EXAMINATION 1

Name:

Chemistry 3A Kim Lavoie Peter Vollhardt February 27, 2003

[Print first name before second! Use capital letters!]

Please check the name of your TA and corresponding section number. Complete the remaining information if applicable.

110	Olga Fedin	270	Staffan Westerberg
120	Sean Wiedemann	310	Cindy Chang
130	Kristie Koski	320	Raja Sivamani
140	Stephanie Chan	360	Ravi Chandrasekaran
160	Zack Fresco	370	Douglas Mitchell
170	Andrew Chi	410	Amish Patel
180	Hany Eitouni	420	Matt Banghart
190	Jenn Barbarow	460	Nicholas Ohler
210	Nicholas Agard	470	Greg Watkins
220	Jimmy Blair	510	Ben Huang
230	Scott Laughlin	520	Tanya Leavy
240	Carl Mieczkowski	560	Josh Goldberger
260	Jessica Defreese	570	Josh Gilmore

Making up an I Grade

(If you are, please indicate the semester during which you took previous Chem 3A _____).

Please write the answer you wish to be graded in the spaces provided. *Do scratch work on the back of the pages.* This test should have 15 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! <u>Good Luck</u>!

DO NOT WRITE IN THIS SPACE		
	I.	 (20)
	II.	 (20)
	III.	 (40)
	IV.	 (30)
	V.	 (40)
	VI.	 (30)
	VII.	 (20)
	Total:	 (200)

I. [20 Points] Name or draw, as appropriate, the following molecules according to the IUPAC rules. Indicate stereochemistry where necessary (*cis, trans, R, S,* or *meso*). Indicate with a circle whether the molecule is chiral or achiral.



chiral	achiral





c. trans-1,2-Diethylcyclohexane





II. [20 Points] Write the best Lewis resonance structure for each of the following molecules. Remember to assign charges!

a.	F F F
b.	<pre>-</pre>
C.	[в о] ⁻
d.	[N H С Ц Н N Н]

TABLE	1-1 Par	rtial Period	ic 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	uperscripts inc	licate the numb	er of electrons	s in each prin	cipal shell o	f the atom.		

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- III. [40 Points] Ozone, O₃, exists in the acyclic form, but has a cyclic isomer.
 - a. Draw both in their best Lewis octet versions.



b. Show the orbital overlap picture for one of the O–O bonds in cyclic ozone.
 Label clearly the overlapping orbitals (e.g. s, p, sp³, etc.). Hint: Recall cyclopropane!



c. Show the orbital splitting associated with the O–O σ bond above in an energy level diagram. Label each level clearly [e.g. *s*, *p*, *sp*³, bonding molecular orbital (MO), etc.].



d. Place an "X" in the box for the two most plausible reasons why ozone is acyclic, but cyclopropane is not. Note: only two marks are allowed!





the O–O bond is much weaker than the C–C bond



acyclic O3 is stabilized by resonance



the entropy for ring opening in cyclic O_3 is much more positive than in cyclopropane



cyclic O₃ has no eclipsing hydrogens



the barrier to ring closure of ozone is too high

IV. [30 Points] Consider the rotation about the C1-C2 bond in 1-bromo-2-methylpropane, illustrated by the series of Newman projections A-F generated by the sequential clockwise motion of the back-carbon (C2) in increments of 60°.



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

anti:	A	gauche:	A	eclipsed:	A	staggered:	A
	В		В		В		В
	С		С		С		С
	D		D		D		D
	Е		Е		Е		E
	F		F		F		F

b. Draw a potential energy diagram for this movement. Start by assigning a relative energy to each rotamer on the diagram 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.



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



V. [40 Points] We have learned that ethane undergoes radical bromination to bromoethane, by attack of Br[•] on a methyl hydrogen.

$$CH_3 - CH_3 + Br_2 \rightarrow CH_3 - CH_2 - Br + HBr$$
 (1)

In principle, however, the reaction could have taken a different path, namely attack on carbon to give bromomethane.

$$CH_3 - CH_3 + Br_2 \rightarrow 2 CH_3 - Br$$
 (2)

a. Using the Tables provided on p. 9, calculate the ΔH° values for reactions (1) and (2).

 $\Delta H^{\circ}(1)$:

 ΔH° (2) :

b. Is the reaction (2) thermodynamically feasible? Circle the right answer.

