## Protein Modeling C - Protein Modeling - UT C - 03-27-2021

Per Texas Science Olympiad rules, you must have printed notes for this event. If you are communicating with your partner through a voice or video call, please start it before you begin the test itself.

Significant time spent outside of the browser window is grounds for a penalty or disqualification per TSO policies.

● True ○ False

1. (1.00 pts) Which of the following isotopes could be used to radioactively label DNA? Select all that apply.
(Mark ALL correct angulars)
(Mark ALL correct answers)  ✓ A) 14-C
☑ B) 32-P
□ C) 12-C
□ D) 32-S
2. (1.00 pts) Select all of the following amino acids with hydrophobic side chains.
(Mark ALL correct answers)
✓ A) Valine
B) Arginine
C) Proline
D) Citrulline
☑ E) Isoleucine
3. (1.00 pts) Which of the following levels of structure would most accurately define the 3D structure of a protein?
O A) Primary
O B) Secondary
C) Tertiary
O D) Quaternary
4. (1.00 pts) High temperature, pH, or solute concentrations can all do which of the following to a protein?
O A) Combust
O B) Hydrolyze
O C) Fuse
D) Denature
5. (1.00 pts) Protein folding can occur in the nascent peptide prior to the release of the entire peptide from the ribosome.
● True ○ False
6. (1.00 pts) A protein with a low isoelectric point (pI) would have a highly negative charge at neutral pH
True  False
7. (1.00 pts) Proteins with transmembrane regions such as GPCRs will likely have high proportions of hydrophobic amino acids in regions in the membrane.

8. (1.00 pts) What does CRISPR stand for?
O A) Clustered Regularly Interspaced Short Pam-adjacent Repeats
B) Clustered Regularly Interspaced Short Palindromic Repeats
○ C) Clustered Regularly Immune Short Protective Repeats
D) Clustered Regularly Inserted Short Partial Repeats
9. (1.00 pts) Select all of the following that can be found in Cas9
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(Mark ALL correct answers)
☑ A) RuvC
B) HNH
C) 5' exonuclease domain
□ D) 3' exonuclease domain
<b>10. (1.00 pts)</b> What is the PAM sequence recognized by SpyCas9?
● A) 5' NGG 3'
○ B) 3' NGG 5'
○ C) 5' NCC 3'
O D) 3' NCC 5'
11. (1.00 pts) Which of the following DNA edits would Cas12a (cpf1) perform?
(
(Mark ALL correct answers)
A) Blunt end DSB
B) ssDNA nick
☑ C) Staggered DSB
<b>12. (1.00 pts)</b> Generation of catalytically inactive Cas9, or dCas9, could be performed with two total mutations in which lobes?
○ A) REC1, RuvC
O B) HNH, REC1
O C) RuvC, PAM-interacting
RuvC, HNH
○ E) REC1, CTD
Use the image below to assign structure names to the numerical labels provided.
Cas9
1 2 9
1 2 seed 9
(3)
5 + 10 .
(  6  ) RRR
6 RRR 7

	2.	
;	3.	
	4.	
	5.	
	6.	
	7.	
	8.	
!	9.	

10.

Expected Answer: NUC REC HNH Hel-II Hel-I RuvC Hel-III CTD crRNA tracrRNA

**14. (2.00 pts)** Select all of the following structures that contain an amide bond.

(Mark ALL correct answers)

- ✓ A) A
- ✓ B) B
- C) C
- ✓ D) D
- ✓ E) E
- ✓ F) F

15. (2.00 pts)	Select all of the amino acids have hydrophobic side chains.
(Mark <b>ALL</b> correct	answers)
□ A) A	
☑ B) B	
□ C) C	
D) D	
□ E) E	
✓ F) F	
16. (2.00 pts)	Select all of the amino acids that are frequently phosphorylated.
(Mark ALL correct	answers)
□ A) A	
□ B) B	
□ C) C	
D) D	
☑ E) E	
□ F) F	
17. (2.00 pts)	Which amino acid has a structural isomer that is indistinguishable using mass spectrometry?
(Mark ALL correct	answers)
□ A) A	
□ B) B	
□ C) C	
□ D) D	
□ E) E	
✓ F) F	
18. (2.00 pts)	Which amino acid will absorb radiation of wavelength 280 nm?
(Mark ALL correct	answers)
□ A) A	
□ B) B	
□ c) c	
□ D) D	
☑ E) E	
□ F) F	
40 (0.00 4-)	What along of any way from white at an atomic page 52
19. (2.00 pts)	What class of enzymes frequently act on structure E?
Expected Ansv	wer: phosphorylase or equivalent
20. (2.00 pts)	The carbamylation of the side chain of structure C would have what effect on the pl?
. ,	· · · · · · · · · · · · · · · · · · ·

