EXAMINATION 1 Chemistry 3A

Key

Name:__

SID #:_____

Print first name before second! Use capital letters!

GSI (if you are taking Chem 3AL): _____

Peter Vollhardt February 28, 2017

Please provide the following information if applicable.

Making up an I Grade

If you are, please indicate the semester during which you took previous Chem 3A and the instructor:

Semester

Instructor

Auditor _____

Please write the answer you wish to be graded in the boxed spaces provided.

Do scratch work on the back of the pages. This test should have **16** numbered pages. Check to make sure that you have received a complete exam. A good piece of advice: **Read** carefully over the questions (at least twice); make sure that you understand exactly what is being asked; avoid sloppy structures or phrases. It is better to be pedantic in accuracy now than sorry later! *Good Luck*!

- I. [30 Points] Name or draw, as appropriate, the following molecules according to the IUPAC rules. Indicate stereochemistry where necessary (*cis*, *trans*, *R*, *S*, or dashed/wedged lines).
- a.



cis-1,1,4,5-Tetrachlorocycloheptane

b.

trans-1,3-Bis(1-methylethyl)cyclopentane



c.



(1*R*,3*R*)-1-(2-Bromoethyl)-3-fluorocyclohexane

d.

(R)-4-(1-Chloroethyl)-1,7-diiodoheptane





(4*S*,6*R*)-6-Ethyl-2,4-dimethyl-4-propyldecane

II. [30 Points] Write the best Lewis structure for each of the following molecules. **Remember to assign charges, if any, to atoms!**

TABLE 1-1 Partial Periodic Table								
Period							Halogens	Noble gases
First	H ¹							He ²
Second	Li ^{2,1}	Be ^{2,2}	B ^{2,3}	C ^{2,4}	N ^{2,5}	O ^{2,6}	F ^{2,7}	Ne ^{2,8}
Third	Na ^{2,8,1}	Mg ^{2,8,2}	Al ^{2,8,3}	Si ^{2,8,4}	P ^{2,8,5}	S ^{2,8,6}	Cl ^{2,8,7}	Ar ^{2,8,8}
Fourth	K ^{2,8,8,1}	-					Br ^{2,8,18,7}	Kr ^{2,8,18,8}
Fifth							I ^{2,8,18,18,7}	Xe ^{2,8,18,18,8}
Note: The s	superscripts ind	licate the numb	er of electrons	s in each prin	cipal shell o	f the atom.		

Pauling Electronegativities

(H							He
2.1					7		~0
Li	Ве	В	C	Ň	0	F	Ne
1.0	1.5	2.0	2.5	3.0	3.5	4.0	~0
Na	Mg	AI	Si	P	Ś	CI	Ar
0.9	1.2	1.5	1.8	2.1	2.5	3.0	~0
K	Ca	Ga	Ge	As	Se	Br	Kr
0.8	1.0	1.6	1.8	2.0	2.4	2.8	~0

a.



What is the geometry of NON? Place an "x" mark into the box next to your answer.

Linear I 🗙

Bent

b.



c. The following structure has three octet resonance forms. Circle the best one.



d. The following structure has three octet resonance forms. Circle the best one.



e. Enter the hybridization (sp, sp^2 , sp^3) of the indicated atom in the box provided.



a. Draw the octet structure of the anion shown below. Hint: There will be a Be-Cl double bond.

b. Draw the orbitals on Be and CI that give rise to the double bond and draw the location of the bonding electrons using dots. Clearly label these orbitals (e.g. 1s, 2s, 2p, 3s, 3p, sp, sp^2 , sp^3 , etc.)



c. Draw (separately) orbital energy splitting diagrams for the formation of each part of the double bond: σ and π . Clearly depict the energy levels of the orbitals entering into overlap and label them (Be on the left, Cl on the right), and label the resulting bonding and antibonding molecular orbital levels. Place the relevant electrons into the various levels.





