PROPOSED MARKING GUIDE

Candidate's Name	••••••	••••••	•••••
School:	•••••	•••••	•••••

P525/3

CHEMISTRY (PRACTICAL)

Paper 3 July/Aug. 2022

3 1/4 hours

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48.1



UGANDA TEACHERS' EDUCATION CONSULT (UTEC)

Uganda Advanced Certificate of Education

CHEMISTRY PRACTICAL

Paper 3

3 hours 15 minutes

INSTRUCTIONS TO CANDIDATES:

Answer ALL questions.

Record your answers on this question paper in the spaces provided.

Mathematical tables and silent non-programmable calculators may be used.

Reference books (i.e. text books, books on qualitative analysis, etc) should not be used.

Candidates are **not** allowed to start working with apparatus for the first 15 minutes. This time is to enable candidates to read the question paper and make sure they have all apparatus and chemicals that they may need.

1. You are provided with the following;

FA1, which is 0.4M hydrochloric acid

FA2, which is sodium hydroxide solution

Solid Q which in impure acid

You are required to

- standardize FA2
- Determine the percentage purity of acid Q

PART A

Procedure

Measure accurately 20 cm³ of **FA1** into a 50 cm³ measuring cylinder. Add distilled water carefully to make 50 cm³ of the total solution. Transfer the solution into a clean beaker and label it **FA3**.

Questions:

 (a) Calculate the, (i) number of moles of the acid in 20 cm³ of FA1. 	(1 ½ marks)
1000cm³ of FAI Contain 0.4 Moles of HC	L:
1000cm³ of FAI Contain 0.4 Moles of HC 20cm³ of FAI Contain (0.4x20) Moles	<i></i>
= 0.008 moles V	7
(ii) The molarity of hydrochloric acid in FA3 .	(02 marks)
500m³ of FA3 Contain 0.008 mole 1000 m³ of FA3 Contain (0.008 x100	3
\ 50	,
= 0.16 M	V

3/2

PART B

Procedure

Pipette 20.0 or 25.0 cm³ of **FA3** into a clean conical flask add 2-3 drops of phenolphthalein indicator. Titrate with **FA2** from the burette until the end point. Repeat the titration to obtain consistent results. Record your results in the Table 1 below.

Results:		X		
Volume of pipette used	20.0 L	······································	cı	n ³ . (½ mark)
	IATTA TATA TATA TATA TATA TATA TATA TAT	1	. 2	3
Final burette reading (cm ³)	•	20	0.0.1.0	64 16
Initial burette reading (cm ³)		32.40	32.10	32.10
		0.00	0.00	0.00
Volume of FA2 used (cm ³)		32.40	32.10	31.10
T:4	-	VÍ	VX	(4 ½ marks)
Titre values used to calculate av 32.10	rerage volume	of FA2 used		(½ marks)
		-4- (O. I	::/://	•••••
Average volume of FA2 used) +62°	cm ³ .	(2½ marks)
		+03	CK,	
Questions		せつか	7	
(a). Calculate the molar concen	tration of sodi	um hydroxid	e in FA2.	(5 marks)
Moles of HCI Ital .	earlad	0 11 2 20	······································	
Moles of HCl that r NaOHap + HClap Reaction ratio	DICTO =	0.10736	%:: 0.002	Moles
Naut Aclas	Naclti	(2) 1000		
Keachion ratio	Naoh: HI	Cl in	1:124	
Moles of NaoH = M	oles of H (C = 0	0032/	wes
,		• • • • • • • • • • • • • • • • • • • •		
32.10 cm3 of FA	2 Contai	 u D. DC		SP V OU
10000 3			······································	of Nauti
1000 cm3 of FA	2 Contai	n/ 0.00	32 x 1000) moler
·		1 2	3-10	/
•••••		= 0.0	997 M	N
	••••••••••••••		5(dkkkd	• • • • • • • • • • • • • • • • • • • •
			•••••••	

PART C

Procedure

Weigh accurately about 1.6g of Q and place it in a beaker.

