9729/04

Suggested Solutions

DUNMAN HIGH SCHOOL **Preliminary Examination** Year 6

H2 CHEMISTRY

Paper 4 Practical

25 August 2022 2 hours 30 minutes

Candidates answer on the Question Paper.

READ THESE INSTRUCTIONS FIRST

Write your centre number, index number, name and class at the top of this page.

Give details of the practical shift and laboratory where appropriate, in the boxes provided. Write in dark blue or black pen.

You may use an HB pencil for any diagrams or graphs.

Do not use staples, paper clips, glue or correction fluid.

Answer all questions in the spaces provided on the Question Paper.

The use of an approved scientific calculator is expected, where appropriate.

You may lose marks if you do not show your working or if you do not use appropriate units.

Qualitative Analysis Notes are printed on pages 21 and 22.

At the end of the examination, fasten all your work securely together. The number of marks is given in brackets [] at the end of each question or part question.



For Examiner's Use		
1		
	16	
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Total		
Total	55	

9729/04

Answer **all** questions in the spaces provided.

1 Determination of solubility product, K_{sp} , of magnesium carbonate

The solubility of solid magnesium carbonate, MgCO₃, in water is low. In this experiment, you will determine the solubility product, K_{sp} , of magnesium carbonate using volumetric analysis.

$$MgCO_{3}(s) \rightleftharpoons Mg^{2+}(aq) + CO_{3}^{2-}(aq)$$

A saturated solution of magnesium carbonate was prepared by collecting the filtrate after mixing aqueous solutions of magnesium sulfate and sodium carbonate.

You will then determine the amount of carbonate ions left in the filtrate using hydrochloric acid.

FA 1 is a saturated solution of magnesium carbonate, MgCO₃. **FA 2** is 0.003 mol dm⁻³ hydrochloric acid, HC*l*. methyl orange indicator

Note:

- Read the procedures given in part (a) and part (b)(i).
- Part (a) has been done for you. You are not required to carry out part (a), but you will still need to read part (a) to know how FA 1 is prepared.

(a) Preparation of FA 1

- 1. Use a measuring cylinder to transfer 20 cm³ of 0.100 mol dm⁻³ magnesium sulfate to a clean and dry 250 cm³ beaker.
- 2. Use a measuring cylinder to transfer 50 cm³ of 0.0400 mol dm⁻³ sodium carbonate to the same beaker. A white precipitate forms.
- 3. Stir the mixture thoroughly. Leave this mixture to stand for 15 minutes to allow equilibrium to be reached.
- 4. Filter the reaction mixture into a dry 250 cm³ conical flask, using dry filter funnel and filter paper. The filtrate is labelled as **FA 1**. Do not wash the white precipitate with water.

(b) (i) Titration of FA 1 against FA 2

- 1. Fill a burette with **FA 2**.
- 2. Pipette 10.0 cm³ of **FA 1** into a 100 cm³ conical flask.
- 3. Add a few drops of methyl orange indicator to the conical flask.
- 4. Titrate the solution in the conical flask with **FA 2**. The end-point for this titration is reached when the solution changes colour from yellow to orange.
- 5. Record all burette readings, to an appropriate level of precision, in the space provided on page 3.
- 6. Repeat steps 1 to 5 until consistent results are obtained.

Titration results

Final burette reading / cm ³	17.00	34.00
Initial burette reading / cm ³	0.00	17.00
Volume of FA 2 used / cm ³	17.00	17.00

[3]

(ii) From your titrations, obtain a suitable volume of **FA 2** to be used in your calculations. Show clearly how you obtained this volume.

Average volume of **FA 2** used = $\frac{17.00 + 17.00}{2}$ = 17.00 cm³

volume of FA 2 =	cm ³
	[4]

(c) (i) The equation for the reaction in the titration is shown below.

 $2H^{+}(aq) + CO_{3}^{2-}(aq) \rightarrow H_{2}O(I) + CO_{2}(g)$

Calculate the concentration of CO_3^{2-} ions in **FA 1**.

