heat-killed BLT	No
M	No
Heat-killed BLT +M	Yes
A-ase treated heat-killed BLT +M	Yes
B-ase treated heat-killed BLT +M	No

b) Which molecule is the carrier of genetic information in *A. termiticus*? Justify your answer.

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Question 1, continued

Next, you set out to determine the structure of the alien genetic material molecule. You first determine that it has six types of bases that you name S, V, W, X, Y, Z. You further determine that the alien cell's content of S is the same as its content of each of X and Z; and that the content of V is the same as its content of each of W and Y.

When you determine the structure of this molecule by X-ray crystallography, you are not surprised to find that the molecule consists of 3 interacting strands.

c) What base interaction combinations do you expect for this molecule?

You want to investigate the mechanism of replication of the alien genetic material. You decide to repeat the Meselson-Stahl experiment. Recall that labeled strands are "heavy" (low in the gradient) and unlabeled strands are "light" (high in the gradient). Also recall that in the experiment, the culture is grown on heavy Nitrogen (¹⁵N), and is switched to light Nitrogen (¹⁴N) at time=0.

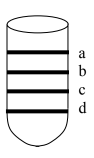
Before proceeding you define three possible models of alien replication:

- <u>conservative</u>, in which, after replication, old strands stay together, and new molecules are made entirely of new strands;
- <u>semi-conservative</u>, in which, after replication, each molecule has one old strand; and
- <u>asymmetric</u>, in which replication creates a molecule with one new strand, and a molecule with two new strands.

Each column in the table below reflects the predictions one of these models makes about the outcome of the experiment. The outcomes are described using symbols a-d to indicate the levels on the gradient as depicted in the figure on the right.

d) Label each column with the name of the appropriate model.

# cycles of replication			
0	d	d	d
1	b, с	a, d	b
2	a, b, c	a, d	a, b
3	a, b, c	a, d	a, b



You determine that each *A. termiticus* mother cell completes one round of genetic material replication, but gives rise to three daughter cells.

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e) Which of the above models is <u>not consistent</u> with this data? Why?

Question 1, continued

Below are the structures of an alien nucleotide X and the earth DNA nucleotide cytosine (C).

- f) For both structures,
 - i. box the sugar
 - ii. circle the base
 - iii. underline the part of the structure used to power the addition of the nucleotide onto the growing chain.
- g) For the DNA nucleotide,
 - i. label the 5' end
 - ii. label the 3' end
 - iii. draw an arrow to the part of the molecule that identifies it as a nucleotide used in DNA rather than in RNA.

Naı	me:			TA:
Qι	aestion 2 (30 poin	nts)		
a)	Transcription			
	i. is the process	that transfers informatio	n from	_ to
	ii. in eukaryotic o	organisms, transcription	occurs in the	
	Nucleus	Ribosome	Membrane	
b)	Translation			
	i. is the process	that transfers informatio	n from	_ to
	ii. in eukaryotic o	organisms, translation oc	ccurs in the	
	Nucleus	Ribosome	Membrane	
hyj bol 5' 3'	pothetical bacteria <i>E</i> ld. The underlined T -TT C CCCTATGG <i>E</i> -AA G GGGATACCT	e of DNA encodes a hyp. hypotheticus. Transcripe I/A base pair indicates the ATGGTCATCTACGATGTACCAGTAGATGCTAC	tion starts at and inches terminator. CCCCCATCACTA CGGGGGTAGTGAT	cludes the C/G base pair in AAGCT <u>T</u> G-3' TTCGA <u>A</u> C-5'
d)	What are the first 3 C- termini.	amino acids of the subse	equent polypeptide?	? Be sure to label the N- and
e)	How many total am	nino acids are encoded ir	this polypeptide?	
	2	bacteria containing a mien it recognizes the code		•
f)	What is the sequence	ce of the anticodon of the	e mutant tRNA? Be	sure to label the 5' and 3'

ends.

Na	me: TA:
Qı	uestion 2, continued
	<u>longer</u>
g)	The Playdo polypeptide would be <u>the same length</u> in the presence of the mutant tRNA.
	<u>shorter</u>
	Why?
pa: a C	bu isolate a mutant bacteria with the Playdo gene sequence below. The bold, boxed G/C base ir is the site of the only difference between wild-type and mutant Playdo—a substitution of G/C base pair for a A/T base pair. As before, the bold C/G base pair indicates the start of inscription, and the underlined T/A base pair indicates the terminator.
5′ 3′	-TT C CCCTATGG G TGGTCATCTACGATGCCCCCATCACTAAAGCT <u>T</u> G-3' -AA G GGGATACC C ACCAGTAGATGCTACGGGGGTAGTGATTTCGA <u>A</u> C-5'
h)	What is the effect of this substitution on the peptide?
i)	Do you expect this peptide to have the same function as the wild-type bacterial peptide? Why or why not?
sec	trigued by Playdo, you search for a similar protein in mice by looking for similar DNA quences in the mouse genome. You find a gene that matches bacterial Playdo almost rfectly but contains a 36 DNA base pair insertion in the center of it.
	hen you purify the Playdo polypetide from mouse cells you are shocked to learn that buse Playdo is the same length in amino acids as bacterial Playdo.
j)	Explain how is it possible that mouse Playdo and bacterial Playdo are the same polypeptide length even though they have substantially different gene lengths.
k)	Do you expect bacterial and mouse Playdo to have the same promoter and terminator sequences? Why or why not?

