Exam 3 Chm 205 (Dr Mattson) 6 April 2016

Academic Integrity Pledge: In keeping with Creighton University's ideals and with the Academic Integrity Code, I pledge that this work is my own and that I have neither given nor received inappropriate assistance in preparing it.

Name:

Chemistry Student Number:

Signature:

Instructions: Show all work whenever a calculation box is provided! Write legibly. Include units whenever appropriate. You will receive credit for how you worked each problem as well as for the correct answer. If you need more space, you may use the back of the data sheet provided — Write: "See data sheet" in the answer box – then write your name on the data sheet. On your desk you are allowed only pencils (but no pencil pouch), an eraser, and a non-programmable calculator without a slipcover. Backpacks, bags, and purse-like items must be stored in the rear section of the room. Cell phones must be silent and placed in your backpack/bag/purse - not in your pocket.

Note: All of these questions refer to aqueous solutions.

1. Phenol or carbolic acid, HC₆H₅O, shown here, is an important industrial chemical used in the production of plastics, cosmetics, sunscreens, and pharmaceuticals, aspirin, and antiseptics. Over 7 x 10⁹ kg phenol is produced annually. Phenol has a $pK_a = 9.90$.



1a. (2 pts) What is the numerical value of Ka? Sig fig problem



1b. (3 pts) What is the formula of the conjugate base of phenol and what is the numerical value of pK_h?

cwb:	
pK _b =	

1c. (9 pts) Which equilibrium expression A, B, C, D, E, F describes each equilibrium below? Fill in the arrows, either ← → or ← → for each. Circle A B C D E or F.

A. $K = K_a/K_w$ **B.** $K = K_w$ **C.** $K = 1/K_w$

D. K = K_a **E.** K = K_w/K_a **F.** K = $1/K_a$

HC ₆ H ₅ O + OH⁻		H ₂ O + C ₆ H ₅ O ⁻	Circle one: A B C D E F
HC ₆ H ₅ O + H ₂ O		H ₃ O ⁺ + C ₆ H ₅ O ⁻	Circle one: A B C D E F
H ₃ O ⁺ + OH ⁻		2 H ₂ O	Circle one: A B C D E F
C ₆ H ₅ O ⁻ + H ₂ O		OH⁻ + HC ₆ H ₅ O	Circle one: A B C D E F
C ₆ H ₅ O - + H ₃ O+		H ₂ O + HC ₆ H ₅ O	Circle one: ABCDEF

1d. (5 pts) Which would form a buffer? More than one!

\square 0.10 mol HC ₆ H ₅ O + 0.10	$mol C_6H_5O^- in 1$	L H ₂ O
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 \Box 0.10 mol H₃O⁺ + 0.10 mol C₆H₅O⁻ in 500 mL H₂O

 \Box 0.10 mol HC₆H₅O + 0.040 mol OH⁻ in 250 mL H₂O

 \square 0.10 mol HC₆H₅O + 0.18 mol OH⁻ in 300 mL H₂O

 \Box 5.0 g HC₆H₅O + 4.0 g NaC₆H₅O in 400 mL H₂O

1e. (4 pts) What is the pH of a solution prepared by dissolving 0.40 mol HC₆H₅O and 0.35 mol C₆H₅O⁻ in 250 mL water?

Answer:

1f. (4 pts) What is the pH of the solution in Question 1e if 0.040 mol potassium hydroxide were added?

ore to mer petaceram nyarextae were added.		
	Answer:	

1f. (1 pt) What will happen to the pH if another 50 mL water is added to the solution in the previous problem?

It will: 🔲 Increase 🔲 Decrease 🚨 Stay the same

1g. (1 pt) The solution in Question 1e has a larger buffer capacity towards the addition of...

Circle one: ☐ Strong acid ☐ Strong base ☐ Both

2. (4 pts) Methanoic acid, HCO_2H , has a $K_a = 1.7 \times 10^{-4}$ and ethanoic acid, $HC_2H_3O_2$, has a $K_a = 1.8 \times 10^{-5}$. What is the equilibrium constant, K_c for:

 $HCO_2H + C_2H_3O_2^{-1} + HC_2H_3O_2 + K_c = ?$

r	
Show work for credit!	
	Answer:
	Allawei

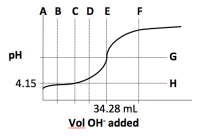
3a. (3 pts) Suppose 40.00 mL 0.1050 M HCl was titrated with NaOH and it took 32.78 mL to reach a nhenolohthalein endogint What is [NaOH]?

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	Answer:

3b. (4 pts) What is the pH of the solution after 30.00 mL NaOH(aq) has been added? Sig fig problem.

Answer:

 An unknown weak acid, HA, was dissolved in 50.00 mL water and titrated with 0.1106 M NaOH as shown in the figure.



4a (3 pts) How many moles of OH⁻ did it take to reach the

equivalence point? Sig fig problem.

Answer with units:	

4b. (2 pts) The intersection of what two lines (for example: B & G) would you use in order to determine the pK_a for the weak acid?

4c. (3 pts) Which lines designate volumes of OH⁻ added where we have a buffer solution? Circle all that apply.

Line A Line B Line C Line D Line E

4d. (4 pts) Suppose Line D represented a volume of 23.00 mL. What is the pH of the solution at this point?

	Answer:
	· · ·

4e. (4 pts) What is the pH at the equivalence point?