Moles of HCl reacted = $\left(\frac{17.00}{1000}\right)(0.003) = 5.1 \times 10^{-5} \text{ mol}$ 2H⁺ = CO₃²⁻ Moles of CO₃²⁻ in 10.0 cm³ of **FA 1** = $\left(\frac{1}{2}\right)(5.1 \times 10^{-5}) = 2.55 \times 10^{-5} \text{ mol}$

Concentration of $CO_3^{2-} = \frac{2.55 \times 10^{-5}}{0.01} = 2.55 \times 10^{-3} \text{ mol dm}^{-3}$

concentration of CO_3^{2-} ions = mol dm⁻³ [1]

(ii) Calculate the total amount of CO_3^{2-} ions present in the total volume of filtrate prepared in (a).

Total moles of CO₃²⁻ in 70.0 cm³ of **FA 1** = $\binom{70}{10}$ (2.55 ×10⁻⁵) = 1.785 x 10⁻⁴ = 1.79 x 10⁻⁴ mol

total amount of CO_3^{2-} ions = mol [1]

(iii) Hence, calculate the amount of CO_3^{2-} ions precipitated as MgCO₃.

Initial moles of $CO_3^{2-} = \left(\frac{50}{1000}\right)(0.04) = 2 \times 10^{-3} \text{ mol}$ Total moles of CO_3^{2-} precipitated as MgCO₃ = 2 x 10⁻³ - 1.785 x 10⁻⁴ = 1.8215 x 10⁻³ = 1.82 x 10⁻³ mol

amount of CO_3^{2-} ions precipitated as $MgCO_3 = \dots$ [2]

- 4
- (iv) Deduce the amount of Mg²⁺ ions removed by precipitation, in step 3 of the procedure in (a). Hence, calculate the amount of Mg²⁺ ions left in FA 1.

 $MgCO_3(s) \rightleftharpoons Mg^{2+}(aq) + CO_3^{2-}(aq)$

Moles of Mg²⁺ removed by precipitation = 1.82×10^{-3} mol

Initial moles of Mg²⁺ = $\left(\frac{20}{1000}\right)(0.1) = 2 \times 10^{-3} \text{ mol}$ Moles of Mg²⁺ left in **FA 1** = 2 × 10⁻³ – 1.8215 × 10⁻³ = 1.785 × 10⁻⁴ = 1.79 × 10⁻⁴ mol

amount of Mg²⁺ ions removed by precipitation = mol

amount of Mg^{2+} ions left = mol

[2]

(v) Write an expression for the solubility product, K_{sp} , of magnesium carbonate. Include units in your answer.

 K_{sp} of MgCO₃ = [Mg²⁺][CO₃²⁻] mol² dm⁻⁶

[1]

(vi) Calculate a value for the solubility product, K_{sp} , of magnesium carbonate.

$$K_{\rm sp} \text{ of } MgCO_3 = \left(\frac{1.785 \times 10^{-4}}{\frac{70}{1000}}\right) \left(\frac{1.785 \times 10^{-4}}{\frac{70}{1000}}\right) = 6.50 \text{ x } 10^{-6} \text{ mol}^2 \text{ dm}^{-6}$$

K_{sp} =[1]

(d) A student follows the procedures described and obtained a higher K_{sp} value compared to the literature value of 6.82×10^{-6} at 25 °C.

Give a possible explanation for the higher K_{sp} value obtained.

The student did not carry out the experiment at 25 °C. He probably carried out the experiment at a temperature at which the equilibrium position of the reaction lies more towards the Mg^{2+} and CO_3^{2-} ions / the forward reaction is favoured. Hence, the K_{sp} value obtained was higher.

[Total: 16]

[1]

2 Planning

Avogadro's constant, *L*, is defined as the number of particles in one mole of a substance. The currently accepted value is 6.02×10^{23} mol⁻¹.

5

The Avogadro's constant, *L*, can be determined through the electrolysis of acidified aqueous potassium iodide.

During electrolysis, the amount of material discharged at each of the electrodes depends solely on the amount of current that has passed through the system.

A student conducted the electrolysis experiment. When a current was passed through a solution of acidified aqueous potassium iodide, hydrogen gas was produced at one electrode, while iodine was produced at the other electrode.

