COMPLETE THIS SECTION: Up to TWO POINTS will be removed for incorrect/missing information!					
Complete in this section . Sp to two points will be removed for incorrections sing information:					
PRINTED FIRST NAME	PRINTED LAST NAME				
Person on your LEFT (or Empty or Aisle)					
Person on your RIGHT (or Empty or Aisle)					
Class you are REGISTERED FOR (onground or hybrid)					
The room where most students will take the test for your class, i.e. LS A-191 for onground and PS H-152 for hybrid)					

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- PRINT YOUR NAME ON EACH PAGE!
- READ THE DIRECTIONS CAREFULLY!
- USE BLANK PAGES AS SCRATCH PAPER

work on blank pages will not be graded...

- WRITE CLEARLY!
- MOLECULAR MODELS ARE ALLOWED
- DO NOT USE RED INK
- DON'T CHEAT, USE COMMON SENSE!

н			He Interacti	on Energies, kcal/mol
Li Be		B C N O F	Ne Eclipsis	ng Gauche
Na Mg		Al Si P S Cl	11	4 Et/Me ~0.95
K Ca S	c Ti V Cr Mn Fe Co Ni	Cu Zn Ga Ge As Se Br	r Kr Et/H ~1	8 i-Pr/Me ~1.1
Rb Sr Y	Zr Nb Mo Tc Ru Rh Pd	Aq Cd In Sn Sb Te I	Vo	t-Bu/Me ~2.7
Cs Ba L	u Hf Ta W Re Os Ir Pt	Au Hg Tl Pb Bi Po At	11	2.8
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	range of values broad peak	N-H C=O strong) -	H `C´
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	0	Aromatic Ar—H	··20	-H ₂ C -NR ₂
	О -С-ОН <u>—С-Н</u>	mainly 8 - 6.5	$C = CH_2$ C	
-	11 10 9	8 7 6	5 4 3	——————————————————————————————————————
(δ, ppm) –	220 200 180	160 140 120	100 80 60	0 40 20 0
	0	R	-C≡N	$\frac{H_2-}{Alkyl} 3^{\circ} > 2^{\circ} > 1^{\circ} .$
⊢	R-C-OH O	R ₂ C=CR ₂	RC≡CR, ,	C-X
	<u> </u>	Aromatic	——————————————————————————————————————	C-NR ₂

YOU MUST COMPLETE THIS PAGE WITH YOUR NAME (EVEN THOUGH YOU ALREADY DID THIS ON THE COVER PAGE) AND ALSO GIVE YOUR ASU OR POSTING ID NUMBER WE NEED THIS NUMBER BECAUSE YOU WOULDN'T BELIEVE THE NUMBER OF STUDENTS WHOSE NAMES WE CAN'T READ!

PRINTED FIRST NAME	PRINTED LAST NAME		ASU ID or Posting ID	
	Points by que	estion		
	1	/14		
	2	/18		
	3	/18		
	4	/8		
	5	/34		
	6	/28		
	7	/27		
	8	/28		
	Points Removed for cover en	rors/2		
	Extra Credi	t/5		
	Total (incl Extra)_	/175+5		

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Question 1 (14 pts.) Give the IUPAC name for the following structure.

Question 2 (18 pts.)

Which has a higher frequency bond vibration observable in an infrared spectrum, as a peak at higher frequency, the C-N triple bond of a nitrile functional group (**A**) or the C-N double bond of an imine (**B**)? Give a BRIEF explanation.

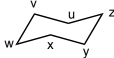
A
$$H_3C$$
— C $\equiv N$:

$$C = N$$

Question 3 (18 pts). Between the C=N double bond in an imine and the C=O double bond in a ketone, which would you expect to have a stronger or more intense peak in an IR spectrum? Give an explanation for your choice.

Question 4 (8 pts.)

a) Shown below are two conventional drawings of a cyclohexane chair. In one, the carbon atoms are labeled u through z. Label the atoms on the **other chair** with u, v, w, x, y and z according to the normal understanding of which atom is which in each chair, i.e. atom u in the first chair should be labelled u in the other chair etc.





b) Which of the 6 atoms, u, v, w, x, y or z, are understood to be ABOVE the plane of the paper in the conventional drawings of a cyclohexane chair, i.e. which atoms are understood to be closer to you?

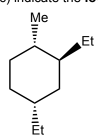
Question 5 (34 pts.) For rotation around the C2-C3 bond of 2,2,3-trimethylpentane:

a) Draw Newman projections AND 3D (sawhorse) structures for BOTH the HIGHEST and the LOWEST energy conformations, (clearly indicate which is which). **Your Newman projections should look FROM carbon 2 TO carbon 3.**

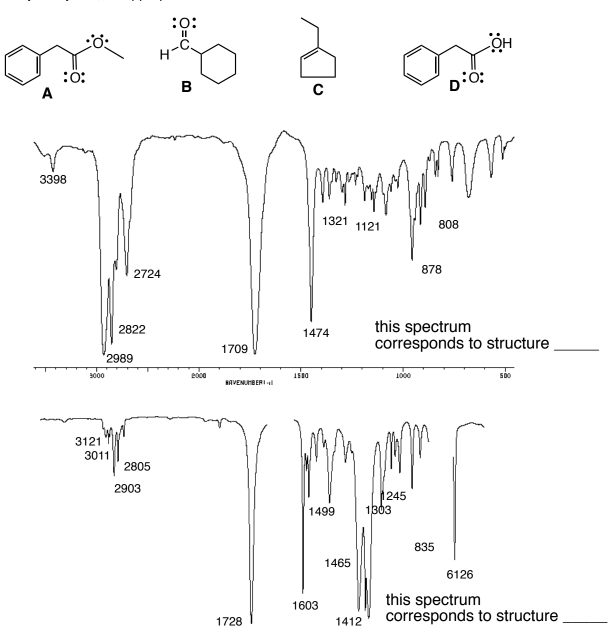
c) Determine the energy difference between these two conformations. CLEARLY indicate the energy factors that you included in your calculation of this energy difference

Question 6 (28 pts) For the following structure

- a) draw both chair conformations
- b) determine the **energy difference** between the two chair conformations. CLEARLY indicate the energy factors that you included in your calculation of this energy difference
- c) indicate the lower energy chair



Question 7 (27 pts.) Assign the BOTH IR spectra to ONE of the FOUR provided structures A - D. Two of the structures do not have a provided spectra. On each spectrum, identify the peaks that are associated with a specific functional group or type of C-H bond by drawing the functional g or bond and drawing an arrow from the specific bond in the functional group that vibrates to t absorption peak, as appropriate.



1500

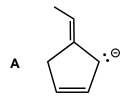
HAVENUMBER! -:!

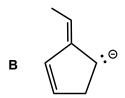
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Question 8 (28 pts) For the following two anions A and B

- a) Draw all reasonable resonance contributors including curved arrow pushing and resonance arrows/brackets as appropriate, and **idendity** the major contributor if there is one.
- b) Which of the anions **A** or **B** would be more chemically reactive? Give a brief explanation that includes the term "energy of the electrons"
- c) For both **A** and **B**, draw an "actual" structures and use the delta symbol to indicate the atoms with fractional charges

- 5 -





d) Anion **A** is drawn for you again below. In this drawing (below), circle all of the carbon atoms that are part of the conjugated system.

e) Anion ${\bf A}$ is drawn for you yet again below. In this drawing, give the hybridization of the 3 atoms indicated by the arrows

EXTRA CREDIT (5 pts) Which types of structures are the primary constituent of gasoline?

alkenes

ketones

alkanes

aromatics