

NAME _____

ID # _____

ORGANIC CHEMISTRY I (2301)

9:30 – 10:20 am, June 30, 2015

Exam 1

If you want to pick this exam up on Thursday in class (in public), please check the box on the right:

If you do not check the box, I will not bring your exam to class on Thursday, and you will need to pick up your exam in private from Chemistry department staff in 115 Smith beginning Thursday afternoon. Exams that are not picked up within two weeks will be disposed of.

A periodic table is attached to the back of this exam as an aid. Otherwise, you are not permitted to use any other materials (including notes, books, or electronic devices of any kind).

Right now, write your name and student ID number at the top of this page. When the exam begins, please write your name at the top of the next page.

You may use pen or pencil. However, re-grades will be considered only for exams completed in pen.

Please write your answers in the boxes/spaces provided. If your answer is not in the appropriate space (say, for example, it's on the back of the page), draw us an arrow and/or note telling us where to look.

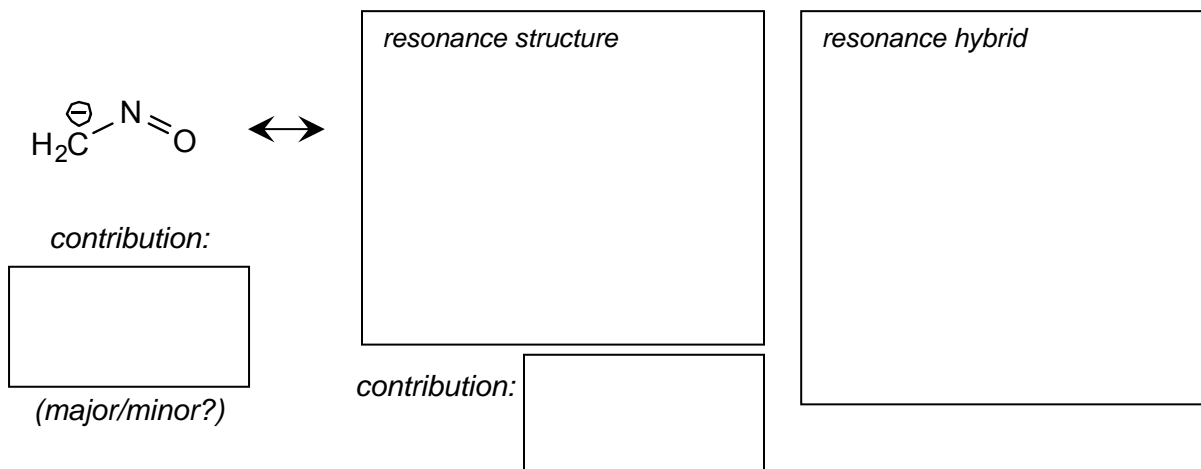
NAME _____

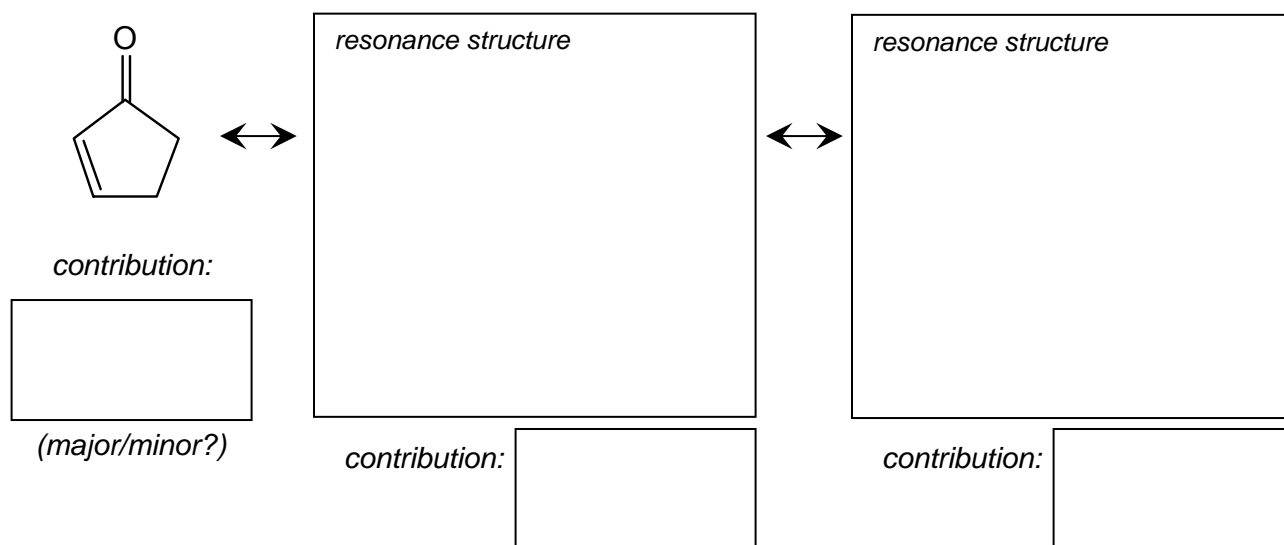
Scoring: 1. _____ / 6 5. _____ / 18
 2. _____ / 20 6. _____ / 20
 3. _____ / 12 7. _____ / 6
 4. _____ / 18

Total Score: _____ / 100

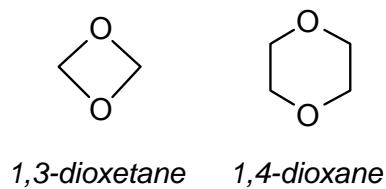
1. (6 pts) **Draw Lewis dash-bond structures** for two constitutional isomers that have molecular formula CH_4N_2 , and that have **no formal charges** on any atom. Draw all atoms and lone pairs of electrons.

2. (20 pts) For each of the molecules on the left, draw as many of the best Lewis dash-bond resonance structures as there are boxes to put them in. (Feel free to omit lone pairs and C-H's, or draw them—your choice.) Then, below each resonance structure, describe whether each would be a major or minor contributor. Finally, draw a resonance hybrid that illustrates partial charges and multiple bonds.

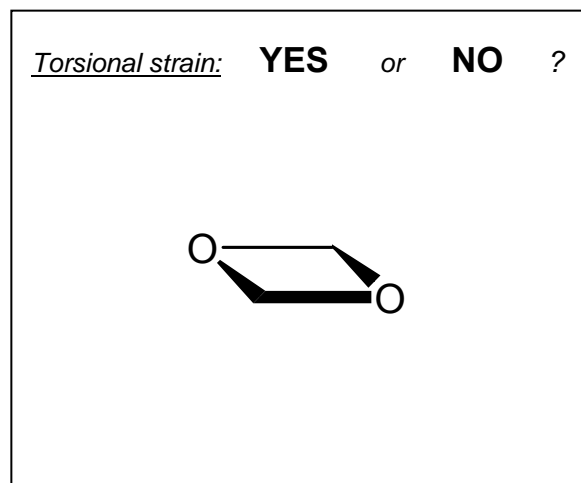
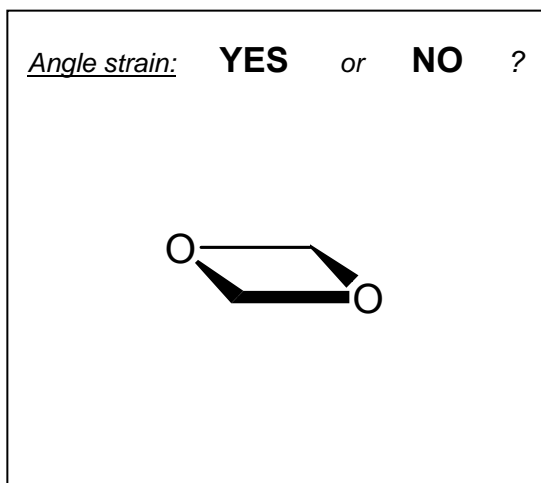




3. (12 pts) This problem compares 1,3-dioxetane and 1,4-dioxane (shown at right).