The amount of hydrogen produced can be determined from the volume of hydrogen gas collected by downward displacement of water. The amount of iodine produced can be determined by titration of a portion of the resultant solution with aqueous sodium thiosulfate.

$$I_2 \textbf{ + 2S}_2 O_3{}^{2-} \rightarrow 2I^- \textbf{ + S}_4 O_6{}^{2-}$$

(a) Explain why the volume of gas collected was higher than the theoretical volume of hydrogen gas that can be produced in the electrolysis.

[1]

The volume of the hydrogen gas that is measured using the downward displacement of water also includes the volume of water vapour.

(b) (i) Plan an investigation to determine the amount of iodine produced in the electrolysis of acidified aqueous potassium iodide, if a current of 0.50 A is passed for 10 minutes, and hence determine a value for the Avogadro's constant, *L*.

You may assume that you are provided with:

- 500 cm³ of 0.500 mol dm⁻³ acidified aqueous potassium iodide,
- 0.0100 mol dm⁻³ aqueous sodium thiosulfate,
- ammeter (instrument used to measure current)
- wires and crocodile clips
- adjustable DC power source (battery)
- graphite rods
- the equipment normally found in a school or college laboratory.

In your plan you should include brief details of:

- a fully labelled diagram of the electrolytic set–up, showing how you would collect and measure the volume of hydrogen gas,
- the apparatus you would use,
- the procedure you would follow for the electrolysis of aqueous potassium iodide.

You need **not** include details of:

- the adjustment of the current setting of the power source to 0.50 A,
- the titration of the resultant solution with aqueous sodium thiosulfate.

Diagram:



- 1. Assemble the apparatus as shown in the diagram above.
- 2. Record the initial reading of the measuring cylinder / burette.
- 3. Switch on the power supply. Adjust current setting to 0.50 A. Start the stopwatch when the current reading is 0.50 A.
- 4. At the end of 10 minutes, stop the stopwatch.
- 5. Switch off the power supply and record the final reading of the measuring cylinder / burette.

[5]

(ii) Table 2.1 gives information about the titration of the resultant solution with aqueous sodium thiosulfate.

total volume of resultant solution after electrolysis / cm ³	500
volume of portion of resultant solution used in titration / cm ³	25.0
volume of sodium thiosulfate used to reach end-point / cm ³	V

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l a	D	е	Ζ.	1

By means of calculations, show how you would use the information in Table 2.1 to determine the Avogadro's constant, *L*, in terms of *V*. Show your working clearly.

[electronic charge, $e = -1.60 \times 10^{-19}$ C]

[3]

titre = $V \text{ cm}^3$ Moles of S₂O₃²⁻ = $\frac{V}{1000} \times 0.01 = 1 \times 10^{-5} V$ mol

 $I_2 \equiv 2S_2O_3^{2-}$

Moles of $I_2 = 1 \times 10^{-5}V \times \frac{1}{2} = 5 \times 10^{-6}V \text{ mol}$ At the anode, $2I^- \rightarrow I_2 + 2e^ \Rightarrow I_2 \equiv 2e^-$ Moles of electrons, n_e , transferred for 25 cm³ = 2 × 5 × 10^{-6}V $= 1 \times 10^{-5}V \text{ mol}$ Moles of electrons, n_e , transferred for 500 cm³ = 1 × 10^{-5}V × $\frac{500}{25}$ $= 2 \times 10^{-4}V \text{ mol}$ $Q = It = 0.50 \times (10 \times 60) = 300 \text{ C}$ $Q = n_eF$ $F = \frac{Q}{n_e} = \frac{300}{2 \times 10^{-4}V}$ F = Le $L = \frac{F}{e} = \frac{300}{2 \times 10^{-4}V} \times \frac{1}{1.60 \times 10^{-19}}$ $= \frac{9.38 \times 10^{24}}{V} \text{ mol}^{-1}$

(c) A student suggested that the percentage error in titre value can be reduced if 0.0500 mol dm⁻³ aqueous sodium thiosulfate was used instead.

Do you agree with this student? Explain your answer.

I do not agree.

With a higher concentration of sodium thiosulfate, the titre value will be lower, resulting in a higher percentage error associated with it.

