EXAMINATION 2 Chemistry 3A

Name:___

Print first name before second! Use capital letters!

SID #:_____

Make sure the number is correct!

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

Peter Vollhardt April 9, 2019

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 **14** 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!

Do not remove this (or any other) page from the exam. It will compromise the consonance of scanned copies in Gradescope.

You probably will not need them for the exam, but here they are for added comfort:

21					cuone	Jacivicy	value	s of the	eleme	ents (P	auling	scale))				He
LiI	Be											В	С	N	0	F	Ne
1.0 1	1.5											2.0	2.5	3.0	3.5	4.0	
Na I	Mg	AL SI P S CL Ar							Ar								
0.9 1	1.2											1.5	1.8	2.1	2.5	3.0	
K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
0.8	1.0	1.3	1.5	1.6	1.6	1.5	1.8	1.8	1.8	1.9	1.6	1.6	1.8	2.0	2.4	2.8	3.0
Rb	Sr	Υ	Zr	Nb	Мо	Tc	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	1	Xe
0.8 1	1.0	1.2	1.4	1.6	1.8	1.9	2.2	2.2	2.2	1.9	1.7	1.7	1.8	1.9	2.1	2.5	2.6
Cs I	Ва	La	Hf	Та	w	Re	Os	lr	Pt	Au	Hg	Ti	Pb	Bi	Po	At	Rn
0.7 0	0.9	1.1	1.3	1.5	1.7	1.9	2.2	2.2	2.2	2.4	1.9	1.8	1.8	1.9	2.0	2.2	2.4
Fr	Ra	Ac															
0.7 0	0.7	1.1															

Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Yb	Lu
1.1	1.1	1.1	1.1	1.1	1.1	1.1	1.1	1.1	1.1	1.1	1.1	1.1	1.2
Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr
1.3	1.5	1.7	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	

Table 2.2

Relative Acidities of Common Compounds (25°C)

Acid	K _a	pK _a
Hydrogen iodide, HI (strongest acid) Hydrogen bromide, HBr Hydrogen chloride, HCl Sulfuric acid, H ₂ SO ₄ Hydronium ion, H ₃ O ⁺ Nitric acid, HNO ₃ Methanesulfonic acid, CH ₃ SO ₃ H Hydrogen fluoride, HF Acetic acid, CH ₃ COOH Hydrogen cyanide, HCN Ammonium ion, NH ₄ ⁺ Methanethiol, CH ₃ SH Methanol, CH ₃ OH Water, H ₂ O Ethyne, HC=CH Ammonia, NH ₃ Ethene, H ₂ C=CH ₂ Methane, CH ₄ (weakest acid) <i>Note:</i> $K_a = [H_3O^+][A^-]/[HA] \mod L^{-1}$. ^a First dissociation equilibrium	$\begin{array}{c} \sim\!\! 1.0\times10^{10}\\ \sim\!\! 1.0\times10^9\\ \sim\!\! 1.0\times10^8\\ \sim\!\! 1.0\times10^3\\ 50\\ 25\\ 16\\ 6.3\times10^{-4}\\ 2.0\times10^{-5}\\ 6.3\times10^{-10}\\ 5.7\times10^{-10}\\ 1.0\times10^{-10}\\ 3.2\times10^{-16}\\ 2.0\times10^{-16}\\ 2.0\times10^{-16}\\ \sim\!\! 1.0\times10^{-35}\\ 1.0\times10^{-35}\\ \sim\!\! 1.0\times10^{-50}\\ \end{array}$	$\begin{array}{c} -10.0\\ -9.0\\ -8.0\\ -3.0^{a}\\ -1.7\\ -1.4\\ -1.2\\ 3.2\\ 4.7\\ 9.2\\ 9.3\\ 10.0\\ 15.5\\ 15.7\\ -25\\ 35\\ -44\\ -50\end{array}$

I. [30 Points] Name or draw, as appropriate, the following molecules according to the IUPAC rules. Indicate stereochemistry where necessary.



Br O



II. [80 Points] Add the missing starting materials, reagents, or products (aqueous work-up is assumed where necessary). **Caution:** Do not forget to consider **stereochemistry**!



For the following questions, circle your choice of an answer:

ls	the pro	duct chi	iral	?		Yes	No	
_								

Is the product optically active? Yes No



For the following questions, circle your choice of an answer:

Is the product chiral?	Yes	No
Is the product optically active?	Yes	No



For the following questions, circle your choice of an answer:

Is the product chiral?	Yes	No	
Is the product optically active?	Yes	No	

d.