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Question 3 (25 points)

- a) Which of the following could alter gene regulation (circle all that apply)
 - i. Deleting a promoter
 - ii. Moving a yeast culture to a new food source
 - iii. Raising the temperature of a bacterial culture
 - iv. Mutating a repressor gene, such that the resulting protein no longer functions

The B operon contains the genes involved in the breakdown of sugar B in the bacteria *E. fake*. Sugar A is the preferred sugar in *E. fake*. In the absence of sugar A, *E. fake* can also use sugar B. The B operon is subject to both negative and positive regulation.

b) Indicate with a Low or a High the expected level of B operon expression when *E. fake* cells are grown in the presence (+) or absence (-) of the sugars A and/or B.

B operon expression	Sugar A only	Sugar B only
	-	-
	-	+
	+	-
	+	+

Below is the diagram of the B operon. B-R encodes B Repressor, the repressor of the B-XYZ genes. Ter sequences denote transcription terminators. P sequences denote promoters. O denotes an operator, and Enh—an enhancer. Recall that the B operon is subject to both negative and positive regulation.

P_R	B-R	TerR	Enh Pxyz O	B-X	B-Y	B-Z	Terxyz

- c) How many in frame translation stop signals (stop codons) are present in the mRNA transcript originating with P_{XYZ} ?
- d) To which element does the B Repressor protein bind?
- e) What is the effect of the B Repressor binding on
 - i. transcription of B-XYZ?

decrease no change increase

ii. translation of any B-XYZ transcripts already made?

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decrease	no change	increase		
uecrease	no change	nicrease		

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Question 3, continued

You have isolated three <u>loss of function</u> mutants in the B operon.

f) For each mutant in the table below, for each condition, indicate (with a Yes or a No) whether the repressor protein and the activator protein are each bound to their respective binding sites, and what the resulting level of expression (None, Low, or High) of the B operon is. Data for the wild-type strain is filled in for you.

Strain	Mutation	Sugar A only			Sugar B only			Sugars A and B		
in		Repressor	Activator	Expression	Repressor	Activator	Expression	Repressor	Activator	Expression
WT		Yes	No	None	No	Yes	High	No	No	Low
M1	B-R									
M2	Enh									
M3	P _{XYZ}									

Question 4 (16 points)

You hope to understand the lysine biosynthesis pathway, so you decide to look for mutants that can not survive without supplementation with the amino acid lysine.

You mutagenize some bacteria, and plate them on rich media. You then replica plate from your original plate onto three plates: one containing complete media, one containing minimal media, and one containing minimal media plus lysine.

By comparing the minimal and the +lysine plates, you identify six colonies that are lysine auxotrophs, that is, they require lysine from the media to grow.

Below are the results of the complementation test, where + means growth and – means no growth.

Mutant	lys1	lys2	lys3	lys4	lys5	lys6	wild-type
lys1	_	+	+	_	+	-	+
lys2		-	-	+	+	-	+
lys3			_	+	+	-	+
lys4				-	+	ı	+
lys5					-	-	+
lys6						-	_
wild-type							+

a) For each mutant, circle whether the mutation is dominant or recessive.

lys1	dominant	recessive
lys2	dominant	recessive
lys3	dominant	recessive
lys4	dominant	recessive

Question 4, continued

- b) Place the recessive mutants into complementation groups.
- c) How many genes (at least) must there be in the lysine biosynthesis pathway?

You find some lysine precursors (X, Y, and Z) that, when added to the media, allow the growth of some mutants. You try growing several mutants on minimal media with X, Y, or Z added, and get the following results. + means growth and - means no growth.

	Precursor supplement						
Mutant	X	Y	Z				
lys1	-	+	-				
lys2	+	+	-				
lys5	-	-	-				
wild-type	+	+	+				

d) Circle the pathway(s) consistent with data.

$$Y \xrightarrow{lys2} X \xrightarrow{lys1} Z \xrightarrow{lys5} lysine$$

$$Y \xrightarrow{lys5} X \xrightarrow{lys1} Z \xrightarrow{lys2} lysine$$

$$X \xrightarrow{lys1} Z \xrightarrow{lys5} Y \xrightarrow{lys2} lysine$$

$$Z \xrightarrow{lys2} X \xrightarrow{lys1} Y \xrightarrow{lys5} lysine$$

$$Y \xrightarrow{lys5} Z \xrightarrow{lys2} X \xrightarrow{lys1} lysine$$

$$X \xrightarrow{lys2} Y \xrightarrow{lys1} Z \xrightarrow{lys5} lysine$$

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$$Z \xrightarrow{lys5} Y \xrightarrow{lys1} X \xrightarrow{lys2} lysine$$

The Genetic Code

	The Genetic Code										
		U		С		A		G			
U	UUU	phe	UCU	ser	UAU	tyr	UGU	cys	U		
	UUC	phe	UCC	ser	UAC	tyr	UGC	cys	C		
	UUA	leu	UCA	ser	UAA	STOP	UGA	STOP	Α		
	UUG	leu	UCG	ser	UAG	STOP	UGG	trp	G		
C	CUU	leu	CCU	pro	CAU	his	CGU	arg	U		
	CUC	leu	CCC	pro	CAC	his	CGC	arg	C		
	CUA	leu	CCA	pro	CAA	gln	CGA	arg	Α		
	CUG	leu	CCG	pro	CAG	gln	CGG	arg	G		
A	AUU	ile	ACU	thr	AAU	asn	AGU	ser	U		
	AUC	ile	ACC	thr	AAC	asn	AGC	ser	C		
	AUA	ile	ACA	thr	AAA	lys	AGA	arg	Α		
	AUG	met	ACG	thr	AAG	lys	AGG	arg	G		
G	GUU	val	GCU	ala	GAU	asp	GGU	gly	U		
	GUC	val	GCC	ala	GAC	asp	GGC	gly	C		
	GUA	val	GCA	ala	GAA		GGA	gly	Α		
	GUG	val	GCG	ala	GAG	glu	GGG	gly	G		

STRUCTURES OF AMINO ACIDS at pH 7.0