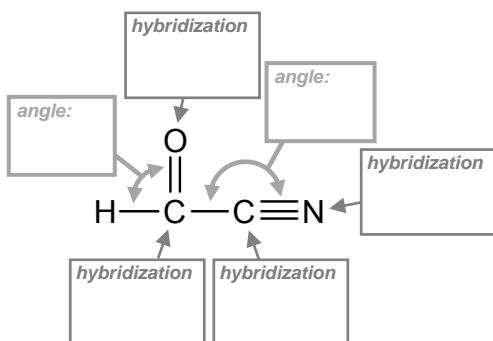
- a. *Angle strain* and *torsional strain* are both repulsive forces between pairs of electrons that make some cyclic molecules less stable than others. **Does 1,3-dioxetane show angle strain and/or torsional strain?** Circle your answers in the boxes below. Then, if you circled “YES” in either box, **illustrate the electron-pair repulsion** (the strain) on my skeleton structure **with a double-headed arrow**. You may have to add to my drawing in order to show the repelling electrons. (If you circled “NO”, leave the rest of that box blank.)



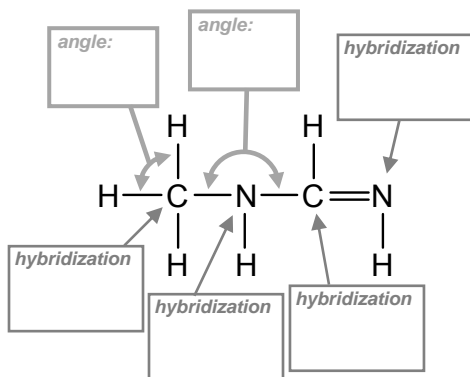
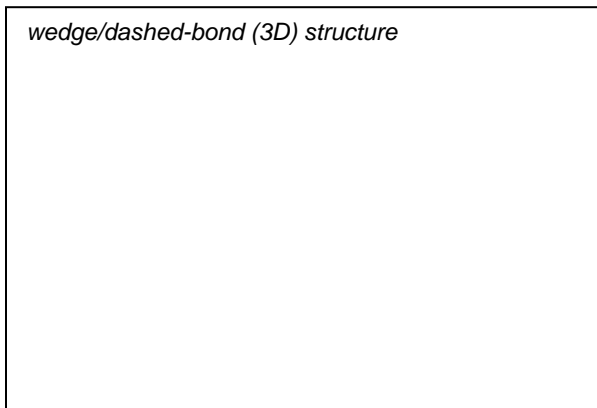
- b. 1,4-Dioxane puckers from planarity in order to avoid angle and torsional strain. In the box on the right, draw the most stable conformation of 1,4-dioxane. In your drawing, **include all H atoms**, but feel free to omit lone pairs.



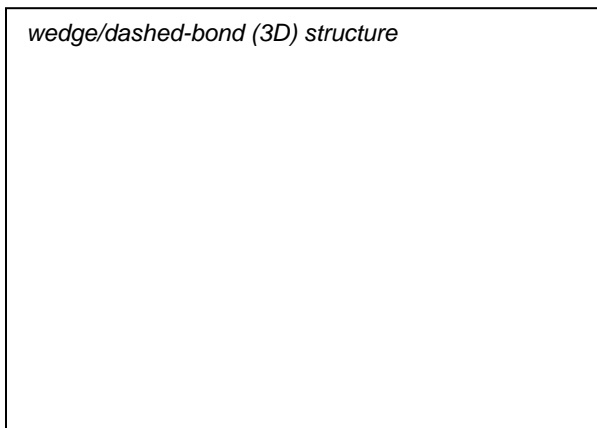
4. (18 pts) For each of the Lewis structures drawn below, in the boxes provided:
- Draw Lewis wedge/dashed-bond structures that illustrate the most stable three-dimensional structure of the molecule. Draw all atoms, but feel free to omit lone pairs.
 - In the boxes provided, write the hybridization state for any atom heavier than hydrogen.
 - In the boxes provided, give any bond angle indicated by curved arrows in the original Lewis structure.



