الدورة الإستثنائية للعام ٢٠٠٨	امتحانات الشهادة الثانوية العامة الفرع : علوم عامة	وزارة التربية والتعليم العالي المديرية العامة للتربية دائرة الامتحانات
الاسم: الرقم:	مسابقة في مادة الكيمياء المدة ساعتان	

This Exam Includes Three Exercises. It Is Inscribed on 3 Pages Numbered From 1 to 3. The Use of a Non-programmable Calculator Is Allowed. Answer the three Following Exercises:

First Exercise (6 points) Identification of certain Organic Compounds

The mild oxidation of an alcohol (A) leads to the formation of a monocarboxylic acid (B) with a saturated open carbon chain.

The aim of this exercise is to identify (A) and (B) and to study their esterification reaction.

Given:

- M(H) = 1 g.mol⁻¹; M(C) = 12 g.mol⁻¹; M(O) = 16 g.mol⁻¹
- The yield of the esterification of an equimolar mixture of carboxylic acid and a primary alcohol is 67 %.

1- Identification of (A) and (B)

The percentage by mass of oxygen in the acid (B) is 36.36 %

- 1.1-Show that the molecular formula of (B) is $C_4H_8O_2$. Deduce the molecular formula of (A).
- 1.2-Knowing that the carbon chain of (A) is branched, identify (A) and (B),

2- Esterification of Alcohol (A)

A mixture of 1 mol of (A) and 2 mol of (B) is heated in the presence of few drops of concentrated sulfuric acid. After a certain time, a homogeneous equilibrium is established with a constant $K_C = 4$.

- 2.1- Write the equation of this reaction. Give the systematic name of the organic compound (C) formed.
- 2.2- Determine the number of moles of (C) formed at equilibrium. Deduce the yield of this esterification.
- 2.3- This yield exceeds 67 %. Explain, if each one of the three following proposals justifies this fact:
 - 2.3.1- Heating the reacting medium.
 - 2.3.2- Presence of few drops of sulfuric acid in the reacting medium.
 - 2.3.3- Using an initial mixture of reactants which is not equimolar.
- 2.4-In order to make the formation reaction of (C) complete, (B) is replaced by one of the two compounds (B[']) and (B"). The equations of the two reactions are the following:

$$\begin{array}{ccc} A & +B' & \rightarrow & B & + & C \\ A & + & B'' & \rightarrow & C & + & HCl \end{array} \tag{1}$$

Identify (B) and (B").

Second Exercise (7 points) Decomposition Reaction of N₂O₅

Nitrogen pentoxide decomposes completely when heated, in the gaseous phase, according to the following equation: $2 N_2 O_{5(g)} \rightarrow 4 NO_{2(g)} + O_{2(g)}$.

Given:

- All gases are considered ideal.

- The ideal gas constant is R = 0.082 atm.L.mol⁻¹.K⁻¹.

1- Decomposition of N₂O₅ in a Container of Constant Volume

 $n_0 = 1$ mol of N_2O_5 is introduced into a container of constant volume V = 10 L, maintained at a temperature T = 413 K.

1.1- Calculate, p_0 , the initial pressure in the container.

- 1.2- Show that the total pressure p in the container increases when the reaction of decomposition of N_2O_5 advances.
- 1.3- Calculate the maximum value of p.
- 1.4- Show that the concentration of N_2O_5 , at instant t, is given by the following relation:

$$[N_2O_5]_t = \frac{5p_0 - 2p}{3RT}$$

2- Kinetics of the Decomposition Reaction of N2O5

The above reaction is slow, to study its kinetics, a certain quantity of N_2O_5 is introduced, at constant temperature, into an evacuated container which is connected to a pressure gauge. The measurement of the pressure in the container makes it possible to calculate the concentration of $[N_2O_5]_t$ at instant t.

The results are given in the following table:

t(s)	0	200	400	600	800	1000	1200	1400	1600	1800	2000
$[N_2O_5]_t (mol.L^{-1})$	1	0.88	0.78	0.69	0.61	0.54	0.48	0.43	0.38	0.34	0.30

2.1- Plot, on the graph paper, the curve $[N_2O_5]_t = f(t)$. Take the following scale: abscissa 1 cm for 200 s and ordinate 1 cm for 0.1 mol.L⁻¹.

- 2.2- Determine graphically the half-life of the reaction.
- 2.3- Determine the rate of disappearance of N_2O_5 when its concentration becomes 0.8 mol.L⁻¹. Deduce the rate of the reaction at the same instant.

Third Exercise (7 points) Acid-base Reactions

Given:

Conjugate acid/base pair	CH ₃ NH ⁺ ₃ /	NH_4^+/NH_3	HF/ F	HCOOH/ HCOO ⁻
	CH_3NH_2			
pK _a	10.7	9.2	3.2	3.8

- The ionic product of water: $K_w = 1.0 \times 10^{-14}$

- Sodium methanoate HCOONa and ammonium chloride NH₄Cl are ionic compounds highly soluble in water.

