MTW5 9-26

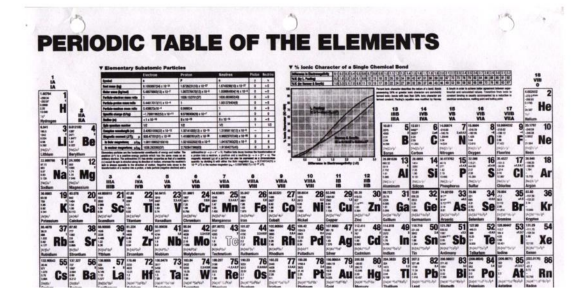
Tuesday, September 20, 2022 2:58 PM

Fl22Exam1Ques	
NAME (Print):	Chemistry 520W 520W
EID	Dr. Brent Iverson 1st Midterm September 22, 2022
SIGNATURE:	
Please print the	
first three letters of your last name in the three boxes	
though it is really a one hour exam. The id	are giving you three hours to take this exam even dea is to give you enough time to show us what
you know, not how fast you can draw struc the best possible structures that you can! the exam long before 9 PM. That is to be	ctures. Please take all the time you need to draw Do not be surprised if you are comfortable leaving expected!
ALLOWED TO INTERACT WITH YO TOUCH YOUR CELL PHONE DURI	JNATE RECENT INCIDENCTS YOU ARE NOT DUR CELL PHONE IN ANY WAY. IF YOU NG THE EXAM YOU WILL GET A "0" NO
MATTER WHAT YOU ARE DOING V IT THERE!!!	VITH THE PHONE. PUT IT AWAY AND LEAVE

Student Honor Code

"As a student of The University of Texas at Austin, I shall abide by the core values of the University and uphold academic integrity."

(Your signature)

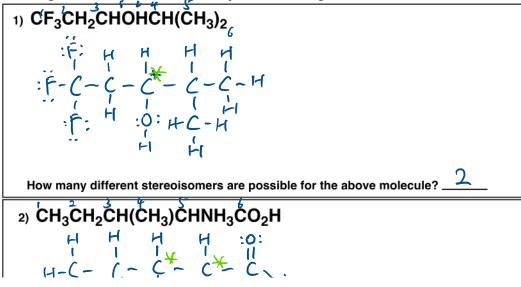


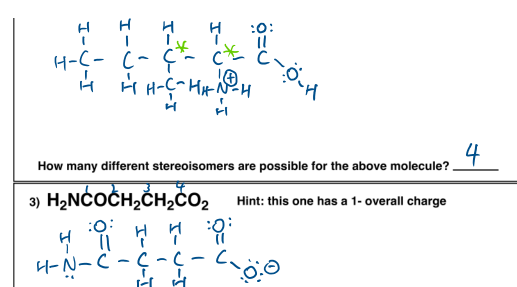
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1. (4 points) What is the most important question in Organic Chemistry?

2. (8 pts each) For the following molecular formula, draw complete Lewis line structures in which all atoms (even H atoms) are drawn, lines are used as bonds, all lone pairs are drawn AND ALL FORMAL CHARGES ARE INDICATED. Note you must infer the formal charges as we do not indicate them on the chemical formulas given. You only have to draw one important contributing structure if that is relevant.

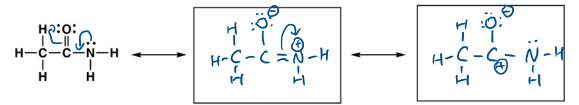




How many different stereoisomers are possible for the above molecule? $\frac{2^{\circ}}{2}$

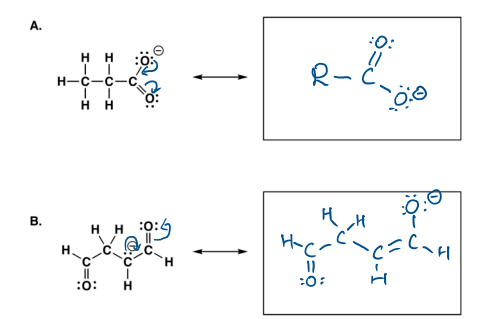
Signature_____ Pg 2 _____(12)

3. (6 pts) I told you this would be here. The following amide molecule is best represented as the hybrid of three contributing structures. **Draw the second and third important contributing structures** in the spaces provided, including all lone pairs and formal charges.