wedge/dashed-bond (3D) structure

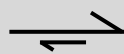
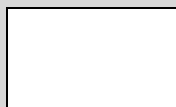
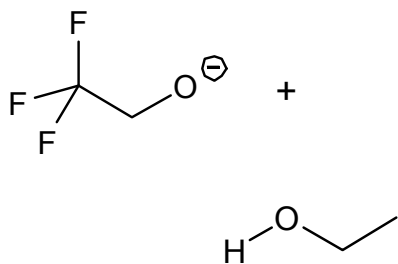


wedge/dashed-bond (3D) structure

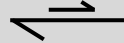


5. (18 pts) For each of the sets of molecules below:

- Using “electron pushing” (with double-barbed arrows), show how the molecules on the left would react in an acid-base reaction to transfer a proton from one to the other.
- In the box on the right, draw the conjugate acid and base products of each reaction.
- In the middle, draw an equilibrium arrow that shows whether you feel the acid-base equilibrium would lie on the left or the right.

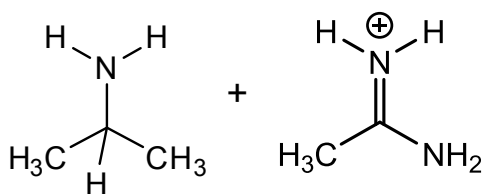
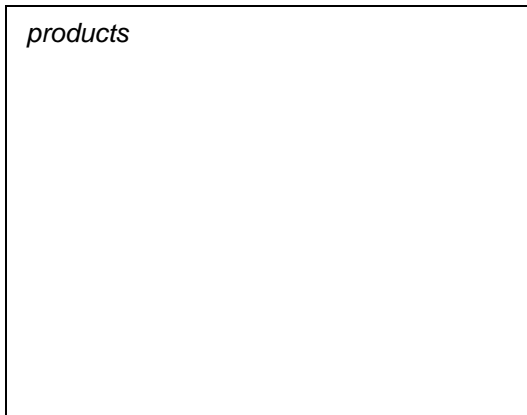


or

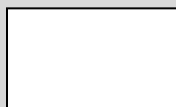
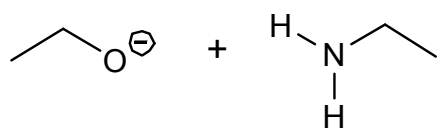
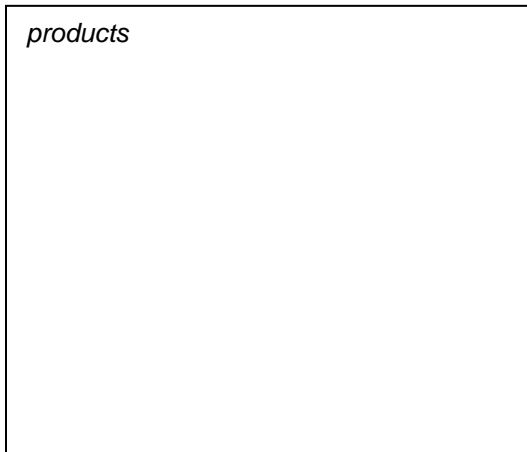


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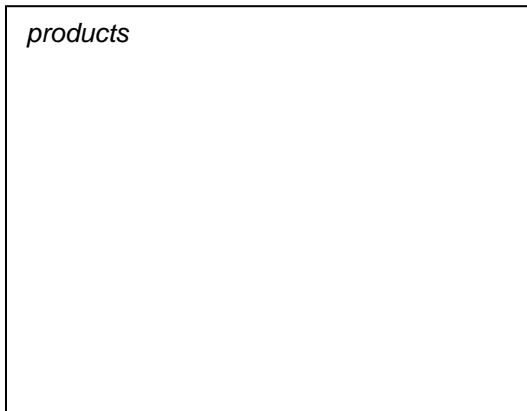
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products