Four beakers contain each an aqueous solution of one of the chemical compounds given in the table below. The solutions have all the same molar concentration C_0 .

Number of the beaker	Chemical compound	pН
1	Sodium methanoate	pH_1
2	Ammonium chloride	pH ₂
3	Methylamine CH ₃ NH ₂	pH ₃
4	Hydrogen fluoride HF	pH ₄

1- <u>pH of the Aqueous Solutions</u>

- 1.1- Classify the pH of these four solutions by ascending order.
- 1.2- The pH of the solution in beaker N° 4 has the value of 2.65.
- 1.2.1- Write the equation of the reaction between HF and water.
- 1.2.2- Show that the molar concentration C_0 is equal to 1.0×10^{-2} mol.L⁻¹.
- 1.3- Distilled water is added to a volume $V_0 = 10$ mL of the solution of beaker N° 3 until we obtain a solution S of volume V = 100 mL. The pH of the solution of beaker N° 3 and that of solution S are measured. The results are given in the following table:

Solution	Beaker Nº 3	S
$C (mol.L^{-1})$	1.0×10^{-2}	1.0×10^{-3}
pН	11.3	10.7
α	0.2	-

Where α is the coefficient of conversion of methylamine during its reaction with water.

- 1.3.1- Write the equation of the reaction between methylamine and water.
- 1.3.2- Name the glassware used to measure with precision the two volumes V_0 and V used in the preparation of solution S.
- 1.3.3- Determine the value of α in solution S missing in the above table. Conclude?

2- Preparation of a Buffer Solution

To prepare a buffer solution of pH = 9.0, a volume V₁ of sodium hydroxide solution of concentration $C = 1.0 \times 10^{-2}$ mol.L⁻¹ is added to a volume V₂ = 40 mL of the solution of beaker N° 2 (ammonium chloride solution).

2.1- Write the equation of the reaction which takes place in this mixture.

- 2.2- Show that this reaction is total.
- 2.3- Determine V_1 .

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	First Exercise (6 points)						
Part of the Q			Answer			Mark	
1.1	The formula of	a carboxylic a	cid with saturat	ted acyclic cart	oon chain is	1	
	32 100 2626						
	$C_n \Pi_{2n} O_2$. The	% of oxygen is	$\frac{12n+2n+32}{12n+2n+32}$	$\frac{-100}{2} = 30.30$			
	3200 = 509.04	n + 1163.52;	n = 4. The for	mula of (B) is	then C ₄ H ₈ O ₂		
	and that of	(A) is C_4H_{10}	O, because by	v mild oxidati	on the carbon		
	skeleton rema	uns the same.					
1.2	The carbon cha	in is branched,	, the condensed	structural form	nulas are then:	1	
		Н.ОН		с он			
	$CH_3 - CH - C$	112011	$CII_3 - CII -$				
	(A)		(B)	Ш О			
	2-methyl-1-pro	panol	2-methylprop	anoic acid			
		L					
2.1	The equation of	f the reaction b	etween (A) and	d (B) is:		0,75	
	СЧ	СЧ		СЧ	СЧ		
	CH ₃ - CH - COOH	$I + CH_3 - CH-CH_3$	$_2OH \Rightarrow H_2O + CH$	$H_3 - CH - C - O - H_{H_3}$	$CH_2 - CH - CH_3$		
	(C) is 2-methy	/lpropyl-2-met	hylpropanoate				
2.2	The law of me		[ester][wat	er]		1,25	
	The law of mas	ss action is: \mathbf{K}_{c}	$_{2} = \frac{1}{[acid][alcoh]}$	nol]			
	The composition	on of the system	n is:	-			
		•					
		Acid	Alcohol	Ester	Water		
	Initial state	2	1	0	0		
	⇒ state	2 - x	1 - x	Х	Х		
		$(X_{)^2}$					
	We have then	<u> </u>	= 4 the follow	ing equation ·			
	$\frac{1}{(1-x)(2-x)} = 4$. The following equation :						
	2	V ²					
	$3x^2 - 12x + 8 = 0$, gives to x the acceptable value : $x = 0.845$. The number						
	of moles of (C)	formed is 0.84	45 mol.				
	The yield of the esterification is: $R = \frac{n(ester)exp}{n(ester)exp}$						
	n(ester)the						
	n(ester)the = $n($	alcohol)ini(lin	niting reactant)				
	$R = \frac{0.845}{1000} = 0.8$	845 or 84.5 %.					
	1						

2.3.1	The reaction of esterification is athermic, thus heating does not affect the equilibrium (the limit does not vary), it simply increases the rate to reach this equilibrium.	0,25
2.3.2	the presence of few drops of sulfuric acid does not change the yield of esterification because, in small quantity, this acid acts as a catalyst and has only kinetic role	0,25
2.3.3	Introducing an initial mixture whch is not equimolar supports the reaction in the direction of formation of ester and consequently increases the yield of esterification to exceed 67 %.	0,25
2.4	The two compounds which can replace this acid in the reaction of synthesis of ester are: $\begin{array}{cccccccccccccccccccccccccccccccccccc$	1.25