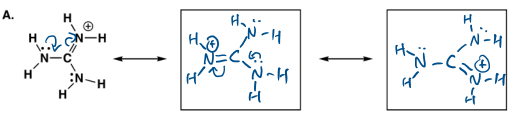
4. (6 pts) The following molecules are best represented as the hybrid of contributing structures. **Draw the second important contributing structure** in the space provided, including all lone pairs and formal charges.

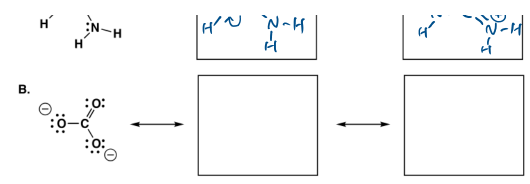
4. (6 pts) The following molecules are best represented as the hybrid of contributing structures. **Draw the second important contributing structure** in the space provided, including all lone pairs and formal charges.



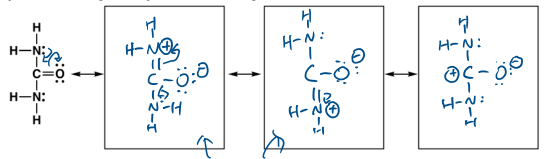


5. (12 pts) The following molecules are best represented as the hybrid of three contributing structures. **Draw the second and third important contributing structures** in the spaces provided, including all lone pairs and formal charges.

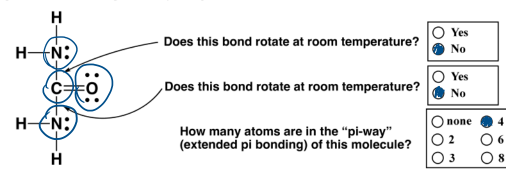




6. (9 pts) The following molecule is called urea and is best represented as the hybrid of four contributing structures. **Draw the second, third and fourth important contributing structures** in the spaces provided, including all lone pairs and formal charges.



7. (10 pts) On the following molecule, circle all the atoms that are sp^2 hybridized and answer the three questions in the boxes provided by filling in the circles next to the correct answers.



Signature

Pg 4 _____(17)

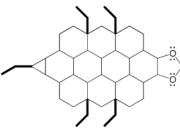
8. (1 pt each) Fill in each blank with the word or words that best completes the sentences.

8. (1 pt each) Fill in each blank with the word or words that best completes the sentences. For organic chemistry, it is best to think of electrons as The electron density in molecules can be described mathematically by adding the functions of all the atomic orbitals for all the atoms in the entire molecule, an approach refered to as The wave functions for the **Valence** atomic orbitals on each atom can be added together first, a process referred to as ______, before looking for overlap with orbitals from other atoms. This aproach is called ______ theory. You need to be able to think about all ______ bonding in molecules as being derived from the overlap of ______ orbitals and all pi bonding as being derived from overlap of unhybridized orbitals. Especially for charged molecules, it is stabilizing to have ______ electron density as well as charges delocalized over more than two atoms. This concept is usually referred to as "stabilization due to de 6 (a (26 for " or simply "resonance stabilization". For pi bonding and therefore pi delocalization to occur over more than atoms (i.e. pi-ways), parallel ______ orbitals are needed on ALL of the adjacent atoms involved, explaining why ALL of these atoms must be (or hybridized and why these sytems are planar.