Expected Answer: decrease

21. (2.00 pts) Provide an explanation for your answer above
Expected Answer: Following carbamylation, the carboxy and amino group will determine the pl, so it will decrease.
You sequence the genome of the rare species M. moopus and discover a highly repeated motif consisting of codons corresponding to hydrophobic and hydrophilic amino acids in a regular pattern. The DNA coding sequence for part of one such motif sequence is provided below:  5' CGC ATG CAG CTG GAA GAT AAA GTG 3'  A codon chart is provided below for reference.
Asp (E)  Asp
22. (3.00 pts) Provide the protein sequence in ONE letter amino acid code (N → C) that would be generated from the sequence. Do not include spaces.
Expected Answer: RMQLEDKV
23. (2.00 pts) At pH 7, what is the net charge on this protein fragment if it were part of a polypeptide backbone?
0

24. (2.00 pts) At pH 12, what is the net charge on this protein fragment if it were part of a polypeptide backbone?

-2
25. (2.00 pts) At pH 1, what is the net charge on this protein fragment if it were part of a polypeptide backbone?
2
26. (3.00 pts) You isolate RNA from a tissue sample of M. moopus and synthesize cDNA. You then decide to amplify fragments of cDNA containing the exact sequence of DNA shown below:  3' GCG TAC GTC GAC CTT CTA TTT CAC 5'  Which of the following primer combinations would be appropriate for this?
<ul> <li>A) 5'GCGTACGTCGACCTTCTATTTCAC 3'; 5' AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA</li></ul>
27. (5.00 pts) Your professor informs you that your method of amplifying cDNA would likely not result in whole protein sequences following cloning and expression. Provide an explanation for why this is the case.
Expected Answer: The first primer is located inside the cDNA sequence and will lack the start codon and the first part of the protein.
After performing a correct protocol, you are left with a set of 2 proteins. You note that when both proteins are mixed together in vitro with DNA, there is DNA binding activity. You tag one protein (A) with rhodamine and another protein (B) with fluorescein to perform FRET.  The peak absorbance / emission wavelengths (nm) for the two dyes are provided below:  • Rhodamine: 490 / 530 nm  • Fluorescein: 530 / 603 nm
28. (2.00 pts) If a solution containing tagged protein A only is excited at 490 nm, what would the wavelength of the peak emission be?
Expected Answer: 530
29. (2.00 pts) If a solution containing tagged protein B only is excited at 530 nm, what would the wavelength of the peak emission be?
Expected Answer: 603 nm

A solution containing tagged protein A and B together is excited at 490 nm. Two peaks of emission are observed, one at 530 nm and the other at 603 nm. You suspect this is due to dimerization. Explain why your hypothesis is consistent with the results above.

30. (5.00 pts)