[1] [Total: 10]

3 Determination of a value for the enthalpy change of reaction between CO₂ and KOH via Hess' Law

The reaction between $KHCO_3(s)$ and HCl(aq) is an endothermic process.

$$\mathsf{KHCO}_3(\mathsf{s}) + \mathsf{HC}l(\mathsf{aq}) \to \mathsf{KC}l(\mathsf{aq}) + \mathsf{CO}_2(\mathsf{g}) + \mathsf{H}_2\mathsf{O}(\mathsf{I}) \qquad \Delta H_1$$

Using ΔH_1 and the standard enthalpy change of neutralisation, you will be able to determine the enthalpy change, ΔH_2 , of the following reaction between CO₂ and KOH via Hess' Law.

$$CO_2(g) + KOH(aq) \rightarrow KHCO_3(s)$$
 ΔH_2

FA 3 is solid potassium bicarbonate, KHCO₃. **FA 4** is 2.00 mol dm⁻³ hydrochloric acid, HC*l*.

(a) Determination of ΔH_1

In this experiment, you will be adding KHCO₃(s) to HC*l*(aq) in a Styrofoam cup. You will then measure the temperature of the reaction mixture over time. Plotting an appropriate graph using the data obtained will allow you to determine ΔH_1 .

In an appropriate format in the space provided on page 11, prepare tables in which to record for your experiment:

- all weighings to an appropriate level of precision,
- all values of temperature, *T*, to an appropriate level of precision,
- all values of time, *t*, recorded to 0.5 min.

It is important that you measure each temperature at the specified time.

Procedure

- 1. Weigh the capped bottle containing solid **FA 3**.
- 2. Place a clean dry Styrofoam cup inside a second Styrofoam cup, then place these in a glass beaker.
- 3. Using a measuring cylinder, add 40 cm³ of **FA 4** into the first Styrofoam cup.
- 4. Stir the solution in the cup with the thermometer. Read and record its temperature, T (time, t = 0.0 min).
- 5. Continue to stir the solution. Read and record *T* every 0.5 min.
- 6. At exactly 2.0 minutes, transfer all the solid **FA 3** into the Styrofoam cup in three portions, taking extra care that the mixture that does not overflow. Stir the mixture but do not read *T*.
- 7. Continue to stir the mixture. Read and record *T* at t = 2.5 min.
- 8. Continue stirring and reading *T* every 0.5 min until t = 7.0 min.
- 9. Reweigh the emptied bottle and its cap.

(i) Results

Mass of capped bottle containing solid FA 3 / g	8.455
Mass of emptied bottle and cap / g	5.423
Mass of FA 3 used / g	3.032

t/min	T/°C
0.0	29.8
0.5	29.8
1.0	29.8
1.5	29.8
2.0	-
2.5	24.6
3.0	24.6
3.5	24.6
4.0	24.7
4.5	24.8
5.0	24.9
5.5	24.9
6.0	25.0
6.5	25.1
7.0	25.2

[3]

(ii) Plot a graph of temperature, *T*, on the y-axis, against time, *t*, on the axis, on the grid in Fig. 3.1.

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Draw a best-fit straight line taking into account all of the points before t = 2.0 min.

Draw another best-fit straight line taking into account all of the points after the temperature of the mixture has started to increase steadily.

Extrapolate both lines to t = 2.0 min.



Fig 3.1

[3]



11

(iii) From your graph, read the minimum temperature, T_{min} , and the maximum temperature T_{max} , at 2.0 min. Record these values in the spaces provided below. Deduce the temperature change, ΔT , at t = 2.0 min.

$T_{min} =$	 	
T _{max} =	 	
$\Delta T =$	 	

 $T_{min} = 24.4 \text{ °C}$ $T_{max} = 29.8 \text{ °C}$ $\Delta T = -5.4 \text{ °C}$

[1]

(iv) Calculate the heat change, q, using the values you deduced in (a)(iii).

You should assume that the specific heat capacity of the solution is 4.18 J g^{-1} K⁻¹, and that the density of the solution is 1.00 g cm⁻³.

 $q = \text{mc}|\Delta T| = (40)(4.18)|-5.4| = 902.88 = 903 \text{ J} (3\text{sf})$

q =[1]

(v) Hence, determine the enthalpy change of reaction, ΔH_1 .

[A_r: K, 39.1; H, 1.0; C, 12.0; O, 16.0]

Moles of KHCO₃ used = $\frac{3.032}{39.1+1+12+16\times3}$ = 0.030290 mol Moles of HC*l* used = $\frac{40}{1000}$ × 2.00 = 0.08 mol KHCO₃ = HC*l* Hence, KHCO₃ is the limiting reagent.