NEWS FLASH: A new species of turtle was just discovered in the turtle pond:

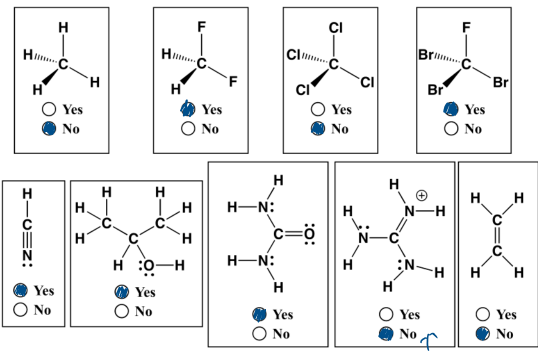
According to IUPAC, its official name is:

(7aR,8aS,11r,13aR,14aS)-7a,8a,11,13a,14apentaethyldotriacontahydro-1*H*-cyclopropa[10,11]ovaleno[3,4*d*][1,3]dioxole

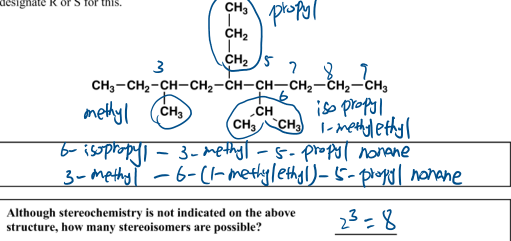


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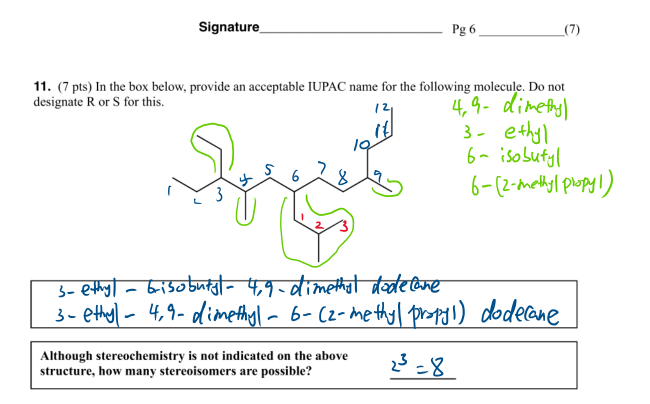
9. (18 pts) Indicate which of the following molecules have an overall molecular dipole moment. You do not need to indicate the direction of the dipole moment, or any of the individual bond dipoles. Fill in the circle next to "Yes" if the molecule has an overall molecular dipole, or "No" if the molecule does not have an overall molecular dipole moment.



10. (7 pts) In the box below, provide an acceptable IUPAC name for the following molecule. Do not designate R or S for this.



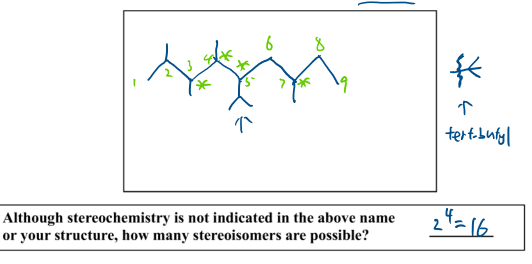
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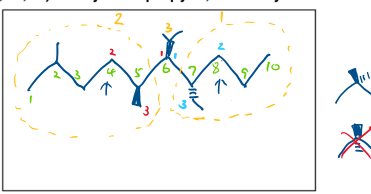


12. (10 pts each) For the following IUPAC name, draw the appropriate line angle drawing. You can ignore R and S for this one.

5-isopropyl-2,3,4,7-tetramethylnonane



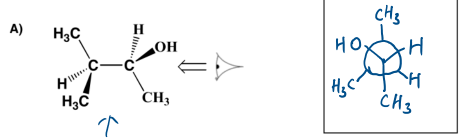
13. (10 pts each) For the following IUPAC name, draw the appropriate line angle drawing. For this one, you need to use wedges and dashes to indicate the appropriate stereochemistry at all chiral centers.