Add to it about 50 cm³ of distilled water and stir to dissolve. Transfer the contents of the beaker into a 250 cm³ volumetric flask. Add distilled water up to the mark. Label the resultant solution **FA4**.

Pipette 20.0 cm³ (or 25.0 cm³) of **FA4** into a conical flask. Add 2-3 drops of phenolphthalein indicator and titrate with **FA2** from the burette until the end point. Repeat the titration until you get consistent results. Record your results in Table 2.

Results:	A (1) /	Y		
Mass of Q and the weighing bottle		` <i>y</i>	g.	
Mass of empty weighing bottle		/ <u> </u>	g.	
Mass of Q used	1.6.	.X	.g.	
Volume of pipette used	20,0	cm ³	(½ mark)	
	1	2	3	
Final burette reading (cm ³)	20.20	40.20	40.50	
Initial burette reading (cm ³)	0.00	20.20	20.50	
Volume of FA2 used (cm ³)	20.20//	20.09	20.00	(v/
Titre values used to calculate average vo	lume of FA2 used	- V ' '	4 ½ markš) 1 ½ marks)	1
20.00, 20.00		1,		
Average volume of FA2 used 20	0.00 tol	$\frac{1}{\sqrt{\chi}}$ cm ³	2 ½ marks)	
Trivolage volume of 1712 about	+03		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
Question	404			
(a) Calculate the number of moles of;	100			
(i) Sodium hydroxide that reacted 1900 cm ³ of FA 2 Con to	ain 0.0997.	Modes of No	(2 marks)	
20.0 cm of FAZ Cont	ain (0.0997)	(20) Mole	T.	
·	1000	···//	<i>/</i> 	
	- 0.00	1994 rud	6.1	
(ii) Acid Q that reacted		(2	½ marks)	
(1 mole of Q reacts with 2 moles 2 Moles of Nathers of 1	of sodium hydroxia	le)	e 51/0	
0.001994 under of N	apil rect	ia # / / Yer	×0019916) Mila
0.001994 moles of N	.400	2 2		7 400
			.0.0.0.17	1 rive
(b) Determine the concentration of Q in I(i) In moldm⁻³	∃ A 4		(2 marks)	
2000 cm3 of FA4 Cont	Til. (0.5009	an wither:	st 🔊	
1000 cm of 1A4 Cont	= $lo poog$	97. × × ×	3/	
1000 cm3 of FA4 Cont	20		MUK)	
	^	0499 M	/	
	= 0.	UTIM	, 	1-
4				

	(ii) In grams per liter (1 mole of Q weigh 90 g)	(1 ½ marks)
	90g of Q W Contain Im Ste	••••••
	Imole of Q Weigh 90g	
	0.0499 miles of a weigh 90 x 0.00	199 /
	4.4.	70 1·1·······
	(c) Calculate the percentage purity of Q. 250 cm ³ of FA4 Contain 169 of Q.	(1 ½ marks)
	1000 cm of 144 Contain 16 X1000)	Percentage Punty = 4.49x100
	= 6.49/2.0°	=70.2%
	,	
2.	You are provided with substance W which contains two cation	s and two anions. Carry
	out the following tests on W to identify the cations and anio evolved. Record your observations and deductions in the Table	n. Identify any gas (es) for al 35 Months

TESTS	OBSERVATIONS	DEDUCTIONS
(a) Heat two spatula ends full of W strongly until there is no further	Pate light green Solid	Ni 2+ Fez+ Gr3+ Cu2+
change.	Colourless liquid Condensati	Hydrated Salf/ water of Crystallisa
	turned Whiste Copper(11)	water of Crystallisa
	Sulphate blue	
	White fumes with a sweet Sm	en Propagione
	turned formed yellow ppt with Brady's reagent.	Vapour
	With Brady's reagent.	Acidet gas,
	Colouress gap turned	CO21, CH3(DO) (U3)
	Colourless gap turned, limewalerimitky and moist blue litmusted	CON CH3(DO) CO3 May be present
	Black residuer	Nig fcu0
b) To one spatula ends full of W in a	rolourless vapour, with	
dry test tube add drops of	a sour Smed of Minegar	Ethanoid ació Vapour y
concentrated sulphuric acid and	turns Moist blugtimus	CH2COO Present
warm	red.	2.132.2