<b>Expected Answer:</b> The donor chromophore, which is rhodamine tagged to protein A, can transfer energy to the acceptor chromophore (fluorescein tagged to protein B) when the two fluorophores are very close together (within several nm). For this to occur, the two proteins must dimerize.
You prepare a solution containing tagged proteins A and B and bleach the fluorescein dye. Upon bleaching, the fluorophore is inactivated and will no longer emit light following excitation. You add tagged protein B to the solution in 5x excess of the original amount then excite at 490 nm and record the emission spectra.
31. (3.00 pts) You suspect that proteins in the A-B dimer can freely exchange with subunit proteins in solution. Following sufficient time to reach equilibrium, identify all emission peaks expected in results consistent with rapid exchange kinetics of dimer subunits.
(Mark <b>ALL</b> correct answers)  A) 490 nm
✓ B) 530 nm
☑ C) 603 nm
<b>32.</b> (4.00 pts) Provide an explanation for the choices selected above.
Expected Answer: Free exchange of dimer subunits would allow for unbleached protein B to dimerize with protein A. Thus, an emission peak would be expected at 603 nm.
33. (3.00 pts) You suspect that proteins in the A-B dimer cannot freely exchange with subunit proteins in solution. Following sufficient time to reach equilibrium, identify all emission peaks expected in results consistent with very slow exchange kinetics of dimer subunits.
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<b>Expected Answer:</b> Bleached protein B will remain bound DNA binding will prevent exchange of subunits.	d to protein A in a dimer binding DNA. There would be no expected fluorescence at 603 nm. Answer should discuss how tight
The Michaelis-Menten constant	
	$K_m$
is often thought of as analogous to the enzymes dissocial	tion constant
	$K_d$
and both are frequently referred to as a measure of the e	nzymes affinity for the substrate.
	ted below. E is a free enzyme, ES is the enzyme-substrate complex, S is the substrate, and P is the product. Reaction rates
$E + S \stackrel{k_1}{\rightleftharpoons} ES \stackrel{k_{\text{cat}}}{\rightarrow} E +$	
$E + S \Longrightarrow ES \longrightarrow E +$	. <b>p</b>
$\frac{1}{\nu}$	•
$\kappa_{-1}$	
Differential equations can be used to describe the rates. F	−or example,
	$rac{dx}{dt}=0.5$
would indicate that the quantity x is increasing by 0.5 unit	s at all t. This would correspond to a linear relationship between x and t.
For each of the quantities below, express the following dif	fferential equation in terms of the values above. Be sure to include brackets if indicating concentrations.
	d[E]
1.	$\frac{1}{dt}$
2.	$rac{d[S]}{dt}$
<del></del> -	
3.	$rac{d[ES]}{dt}$
4.	$rac{d[P]}{dt}$
	$a\iota$
<b>37. (4.00 pts)</b> 1.	
Expected Answer: -k_1[E][S] + k1 [ES] + k_cat [ES]	

**36. (4.00 pts)** Provide an explanation for the choices selected above.

**38. (4.00 pts)** 2.

Expected Answer:  $-k_1[E][S] + k_-1[ES]$ 

39. (4.00 pts)	3.	
Expected Ans	swer: k_1[E][S] - k1[ES] - k_cat[ES]	
40. (4.00 pts)	4.	
Expected Ans	swer: k_cat[ES]	
41. (3.00 pts)	Express the dissociation constant $K_d$	
	in terms of the provided rate constants.	
Expected Ans	:wer: k1 / k_1	
42. (4.00 pts)	The total amount of enzyme, $[E_0] = [E] + [ES]$	
	$[E_0] = [E] + [ES]$ . Assuming the steady-state equilibrium holds, express the dissociation constant K_d in terms of [E_0], [S], and [ES].	
Expected Ans	swer: ([E_0]-[ES])[S] / [ES]	
The formula for	r $K_m$	
is provided belo	ow for reference.	

 $K_{\rm m} = \frac{(k_{-1} + k_{\rm cat})}{k_{1}}$ 

43. (4.00 pts)	In what situation could the value of		
	$K_m$		
	be close to		
	$K_d$		
	?		
Expected Ansv	wer: k_cat << k1; accept "when k_cat is small / very small"		
44. (5.00 pts)	If you choose to use		
	$K_m$		
	instead of		
	$K_d$		
	to show binding affinity, would this consistently underestimate or overestimate the binding affinity to the substrate? Explain your answer.		
Expected Ansv	wer: Constantly underestimate (2). K_m will have a value higher than K_d. Higher values relative to K_d imply that the enzyme has less affinity to the substrate.		
	Burke plot assisted early biochemists in linearizing data to determine enzyme parameters such as K_m and V_max. Despite its usefulness, there are several etting a regression line in this manner.		
	on the right of the graph are collected at [1] substrate concentrations and should thus correspond to [2] reaction rates. If we assume experimental error affects		
	and not substrate concentration, then [3] y-values are highly prone to error.		
The Eadee-Hof	The Eadee-Hofstee plot expresses a different relationship, but maintains the same principle of relating experimental values to a linear plot. The equation is provided as follows:		
v = V_max - K_	m * v / [S]		
Experimental er	rors in measuring reaction rates would result in deviations parallel to the axis plotting [4].		
45. (4.00 pts)	Provide responses to blanks 1-4 below:		
	1.		
	2.		
	3.		
	4.		
Expected Ans	were low alow high account towards or away from the origin		
Expected Ansi	wer: Low, slow, high, accept towards or away from the origin		

## Research Literacy

In this section, you will be presented with excerpts from a few works studying CRISPR. You will be tested on how well you can apply your background knowledge of CRISPR to understand the information presented.