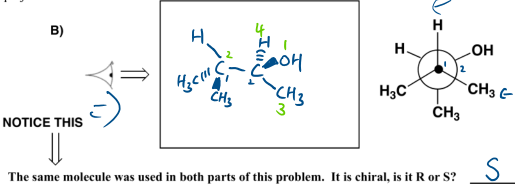




14. (5 pts) Draw the Newman projection for the conformation of 3-methyl-3-butanol as shown.

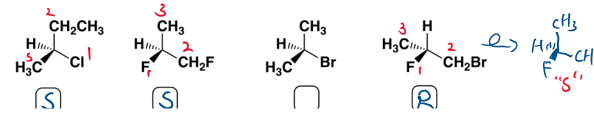


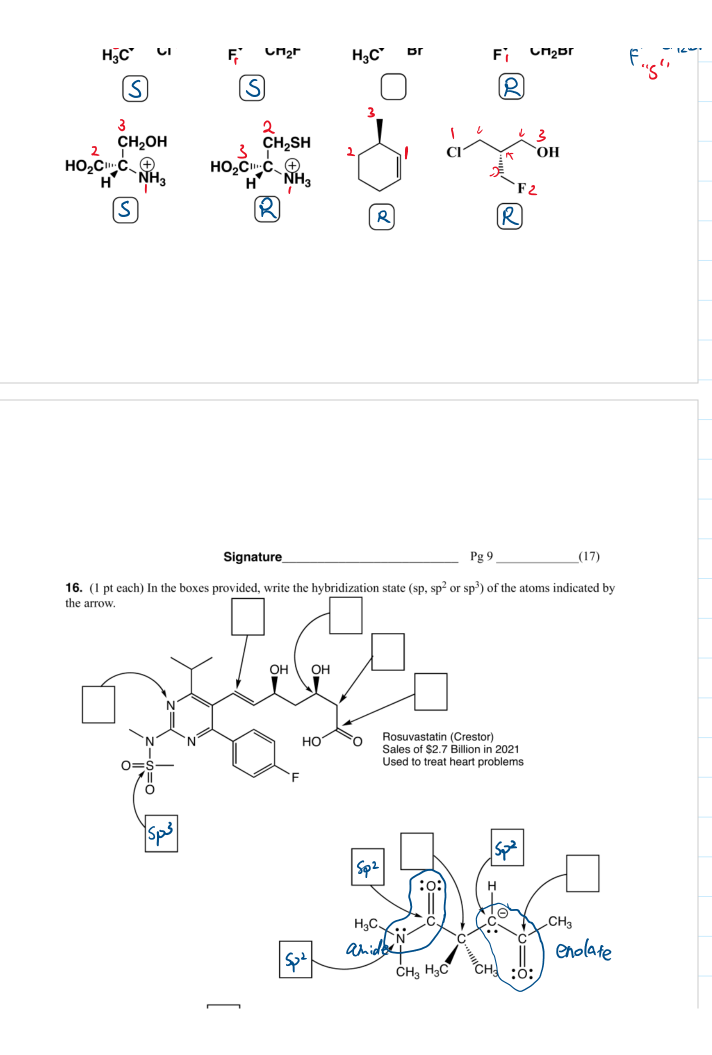
(7 pts) In the empty box draw the conformation of 3-methyl-3-butanol indicated by the Newman projection shown.