Second	Exercise	(7	points))
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Part of the Q	Answer						
1.1	The ideal gas equation	n makes it pos	ssible to calculate	p ₀ :	0.5		
	$p_0 = \frac{n \times R \times T}{V} = \frac{1 \times 0}{V}$	$\frac{.082 \times 413}{10} = 3$	3.387 atm.				
1.2	The change of the pre	ssure can be 1	represented by the	following table:	1		
	Instant	p(N ₂ O ₅)	P(NO ₂)	p(O ₂)			
	0	\mathbf{p}_0	0	0			
	t	$p_0 - 2 x$	4 x	Х			
	The total pressure is: $p = p_0 - 2x + 4x + x = p_0 + 3x > p_0$. The pressure increases throughout the reaction.						
1.3	The maximum value	of p is obtaine	ed when N_2O_5 rea	acts completely and	0.75		
	its pressure is cancelled:						
	$p_0 - 2 x = 0$; $x = \frac{1}{2} p_0$ and $p = p_0 + 3 \frac{p_0}{2} = 2.5 p_0 = 8.467$ atm						
1.4	The total pressure p =	$p_0 + 3 x$, where	ere x = $\frac{p - p_0}{3}$ and	d the pressure of	1.25		



Third Exercise (7 points)

Part of the Q	Answer	Mark
1.1	Starting with the same initial concentration, the smallest value of pKa corresponds to the most strong acid and consequently the lowest value of pH:	1.25
	pK_{a} (HF/F ⁻) < pK _a (NH ₄ ⁺ /NH ₃) thus pH ₄ < pH ₂ < 7.	
	Starting with the same initial concentration, the smallest value of pKa	
	corresponds to the most weak base and consequently the lowest value of pH:	
	pK_a (HCOOH/HCOO ⁻)< pK_a (CH ₃ NH ₃ ⁺ /CH ₃ NH ₂) thus 7 < pH_1 < pH_3 .	
	The order of increasing pH is then: pH_4 , pH_2 , pH_1 , pH_3 .	
1.2.1	The equation of this reaction is: $HF + H_2O \Rightarrow F^- + H_3O^+$	0.5
1.2.2	According to the equation of this reaction, we can write:	1
	$[F^{-}] = [H_3O^+] = 10^{-pH} = 10^{-2.65} = 2.23 \times 10^{-3} \text{ mol.L}^{-1} \text{ and}$	
	$[HF] = C_0 - [F_1].$	
	In addition, $K_a (HF/F^-) = \frac{[H_3O^+] \times [F^-]}{[H_3O^+]}$ and	
	$C_0 = \frac{(2.23 \times 10^{-5})^2}{6.31 \times 10^{-4}} + 2.23 \times 10^{-3} = 1.01 \times 10^{-2} \text{ mol.L}^{-1}.$	
1.3.1	The equation of this reaction is: $CH_3NH_2 + H_2O \Rightarrow OH^- + CH_3NH_3^+$	0.5
1.3.2	To take V_0 , we use a volumetric pipette of 10 mL and to have volume V,	0.5
122	we use a volumetric flask of 100 mL.	^
1.3.3	In solution S:	0.75
	$\alpha = \frac{n(CH_3NH_2)(ransformed)}{n(CH_2)(ransformed)} = \frac{n(CH_2)(ransformed)}{n(CH_2)(ransformed)} = \dots = \frac{[CH_2]}{n(CH_2)(ransformed)} = \dots$	
	$n(CH_3NH_2)$ initial $n(CH_3NH_2)$ initial C	
	$\frac{10^{\text{ph-14}}}{10^{-3}} = 0.5.$	
	10°	
2.1	The equation of this reaction is: $OH^- + NH^+ \Rightarrow NH_0 + H_0O$	0.5
2.2	The constant K_{-} of this reaction is:	0.75
	$K_r = 10^{14-9.2} = 6.4 \times 10^4 > 10^4$; thus this reaction is complete.	0.72
2.3	Calculation of V_1 , knowing that $C = C_0$:	1.25
	Since the solution is buffer, NH_4^+ and NH_3 coexist, HO^- is the limiting	
	reactant.	
	$OH^- + NH_4^+ \rightarrow NH_3 + H_2O$	
	Initial state $C \times V_1$ $C_0 \times V_2$ 0 solvant	
	Obtained solution 0 $C(V_2 - V_1)$ $C \times V_1$ solvant	
	pH (solution) = pK _a (NH ₄ ⁺ /NH ₃) + log $\frac{[NH_3]}{[NH_4^+]}$;	
	so : log $\frac{V_1}{V_2 - V_1} = 9 - 9.2 = -0.2$ and $V_2 = 40$ mL, hence $V_1 = 15.5$ mL.	