Complete the stencil in the box by adding the missing substituents at C1, C2, and C3

For the following question, circle your choice of an answer:

Is the product chiral?	Yes	No
------------------------	-----	----

Is the product optically active? Yes No

III. [40 Points] The following reactions proceed (predominantly) by S_N2, S_N1, E2, or E1 pathways, respectively. Give the major organic product in each case and answer the questions by *circling* the most applicable statement.



Changing the solvent to CH₃CH₂OH has one of the following effects:

 $E2 / S_N1 \text{ increases} \qquad E1 / S_N1 \text{ increases} \qquad Rate \text{ increases} \qquad Rate \text{ decreases}$



Omitting Nal from the starting mixture has one of the following effects:

Starting material stays unchangedStarting material equilibrates with its diastereomerE2 / E1 ratio increases $S_N2 / E1$ ratio increases



Which of the following statements is correct?

Doubling the concentration of starting iodide will double the rate of its disappearance Doubling the concentration of starting iodide will quadruple the rate of its disappearance Doubling the concentration of starting iodide will not change the rate of its disappearance

IV. [40] Points]

a. For the following reaction, provide a detailed mechanism (i.e., write a scheme with structures, arrow pushing, etc.) Do **not** *add* any reagents! This is not a synthesis!



Work from left to right in the following spaces. There is more space than you will need.

b. For the following reaction, provide a detailed mechanism (i.e., write a scheme with structures, arrow pushing, etc.) Do **not** *add* any reagents! This is not a synthesis!



Work from left to right in the following spaces. There is more space than you will need. Do not worry about stereochemistry.

V. [40 Points]

a. Provide a viable conversion of the starting material below as the only carbon source to the product. It will help you if you execute a retrosynthesis on the back of the preceding page (on your left). This problem should look familiar.

HO H₃C−CH₃ ------

Work from left to right in the following spaces. There is much space than you will need.

b. Provide a viable conversion of the starting material below to the product. You may use any additional organic compounds containing four carbons or less and any inorganic compounds in your scheme. It will help you if you execute a retrosynthesis on the back of the preceding page (on your left).

റ HO HO –Н 0 Do not worry about

stereochemistry

Work from left to right in the following spaces. There is more space than you will need.

VI. [20 Points]

a. We learned that the acidity if alcohols *decreases* when progressing from primary to secondary to tertiary, as shown in Table 8.2 below, because of increasing steric hindrance to solvation of the resulting conjugate base.

Table 8.2	pK_a Values of Alcohols in Wate	r
Compound	pK _a	
НОН	15.7	
CH ₃ OH	15.5	
CH ₃ CH ₂ OH	15.9	
(CH ₃) ₂ CHOH	17.1	
(CH ₃) ₃ COH	18	

In contrast, Table 8.3 in your book shows that the acidities of alkyloxonium ions *increase* when progressing from primary to secondary to tertiary.

Table 8.3	p <i>K</i> _a Values of Four Protonated Alcohols
Compound	pK _a
$CH_3 \overset{+}{O}H_2$	-2.2
CH ₃ CH ₂ ⁺ OH	l ₂ -2.4
$(CH_3)_2CHC$	-3.2
$(CH_3)_3 COH$	H ₂ -3.8

Place an *X mark* in the box preceding the most reasonable explanation for this finding.

Along the series (primary to secondary to tertiary), hyperconjugation increases.



Along the series (primary to secondary to tertiary), the alcohols resulting from proton dissociation are increasingly better solvated.



Along the series (primary to secondary to tertiary), the oxonium ions suffer increasing steric hindrance to solvation.



Along the series (primary to secondary to tertiary), the oxonium ions suffer decreasing inductive stabilization.

b. The following four problems list pairs of statements. For each, place an *X mark* in the box preceding the correct one.

1.	Electronegativity increases to the right and up the periodic table.
	Electronegativity increases to the right and down the periodic table.
2.	Nucleophilicity in protic solvents increases down the periodic table.
	Nucleophilicity in aprotic solvents increases down the periodic table.
3.	Leaving group ability increases to the right and down the periodic table.
	Leaving group ability increases to the right and up the periodic table.
4.	The S_N2 reaction of oxacyclopropanes is possible because the ring carbon is relatively unhindered.
	The S_N2 reaction of oxacyclopropanes is possible because ring strain is released.



* The End *