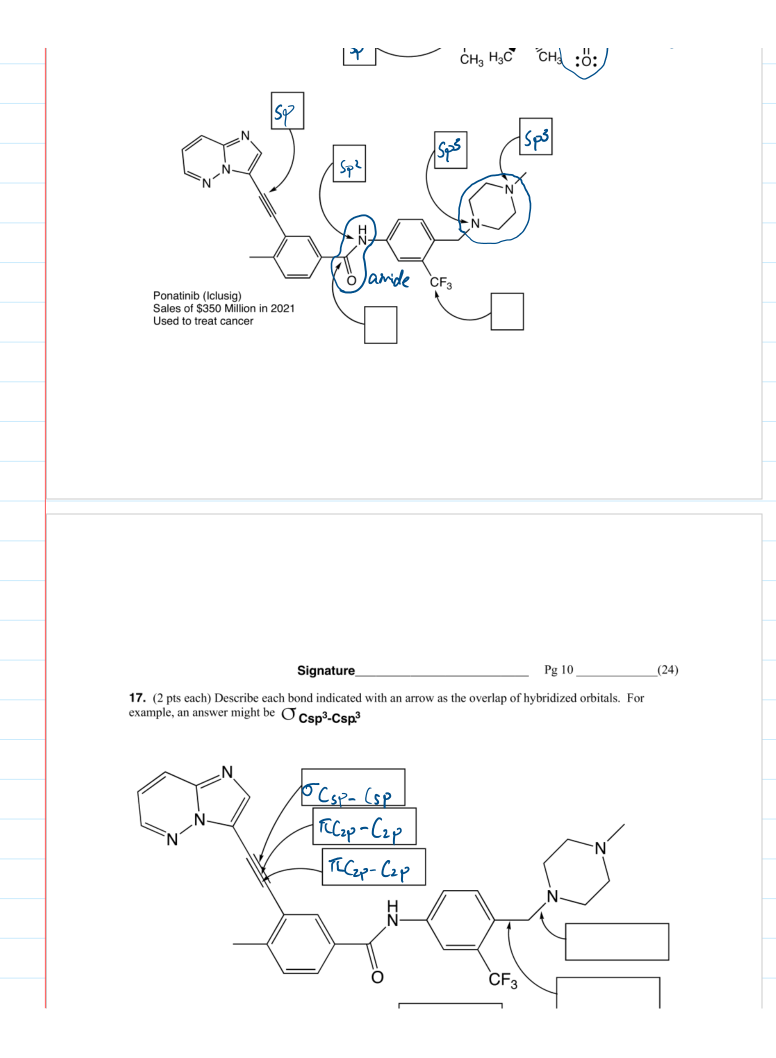


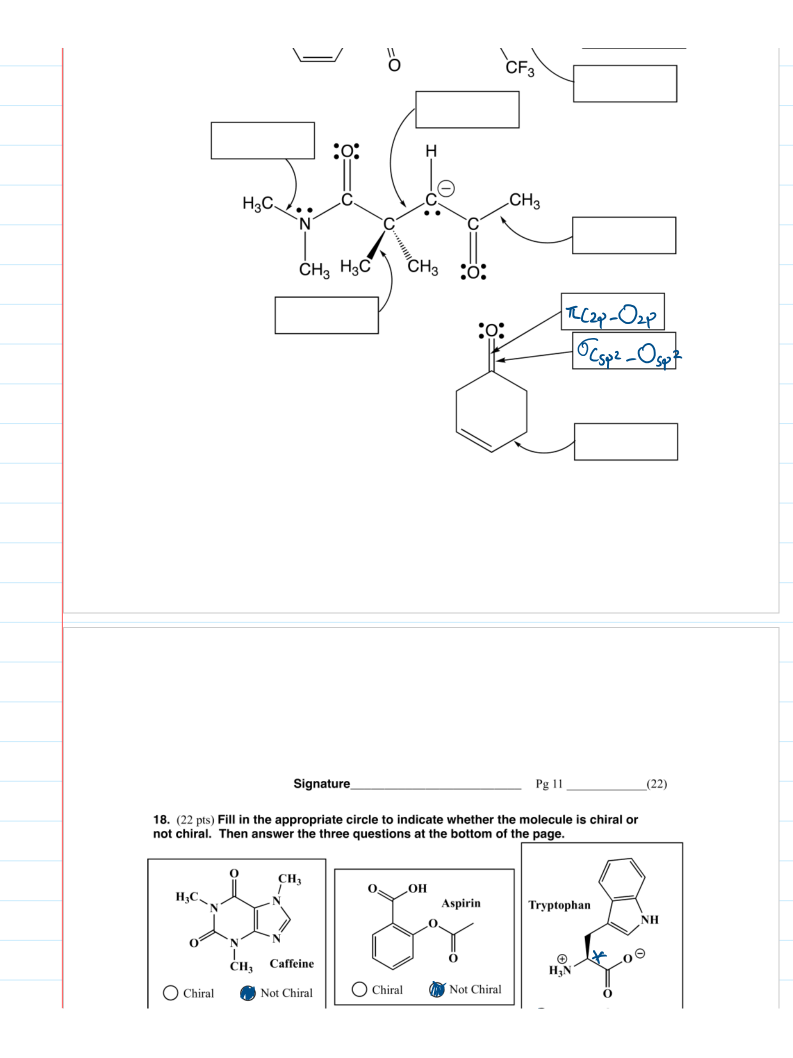
15. (2 pts each) Examine the following structures. For each molecule with a chiral center, assign the