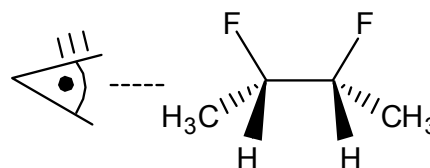


products



6. (20 pts)

- a. For the difluorobutane shown at right, in the boxes below, draw Newman projections that show the most stable, second-most stable, least stable, and second-least stable conformations of the molecule. Draw your projections looking down the central C-C bond, using the perspective I've shown in the drawing.



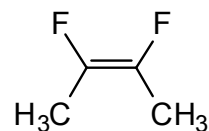
- b. Different conformations can contribute different polarities to the overall, average polarity of a molecule. For each of the two most stable conformations of the molecule above, circle whether the molecule is polar or non-polar when it is in that conformation. If you circle "POLAR", also draw one dipole arrow (\rightarrow) that shows the total dipole moment for that conformer.

<p><i>Newman projection for most stable conformation</i></p>	<p><i>Newman projection for second-most stable conformation</i></p>
<p>POLAR or NON-POLAR ? <i>(circle one)</i></p>	<p>POLAR or NON-POLAR ? <i>(circle one)</i></p>
<p><i>Newman projection for least stable conformation</i></p>	<p><i>Newman projection for second-least stable conformation</i></p>

c. Would you call the 2,3-difluorobutene shown at right a

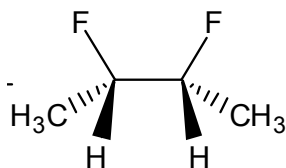
CONFIGURATIONAL ISOMER,

STEREISOISOMER, or NEITHER

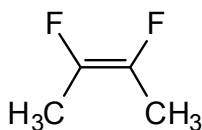


compared to the 2,3-difluorobutane on the previous page? (*Circle one answer.*)

d. Which is more polar:



this difluorobutane?

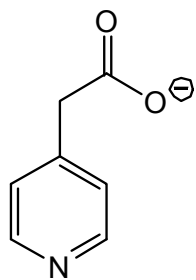


this difluorobutene?

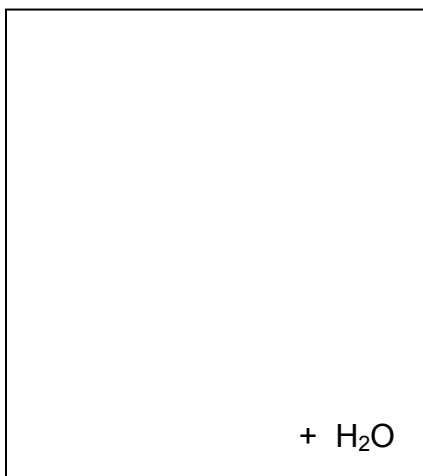
**or are they
equally polar?**

(*Circle one answer.*)

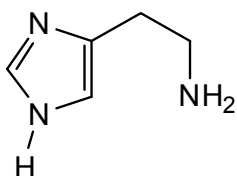
7. (6 pts) Each of the basic molecules below has multiple potential protonation sites. Given the pKa values in the chart on the right, draw the organic product you would expect if each molecule was combined with just one molecule of H₃O⁺.