 π Part of the double bond



IV. [30 Points] Consider the rotation about the C1–C2 bond in the cyclopropane derivative shown.



The Newman projections A-F below illustrate the sequential clockwise motion of the backcarbon (C1) in increments of 60°.



a. Indicate, by circling the appropriate letter, the rotamers that contain substituents that are (with respect to each other) anti or gauche, or rotamers that are eclipsed or staggered:



b. Draw a potential energy diagram for this movement. Start by assigning a relative energy to each rotamer on the diagram below, before drawing the interconnecting curve.



c. Two of the staggered rotamers have the same energy. Show which ones, by placing the appropriate letters in the box.

A, C

d. Two of the eclipsed rotamers have the same energy. Show which ones by placing the appropriate letters in the box.

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D, F
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V. [50 Points] A researcher wants to explore the radical "methylthiolation" of alkanes, as exemplified below for reaction of methane.

$$H_{3}C \xrightarrow{\xi} H + H_{3}CS \xrightarrow{\xi} SCH_{3} \xrightarrow{hv} H_{3}C \xrightarrow{\xi} SCH_{3} + H_{3}CS \xrightarrow{\xi} H (1)$$

$$105 \qquad 74 \qquad 77 \qquad 92 \quad \text{kcal mol}^{-1}$$

a. Using the bond dissociation energies provided in equation (1), calculate the ΔH° value for reaction (1). Show your work.

$$\Delta H^{\circ}(1)$$
 : (105 + 74) - (77 + 92) = +10

b. Is the reaction (1) thermodynamically feasible? Circle your answer.

Yes

Answer:



1st Propagation step : $H_3C - H + SCH_3 \longrightarrow CH_3 + HSCH_3$

2nd Propagation step :

 CH_3 + H_3CS SCH₃ ---- CH_3SCH_3 + $\cdot SCH_3$

d. The normalized selectivity tertiary : secondary : primary in methylthiolations with CH_3S is 50:5:1. In the boxes provided, draw the products of single methylthiolation at all positions of *cis*-1,2-dimethylcyclohexane (a–d), as indicated. Note: The bottom half of the molecule is the same as the top; draw only substitutions at the top. **Caution:** Watch out for diastereomers!



cis-1,2-Dimethylcyclohexane

Position a:



Position b:



Position c:





e. Does the radical methylthiolation of *cis*-1,2-dimethylcyclohexane give any optically active products? Circle your answer.

Answer:

Yes

f. Give the ratio of all of the respective products of methylthiolation at positions a–d in the box provided. (In other words, all of the products in the box labeled "Position a" : all of the products in the box labeled "Position c" : all of the products in the bo

Ratio of products a-d:

3:50:10:10

VI. [15 Points]

a. Using the values in the Table below and the cyclohexane stencils provided, show the structures of the most stable conformers of **A**, **B**, and **C**. For each, calculate the change in free energy on "ring flip" to the less stable conformer. Make sure to "cap off" all axial and equatorial bonds with substituents and H atoms. Enter your calculated energy of "ring flip" in the box provided.



A: cis-1-Bromo-2-fluorocyclohexane





B: cis-1-Methyl-4-(1,1-dimethylethyl)cyclohexane





C: trans-1-lodo-3-methylcyclohexane



VII. [65 Points]

1. Mark the box next to your choice of an answer with an X.

The C-H bond strength deceases in the order CH4, primary, secondary, tertiary, because

steric hindrance decreases

the product radicals increase in stability

	_

the product radicals decrease in stability



hybridization changes from sp^2 to sp^3

2. In the following pairs of compounds, circle which one is more acidic.



3. Consider the following chlorination.



a. What is the absolute configuration of A (R or S)? Circle the correct answer.

r S

b. The products are (circle the correct answer):



In equal amounts (In unequal amounts)

4. Does the arrow pushing in the following molecules **A**–**D** lead to an acceptable resonance form? Circle your answers.



5. The following potential-energy diagram shows the addition reaction of A to B to give E via intermediates C and D.



a. Mark with an "X" the rate-determining step of the transformation of A+B to D.

b. If you were to follow the progress of the transformation, would you observe sizeable amounts of C and/or D? Place an "X" mark in the appropriate box.



c. The reaction of A with B follows second order kinetics. If you double the concentration of A and simultaneously double the concentration of B what would be the effect on the rate of the reaction? Place an "X" mark in the appropriate box.





J The End J