Answer: Yes No

TABLE 3-1	Bond-Diss	ociation En	rergies of	Uarious A	-B Bonds			T981 F 3-4
				B in A-I	_			DH ^o Values for the
A in A-B	Н-	-F	-CI	-Br	I-	HO-	-NH2	Elemental Halogens
H—	104	136	103	87	71	119	108	
CH ₃ -	105	110	85	70	57	93	84	DH°
CH,CH,-	101	111	84	70	56	94	85	Halamn (Faal mal-1)
CH,CH,CH,-	101	110	85	70	56	92	84	IIIalogell (Kcal III01 -)
(CH ₁), CH-	98.5	III	84	11	56	96	86	
(CH ₃) ₃ C-	96.5	110	85	71	55	96	85	F ₂ 38
Note: These numbers are	e being revise	ed continually	because of i	improved meth	nods for their	r measureme	nt. Some of	CI 58
the values given here me	ay be in (smal	II) error.						V12 J0
								Br, 46
								1 36
								27
TABLE 3-2	Bond-D	issociati	ion Ener	rgies for	Some A	lkanes		
		oHo					DH°	
Compound	(ł)	cal mol ⁻¹	0 (1	ompound			(kcal mol ⁻¹)	
CH ₃ ← H		105	0	CH ₃ +CH ₃			90	
C ₂ H ₅ +H		101	0	² H ₅ +CH ₃			89	
C_3H_7 $+H$		101	0	² ,H ₅ +C ₂ H	2		88	
(CH ₃) ₂ CHCH ₂ +H	F	101	9	CH ₃) ₂ CH ₄	-CH ₃		88	
(CH ₃) ₂ CH→H		98.5	9	CH ₃) ₃ C+(CH ₃		87	
(CH ₃) ₃ C+H		96.5	9	CH ₃) ₂ CH+	-CH(CH	3)2	85.5	
			9)	CH ₃) ₃ C+(C(CH ₃) ₃		78.5	
Note: See footnote	for Table 3-	-1.						

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c. Formulate the first propagation step for reactions (1) and (2).

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Reaction (1) — 1^{st} Propagation Step :
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Reaction (2) — 1st Propagation Step :

- d. Calculate the ΔH° values for the two reactions in c.
- (1) ΔH° (1st propagation step) :

(2) ΔH° (1st propagation step) :

e. Considering the results of d., can you think of a reason why ethane reacts with bromine to give bromoethane and not bromomethane?

VI. [30 Points] Compound A undergoes radical bromination to give B and C, among other products.



Substituent	ΔG° (kcal mol ⁻¹)	Substituent	ΔG° (kcal mol ⁻¹)
Н	0	F	0.25
CH ₃	1.70	Cl	0.52
CH ₃ CH ₂	1.75	Br	0.55
(CH ₃) ₂ CH	2.20	I	0.46
(CH ₃) ₃ C	≈ 5	НО	0.94
о НО-С	1.41	CH ₃ O	0.75
O ∥ CH₃O−C	1.29	H ₂ N	1.4

a. Given the values in the Table and using the cyclohexane stencils provided, show the structures of the most stable conformers of A, B, and C. For each, calculate the free energy of "ring flip" to the less stable conformer.



b. The molecule **A** has primary, secondary, and tertiary hydrogens. Considering the relative reactivity in brominations of C-H:

primary : secondary : tertiary = 1 : 80 : 1700,

what would you expect the ratio of all primary to all secondary to all tertiary substitution products to be?

Show your work.

- VII. [20 Points] Place an X mark in the box designating the most accurate statement.
 - a. The ΔH° of an organic reaction:



reflects entropy changes



is highly negative for fast transformations



can be estimated by subtracting the sum of the \textit{DH}° values of the bonds formed from those broken



is the symbol for the rate in the Arrhenius equation

b. When considering the following potential energy diagram:





compound A will convert to B faster than it will to C.

C is the thermodynamically most stable component of the mixture and will form at the greatest rate from **A** or **B**.



B will convert to C faster than A will.



none of the above

c. The compounds *cis*- and *trans*-1,3-dimethylcyclohexane are:

identical
stereoisomers
interconverted by ring flip
rotamers

d. The bromination of 2S,3S-dibromobutane **A** gives 2,2,3-tribromobutane **B**:



	I
	I
	I
	I
	I



as a racemate



as a meso compound



as an achiral molecule



* The End *