(c)	Dissolve three spatula ends full of W in about 5 cm ³ of water to make a solution To 1 cm ³ of solution add iron (ii) chloride solution and heat.	Brown precipitate Brown Solution	CHIZGOO 02	
(ii)	To the remaining solution of W add sodium hydroxide solution drop wise until there is no further change. Filter and keep both the filtrate and the residue.	Colonidess Filtratet	le Ni2t/Fe2+ 03/2 Zn, Ais, Pb Sn	-/
(d)	Add dilute hydrochloric acid to the filtrate until the solution is just acidic. Divide the solution into four portions.	White precipitate Soluble jú exess aud	Zm, Al3+ Pb, Sn2+	-
(i)	To the first portion of the acidic solution add sodium hydroxide solution drop wise until in excess.	White precipitale Schible in access forming Colontes Solution		
(ii)	To the second portion, add potassium iodide solution	No observable change	Zn2+Al3+Sn2+01	
		Blue fatait form sa blue la me solution.	AlitConfirmed 2	
(iv)	To the fifth portion of acidic filtrate add 2-3 drops Barium nitrate solution	White precipitate	SOFFIEL U	
(e)	Wash the residue and dissolve it in minimum dilute sulphuric. Divide the resultant solution into 3 portions	bissolves forming pale green Bolution	Ni ²⁺ /Fe ²⁺ 01	

i)	To the first portion add sodium hydroxide solution drop wise until in excess.		Ni ^{2t}	1/2
ii)	To the third portion add ammonia solution drop wise until in excess.	Soluble firming blud Solution	Ni ^{2†}	02
iii) A OY A	Use the third portion to carry out a test of your own choice to confirm one of the anions. Add bimethylg lyoximal Schulus add KCNag, Unhill In exess	Red precipitale	Nutf	IŁ
		·		26
3.	You are provided with organic su of T . Carry out the following tests in Table 4 below.	abstance T . You are required to det s on W and record your observation	ns and deductions	
	TESTS	OBSERVATIONS	(18 marks) DEDUCTIONS	1
	Burn a small amount of T on a spatula end or on crucible lid.	White Soliff Crystals	Anomatical	The control of the co
			aliphatic lineate compound or Aliphatic compound with high C:H:	

To a spatula end-ful of T in a test tube add 5 cm³ of water. Shake vigorously and warm. Test the solution with litmus paper. Divide into three equal portions.

Slightly Parhally Soluble phend, aromanical carbonahily solution water, Soluble carbonahily carbonahily solution water, soluble carbonahily solution water, soluble carbonahily solution with litmus paper. Divide firming colourless solution, suspected firming colourless solution, suspected turned blue himus red.

(i)	To the first portion of the solution	No observable thange	Carlony Co 1	
	add 2-3 drops of 2,4- dinitrophenyl hydrazine.	or No yellow orange ppt firmed	absent.	
(ii)	To the second portion of the solution add 2-3 drops of neutral Iron(iii) chloride solution.	Violet purple / Colouration 1	Phenot absent 02	
(iii) (i)	To the 2 cm ³ of ethanol add a spatula end-ful of T and shake to desolve. Add 3-4 drops of cone sulphuric acid and warm the mixture.	Sweet-fruity finelling Product or Sweet fruity Smell	Ester, formed Estentication OB occured 2. Calorytic acid Con firmo d	
(b) Comment on the nature T. Anomahic Compound with a hydroxyl group attached II to benzene ring, also Contains the -efoth (Carboxyhic and group)				
	A. Phenol With	a carboxylic acid	group.	
	Compound with	Structure END	ODH CHR-COOL	