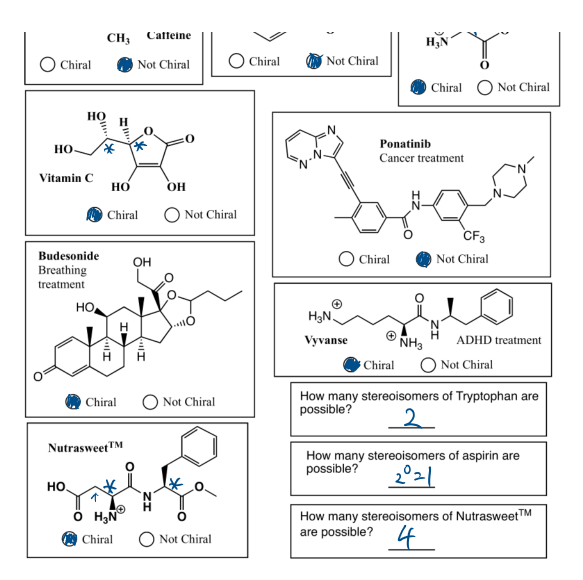
stereochemistry then write "R" or "S" as appropriate in the box provided below each structure. For all molecules that have no chiral centers, do not put anything in the box.





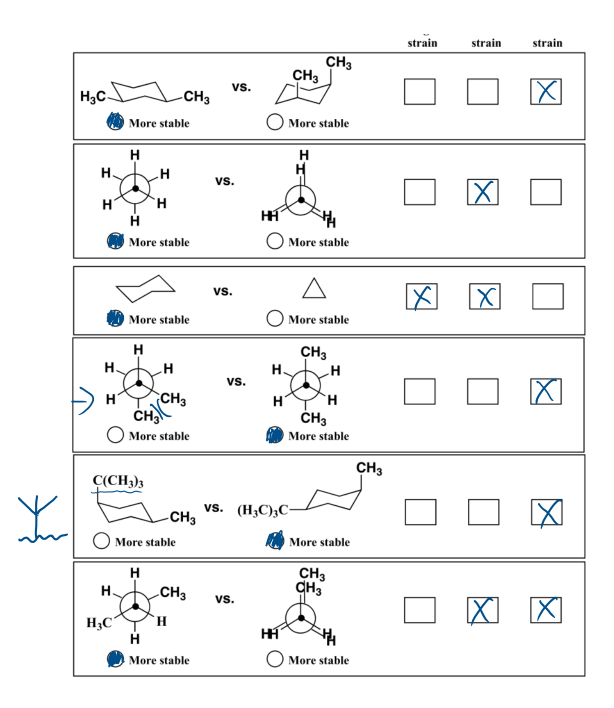






Signature	Pg 12	2	_(24)
19. (4 pts each) For each pair of molecules, fill in the circle under the one put an "X" in the box under all the types of strain that explain(s) your answ		e stable of the tw	o, then
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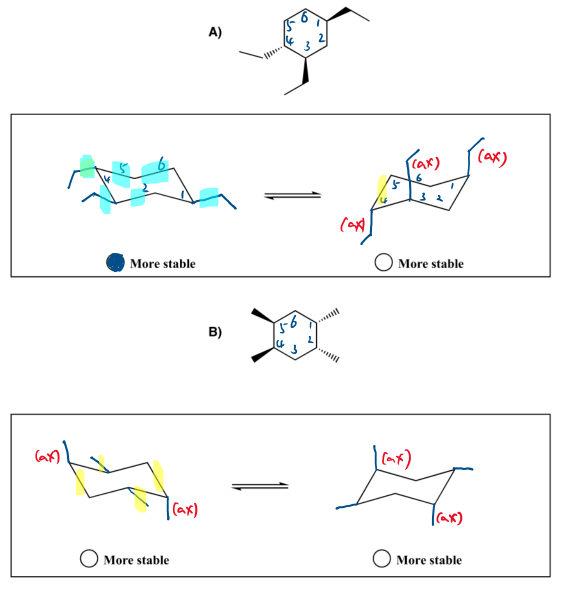
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 $CH_3 \mid$			



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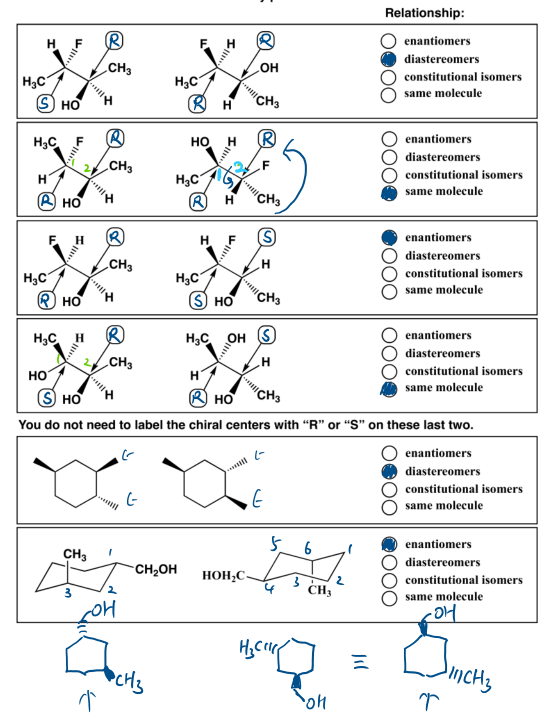
Pg 13 _____(20)

20. (20 pts) For the following cyclohexane derivatives, draw the two alternative chair conformations. IF there is a difference in stability, fill in the circle that says "More stable". If there is not any difference in stability, do not fill in any circle.



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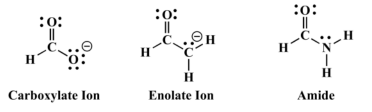
21. (38 pts) For each pair of molecules, on the line provided state the relationship between the two structures. Possible answers could be enantiomers, diastereomers, consitutional isomers, or same molecule. Fill in the circle to indicate the correct relationship between the molecules shown. In the boxes provided next to each chiral center, write "R" or "S" to indicate the absolute stereochemistry present.



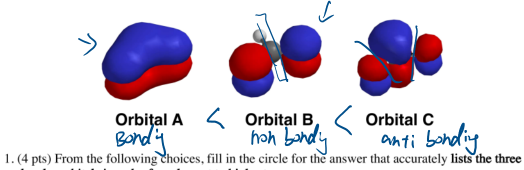
_____ Pg 15 _____ Signature____ (4)

22. (22 points total). Here is an "apply what you know" problem in the form of an MCAT style passage.

We have seen a number of molecules that contains a three-atom, delocalized pi bond (a "pi-way"). In partcular, we have seen a carboxylate ion, enolate ion, and of course, amides. Each of the three atoms donates a 2p orbital that overlap.