## DNA capture by a CRISPR-Cas9-guided adenine base editor

Note: ABE8e refers to the new adenine base editor proposed in the study.

Lapinaite A, Knott GJ, Palumbo CM, Lin-Shiao E, Richter MF, Zhao KT, Beal PA, Liu DR, Doudna JA

sgRNA, or ABE8e complexed with a nontargeting sgRNA. Apo-ABE8e was able to deaminate only ssDNA, whereas ABE8e RNP was able to modify adenines in ssDNA and dsDNA, in contrast to the absence of trans-dsDNA editing by ABE8e engaged in an R-loop complex. Moreover, the dsDNA deamination is sgRNA sequence independent because a nontargeting ABE8e RNP also deaminates dsDNA adenines, suggesting that stable R-loop formation is not required. To test whether Cas9's interaction with the PAM affects the observed dsDNA editing by ABE8e, we performed an in vitro DNA deamination assay with radiolabeled dsDNA devoid of consensus PAMs. To our surprise, ABE8e RNPs containing both the targeting and nontargeting sgRNA deaminated adenine in dsDNA lacking PAMs in the NTS.
46. (3.00 pts)
Deamination of adenine would result in a conversion to, which is recognized by DNA replication machinery as, resulting in a base pair edit to following round(s) of replication.
Expected Answer: hypoxanthine, guanine, G-C, 1 (0.5 pts each for last 2)
47. (5.00 pts) The general structure of ABE8e is an ssDNA deaminase fused to modified Cas9. Provide a hypothesis as to why apo-ABE8e is only able to modify ssDNA.
Expected Answer: apo-ABE8e will lack the sgRNA (2) which would melt a short stretch of DNA with homology (1) The attached deaminase will have no opportunity to act on dsDNA
without local unfolding (2)
48. (5.00 pts) The researchers chose to use a nickase Cas9 instead of dCas9. What was the purpose of including partial catalytic activity in ABE8e?
Expected Answer: Allows the recruitment of DNA repair machinery that would replicate the stretch of DNA that was modified to preserve the edit.
Expense 7 mente in a formal manufactory and media replicate the earter of Britain and media to present the earter
49. (10.00 pts) You hypothesize that the search mechanism of this particular Cas9 protein involves transient melting of DNA as the protein "scans" for a PAM motif. Is this consistent with the results
presented in the excerpt? If so, provide a thorough explanation as to why, referring back to the experimental results presented.
Expected Answer: Yes (1). ABE8e with nontargeting sgRNA (implying no homology to DNA sequence) was able to deaminate adenine even in the absence of a PAM motif. (3) Apo-ABE8e was unable to deaminate dsDNA, which proves that deamination will not occur when Cas9 cannot bind DNA. (1). Thus, for deamination to occur, there must be local melting of the DNA sequence to expose ssDNA. The hypothesis is consistent with the presented results since transient melting would expose ssDNA and allow for the deaminase to act. (5)
50. (3.00 pts)  Fusion of Cytidine deaminase was used similarly to catalyze the conversion of cytosine bases to uracil. The immediate correction of the mismatch is prevented by Uracil DNA Glycosylase Inhibitor. If a different deaminase targeted 5-methylcytosine, what protein would have to be fused to the base editor for maximum efficacy?
Expected Answer: thymine-DNA glycosylase inhibitor / TDG inhibitor

Deamination assays were conducted using either radiolabeled ssDNA or dsDNA substrates and ABE8e in three different states: apo-ABE8e, ABE8e complexed with a targeting

	Thanks for taking this test! If you want to leave any feedback, please do so at https://tinyurl.com/utreg21feedback (https://tinyurl.com/utreg21feedback).
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	Terms of Use (/ToS.html)   Privacy (/Privacy.html)   Cookie Policy (/CookiePolicy.html)   Support (/tx/Support)   Contact (/tx/Home/Contac