Answer:

5a. (12 pts) Urea has formula CON_2H_4 and a melting point of 134 ^{o}C . Its solubility in water increases with temperature. Like most organic compounds, urea can be combusted, forming $CO_2(g)$, $H_2O(g)$ and $N_2(g)$. Predict the sign for ΔH , ΔS and ΔG for each of the following processes. Write in each box: +, 0 or -.

Process involving urea:	ΔН	ΔS	ΔG
Combustion of CON ₂ H ₄			
Breaking covalent bonds at 298 K			
Melting of solid urea at 134 °C			
Dissolving urea in water			

Score	
Δ > 90° B+ > 85° B > 80° C+ > 75° C > 70° D > 60	

6a. (5 pts) Calculate ΔS^o for the reaction: See data sheet. $N_2H_4(g) + 2 N_2O(g) \longrightarrow 3 N_2(g) + 2 H_2O(g)$ Answer with units:

Is this reaction entropy favored? Yes No

6b. (5 pts) Calculate $\triangle G^o$ for the reaction: See data sheet. $N_2H_4(g) + 2 N_2O(g) \longrightarrow 3 N_2(g) + 2 H_2O(g)$ Answer with units:

Is this reaction spontaneous? Yes No

7. (8 pts) Circle + or - for each of these

ΔG is spontaneous	if ∆H is	and ∆S is
at all temperatures.	+ -	+ -
only at high temperatures.	+ -	+ -
only at low temperatures.	+ -	+ -
never.	+ -	+ -

8a. (5 pts) Given that ΔH^0 = +57.0 kJ, ΔS^0 = +176 J/K and ΔG^0 = +4.6 kJ for the reaction below, calculate ΔG at 298 K when P_{N2O4} = 5.0 atm and P_{NO2} = 0.10 atm.

$N_2O_4(g) \rightarrow 2 NO_2(g)$		
Answer with units:		
Is the forward reaction spontaneous? Yes No		

8b (5 pts) Calculate the equilibrium constant at 298 K for the reaction.

the reaction.		
	A	
	Answer:	

8c. (4 pts) Calculate the crossover temperature (when ΔG = 0 kJ) for this reaction.

Answ	ar.	

Data sheet

Useful Formulas:

$$\Delta G = \Delta G^{\circ} + R T InQ$$

 $\Delta G^{\circ} = -R T InK$

 $R = 8.314 \text{ J mol}^{-1} \text{ K}^{-1}$

 $R = 0.0821 L atm mol^{-1} K^{-1}$

	ΔH_f^o	ΔG_f^o	Sº
	kJ/mol	kJ/mol	J/mol K
N ₂ H ₄ (g)	95.4	159.3	238.4
N ₂ O(g)	82.0	104.2	219.7
N ₂ (g)	0	0	191.5
H ₂ O(g)	-241.8	-228.6	188.7

Answers

1a.
$$K_a = 1.3 \times 10^{-10}$$

1b. Conjugate base $HC_6H_5O^-$ and $pK_b = 4.10$

1c.

HC ₆ H ₅ O + OH⁻	_	H ₂ O + C ₆ H ₅ O ⁻	Circle one:
HC ₆ H ₅ O + H ₂ O	←→	H ₃ O ⁺ + C ₆ H ₅ O ⁻	Circle one:
H ₃ O ⁺ + OH ⁻		2 H ₂ O	Circle one:
C ₆ H ₅ O - + H ₂ O	₩	OH⁻ + HC ₆ H ₅ O	Circle one:
C ₆ H ₅ O - + H ₃ O+		H ₂ O + HC ₆ H ₅ O	Circle one:

1d.

X 0.10 mol HC $_6$ H $_5$ O + 0.10 mol C $_6$ H $_5$ O $^{\scriptscriptstyle -}$ in 1 L H $_2$ O

 \square 0.10 mol H₃O⁺ + 0.10 mol C₆H₅O⁻ in 500 mL H₂O

X 0.10 mol $HC_6H_5O + 0.040$ mol OH^- in 250 mL H_2O

 \square 0.10 mol HC₆H₅O + 0.18 mol OH $^{-}$ in 300 mL H₂O

X 5.0 g $HC_6H_5O + 4.0$ g NaC_6H_5O in 400 mL H_2O

1e. 9.84

1f. 9.93

1f. stay the same

1g. strong base

 $2. K_{c} = 9.5$

3a. 0.1281 M

3b. 2.29

4a. 3.791 x 10⁻³

4b. C and H

4c. Lines B, C, and D

4d. 4.46

4e. 8.40

5a.

Process involving urea:	ΔН	ΔS	ΔG
Combustion of CON ₂ H ₄	-	+	-
Breaking covalent bonds at 298 K	+	+	+
Melting of solid urea at 134 °C	+	+	0
Dissolving urea in water	+	+	-

6a. $\Delta S^0 = 274.1 \text{ J/mol rxn K}$; favored

6b. ΔG° = -824.9 kJ/mol rxn; spontaneous

7.

∆G is spontaneous	if ∆H is	and ∆S is
at all temperatures.	-	+
only at high temperatures.	+	+
only at low temperatures.	-	-
never.	+	-

8a. $\Delta G = -10.8 \text{ kJ/mol rxn}$; Yes, spontaneous

8b $K_p = 0.15$

8c. 324 K