When the three 2p orbitals overlap, there are three pi molecular orbitals formed that extend over all three atoms. You have seen these before, and they are shown below:



molecular orbitals in order from lowest to highest energy:

💭 Orbital A Orbital B Orbital C Orbital B Orbital C Orbital A Orbital A Orbital C Orbital B Orbital C Orbital B Orbital A

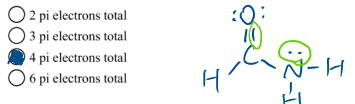
CHz

One of the more difficult parts of the analysis of delocalized pi bonding concerns how many electrons are involved in the pi molecular orbitals. Each of the ions shown above, the carboxylate ion, the enolate ion, and the amide, have the same number of pi electrons in the pi molecular orbitals.

Signature	_ Pg 16	(8)
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22 (cont).

2. (4 pts) Fill in the circle for the answer that **lists how many electrons reside in these pi molecular** orbitals in the carboxylate ion, the enolate ion and the amide?



3. (4 pts) Fill in the circle for the answer that lists which of the orbitals are filled by the electrons you listed in part two above.

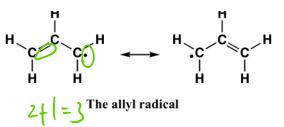
Orbitals A, B and C
Orbitals A and B
Orbital A only
Orbitals A and C

Your answer to part 3. explains the properties of the carboxylate ion, the enolate ion and the amide.

For example, let's consider the carboxylate ion. By understanding which of these orbitals are filled, it explains why there is partial double bond chacacter over all three atoms (O-C-O) and why the negative charge is located on only the two oxygen atoms.

Later this semester you will learn that there are some highly reactive intermediates that have an unpaired electron in their valence shell, and these are called "radicals". One example is shown below, it is called the "allyl radical" and it is best described as the resonance hybrid of two contributing structures. Note that radicals have no formal charge!





It turns out that all of the carbon atoms of the allyl radical are sp^2 hybridized and the three 2p orbitals overlap to create the same three pi molecular orbitals shown above in this problem.



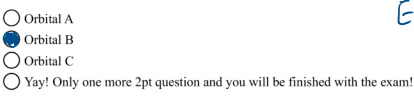
22 (cont).

4. (4 pts) Fill in the circle for the answer that lists **how many electrons reside in these pi molecular orbitals in the allyl radical?**

- 2 pi electrons total
- 🔵 3 pi electrons total
- 4 pi electrons total
 4
- 6 pi electrons total

One of the pi molecular orbitals of the allyl radical is only half-filled, containing the electron density for only a single electron, not a pair of electrons! That is why radicals are so reactive, they react to make bonds that fully fill their valence shell.

5. (4 pts) Given everything you know about delocalized pi molecular orbitals, fill in the circle for the answer that correctly lists **the pi molecular orbital that is half-filled in the allyl radical**.





6. (2 pts) Examine the contributing structures for the allyl radical above, does the orbital you selected in part 5. make sense to you?

6. (2 pts) Examine the contributing structures for the allyl radical above, does the orbital you selected in part 5. make sense to you?

O No, there is no correlation between the molecular orbitals and contributing structures

Yes, the orbital I chose and the contributing structures place the unpaired electron density on the same atoms.

A good way to get ready for a 5K race is to remember that avoiding a running injury means being patient and increasing your distance slowly. Start by running as far as you can comfortably. Do not push it at the beginning. Let's say you can run 1 mile before feeling too out of breath. Run that 1 mile 2-3 times a week at first, making sure you have no foot or leg pain. If you do have foot/leg issues, try new running shoes fit by a professional (The Loop or Rogue Running are great running stores for this). After you are comfortable running 1 mile for a week, try 1.25 miles for 2-3 times the next week. Then run to 1.5 miles, then 2.0 miles, then 2.5 miles each 2-3 times for a week. It will then be time for the race and you will make it!!!