 $\Delta H_1 = + \frac{902.88}{0.030290} \times 10^{-3} = +29.808 = +29.8 \text{ kJ mol}^{-1}$

 $\Delta H_1 = \dots$ [2]

(vi) Calculate the percentage error associated with the measurement of the volume of FA 4.

% error =
$$\frac{\pm 0.5}{40} \times 100 \% = \pm 1.25 \%$$
 [1]

(vii) Suggest the effect that using 80 cm³, rather than 40 cm³, of HC*l* would have on the value for ΔT . Hence, deduce and explain the effect this will have on the value for ΔH_1 .

Since HCl was in excess, the amount of heat produced only depends on the number of moles of KHCO₃ present. Hence, ΔT should be halved as the same amount of heat is dissipated over double the volume.

There should be no change in ΔH_1 as the amount of heat produced and the number of moles of KHCO₃ used is the same.

[1]

(b) Determination of ΔH_2

The enthalpy change of neutralisation between a strong acid and a strong base is $-57.3 \text{ kJ mol}^{-1}$.

Using this information and your answer in (a)(v), draw an appropriate energy cycle to determine ΔH_2 , the enthalpy change of following reaction.

$$CO_{2}(g) + KOH(aq) \rightarrow KHCO_{3}(g) \qquad \Delta H_{2}$$
[2]
$$HCl(aq) + CO_{2}(g) + KOH(aq) \xrightarrow{\Delta H_{2}} KHCO_{3}(g) + HCl(aq)$$

$$\Delta H_{neut} \qquad \Delta H_{1}$$

$$KCl(aq) + CO_{2}(g) + H_{2}O(l)$$

By Hess' Law, $\Delta H_2 = \Delta H_{\text{neut}} - \Delta H_1$ $\Delta H_2 = -57.3 - (+29.808) = -87.108 = -87.1 \text{ kJ mol}^{-1}$

[Total: 14]

4 Inorganic Analysis

13

(a) FA 5 contains two cations and two anions.FA 6 is a solution of ammonium vanadate(V), NH₄VO₃.

You are to perform the tests described in Table 4.1 and record your observations in the table. Unless otherwise stated, the volumes given below are approximate and should be estimated rather than measured.

In all tests, the reagents should be added gradually until no further change is observed unless you are instructed otherwise.

You should indicate clearly at which stage in a test a change occurs, recording your observations alongside the relevant tests. Your answers should include

- details of colour changes and precipitates formed,
- the identities of gases evolved and details of the test used to identify each gas, **except** tests (a)(iv), (vi) and (vii).

If there is no observable change, write **no observable change**.

	test	observations
(i)	Using a measuring cylinder, add 2 cm ³ of FA 5 into a boiling tube.	
	Add aqueous sodium hydroxide slowly, with shaking, until no further change is seen.	Grey-green ppt soluble in excess NaOH giving dark green solution Off-white ppt insoluble in excess
	Filter the mixture. Retain the filtrate for tests (ii) and (iii) .	Off-white residue rapidly turned brown on contact with air / brown residue
	Retain the residue for test (iv).	observed. Filtrate is green.
(ii)	Add 1 cm depth of the filtrate from test (i) into a boiling tube.	
	Add one piece of aluminium foil and warm gently.	Gas evolved turns damp red litmus paper blue. Gas is ammonia.
(iii)	Add 1 cm depth of filtrate from test (i) into a test-tube.	
	Add an equal volume of barium nitrate followed by excess dilute nitric acid.	White ppt seen insoluble in excess
(iv)	Add 1 cm ³ of hydrogen peroxide to the residue from test (i) .	Effervescence observed. Black ppt/ dark brown ppt observed.