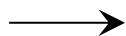
+ H₃O⁺



	<u>pK_a</u>
	10.6
	7.0
	5.3
	4.7



+ H₃O⁺



		1		2		3		4		5		6		7		8		9		10		11		12		13		14		15		16		17		18																																																																																																																																																																																																						
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1	1	H Hydrogen 1.01	2	He Helium 4.00	3	4	Li Lithium 6.94	5	Be Beryllium 9.01	6	7	B Boron 10.81	8	C Carbon 12.01	9	N Nitrogen 14.01	10	O Oxygen 16.00	11	F Fluorine 19.00	12	Ne Neon 20.18	13	Na Sodium 22.99	14	Mg Magnesium 24.31	15	Al Aluminum 26.98	16	Si Silicon 28.09	17	P Phosphorus 30.97	18	S Sulfur 32.07	19	Cl Chlorine 35.45	20	Ar Argon 39.95	21	K Potassium 39.10	22	Ca Calcium 40.08	23	Sc Scandium 44.96	24	Ti Titanium 47.87	25	V Vanadium 50.94	26	Cr Chromium 52.00	27	Mn Manganese 54.94	28	Fe Iron 55.85	29	Ni Nickel 58.69	30	Cu Copper 63.55	31	Zn Zinc 65.39	32	Ga Gallium 69.72	33	Ge Germanium 72.61	34	As Arsenic 74.92	35	Se Selenium 78.96	36	Kr Krypton 83.80	37	Rb Rubidium 85.47	38	Sr Strontium 87.62	39	Y Yttrium 88.91	40	Zr Zirconium 91.22	41	Nb Niobium 92.91	42	Mo Molybdenum 95.94	43	Tc Technetium (98)	44	Ru Ruthenium 101.07	45	Rh Rhodium 102.91	46	Pd Palladium 106.42	47	Ag Silver 107.87	48	Cd Cadmium 112.41	49	In Indium 114.82	50	Sn Tin 118.71	51	Sb Antimony 121.76	52	Te Tellurium 127.60	53	I Iodine 126.90	54	Xe Xenon 131.29	55	Cs Cesium 132.91	56	Ba Barium 137.33	57	La Lanthanum 138.91	58	Ce Cerium 140.12	59	Pr Praseodymium 140.91	60	Nd Neodymium 144.24	61	Pm Promethium (145)	62	Sm Samarium 150.36	63	Eu Europium 151.96	64	Gd Gadolinium 157.25	65	Tb Terbium 158.93	66	Dy Dysprosium 162.50	67	Ho Holmium 164.93	68	Er Erbium 167.26	69	Tm Thulium 168.93	70	Yb Ytterbium 173.04	71	Lu Lutetium 174.97	72	Fr Francium (223)	73	Ra Radium (226)	74	Ac Actinium (227)	75	Rf Rutherfordium (261)	76	Hf Hafnium 178.49	77	Ta Tantalum 180.95	78	W Tungsten 183.84	79	Re Rhenium 186.21	80	Os Osmium 190.23	81	Ir Iridium 192.22	82	Pt Platinum 195.08	83	Au Gold 196.97	84	Hg Mercury 200.59	85	Tl Thallium 204.38	86	Pb Lead 207.2	87	Bi Bismuth 208.98	88	Po Polonium (209)	89	At Astatine (210)	90	Rn Radon (222)	91	Th Thorium 232.04	92	Pa Protactinium 231.04	93	U Uranium 238.03	94	Np Neptunium (237)	95	Pu Plutonium (244)	96	Am Americium (243)	97	Cm Curium (247)	98	Bk Berkelium (247)	99	Cf Californium (251)	100	Fm Fermium (257)	101	Md Mendelevium (258)	102	No Nobelium (259)	103	Lr Lawrencium (262)	104	Db Dubnium (262)	105	Sg Seaborgium (266)	106	Bh Bohrium (264)	107	Hs Hassium (269)	108	Mt Meitnerium (268)	109	Uu Ununennium (289)	110	Uub Ununbium (288)	111	Uut Ununtrium (288)	112	Uuq Ununquadium (289)	113	Uup Ununpentium (288)	114	Uuq Ununhexium (289)	115	Uup Ununseptium (288)	116	Uuq Ununoctium (289)	117	Uup Ununseptium (288)	118	Uuq Ununoctium (289)

Key

11	Na Sodium 22.99
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- Atomic number
- Element symbol
- Element name
- Average atomic mass*

* If this number is in parentheses, then it refers to the atomic mass of the most stable isotope.