Table 4.1

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	lest	observations
(v)	Using a measuring cylinder, transfer 2 cm ³ of FA 6 into a boiling tube.	
	Add 10 cm ³ of dilute sulfuric acid and swirl the mixture gently.	
	The resultant yellow solution is FA 7 , which is an acidified solution of VO_2^+ .	
(vi)	Using a spatula, add a very small quantity of zinc powder to the boiling tube containing FA 7 solution from test (v) . Swirl the mixture gently and record your observations.	Yellow solution turns green.
	Continue to add more zinc powder in small quantities with swirling, until no further colour change is observed. Record all colour changes observed.	Green solution turns blue. Blue solution turns green. Green solution turns violet/purple.
		Effervescence observed. Gas extinguished lighted splint with a 'pop sound. Gas is H ₂ .
	Filter the mixture and retain the filtrate for test (vii) .	
(vii)	To 1 cm depth of the filtrate from test (vi), add an equal volume of aqueous hydrogen peroxide.	Violet solution turns red brown/brown/ orange/orange-brown.
		Effervescence observed. Gas relighted glowing splint. Gas is O ₂ .

- [7]
- (b) (i) Using the results in Table 4.1, deduce the identity of the two cations and two anions present in **FA 5**.

cations

[2]

cations: Cr^{3+} and Mn^{2+} anions: NO_3^- and SO_4^{2-}

(ii) Table 4.2 shows some standard electrode potentials.

Table 4.2

electrode reaction	<i>E</i> ⇔/ V
$VO_2^+ + 2H^+ + e^- \rightleftharpoons VO^{2+} + H_2O$	+1.00

$VO^{2+} + 2H^+ + e^- \rightleftharpoons V^{3+} + H_2O$	+0.34
$V^{3+} + e^- \rightleftharpoons V^{2+}$	-0.26
V ²⁺ + 2e ⁻ ≓ V	-1.20
Zn²+ + 2e⁻ ≓ Zn	-0.76

State and explain which ion is responsible for the final colour of the filtrate in test (a)(vi).

[2]

The E^{\ominus} value for Zn²⁺/Zn of -0.76 V is more negative than all of the above vanadium redox couples except $E^{\ominus}(V^{2+}/V)$. Hence, VO₂⁺ will be reduced to V²⁺.

OR

V²⁺. Zn will not further reduce V²⁺ to V as $E^{\ominus}_{cell} < 0$.

 $E^{\ominus}_{\text{cell}} = (-1.20) - (-0.76) = -0.44 \text{ V} < 0.$

Since $E^{\ominus}_{cell} < 0$, reaction is not spontaneous.

(c) A student performed an analysis using aqueous cobalt(II) chloride, CoC*l*₂. The test and observations are recorded in Table 4.3.

Table 4.3

test	observations
To 1 cm depth of $CoCl_2(aq)$ in a test-tube, add aqueous ammonia dropwise, with shaking, until no further change is seen.	Blue-green ppt formed is partially soluble in excess $NH_3(aq)$ to form a pale brown solution.
Then add about 1 cm depth of aqueous hydrogen peroxide and shake the mixture thoroughly.	Blue-green ppt turned brown. Effervescence observed. Gas relights glowing splint. Gas is O ₂ .

(i) Aqueous $CoCl_2$ contains $[Co(H_2O)_6]^{2+}$ ions.

Write an equation for the formation of the blue-green precipitate.

[1]

 $[Co(H_2O)_6]^{2+} + 2OH^- \rightleftharpoons Co(OH)_2 + 6H_2O OR$ $[Co(H_2O)_6]^{2+} + 2OH^- \rightleftharpoons Co(H_2O)_4(OH)_2 + 2H_2O$

(ii) Using your answer in (c)(i), account for the partial dissolution of the blue-green precipitate to form a pale brown solution when aqueous ammonia was added in excess.

[2]

Ligand exchange occurs where stronger NH_3 ligands displace the weaker H_2O ligands in $[Co(H_2O)_6]^{2+}$ to form pale brown $[Co(NH_3)_6]^{2+}$ complex.

By Le Chatelier's Principle, the decrease in concentration of $[Co(H_2O)_6]^{2+}$ results in position of equilibrium in (c)(i) to shift left, resulting in partial dissolution of Co(OH)₂.

(iii) With reference to changes in oxidation state, explain the formation of the brown precipitate upon the addition of aqueous hydrogen peroxide.

[1]

 $Co(OH)_2$ is oxidised to $Co(OH)_3$ as the oxidation state of Co increases from +2 (in $Co(OH)_2$) to +3 (in $Co(OH)_3$).

[Total: 15]