

**Hale School**

**Year 12 Examination**

**Semester 1, 2020**

Write your name below:

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

**Yr12 ATAR CHEMISTRY**

**UNIT 3**

*Circle your teacher’s initials:*

JWZ PRB KF AD

TIME ALLOWED FOR THIS PAPER

Reading time before commencing: Ten minutes

Working time for paper: Three hours

|  |  |
| --- | --- |
| **For Examiners only** | |
| Section 1 |  |
| Section 2 |  |
| Section 3 |  |
| Total |  |

MATERIAL REQUIRED/RECOMMENDED FOR THIS PAPER

TO BE PROVIDED BY THE SUPERVISOR

This Question/Answer booklet for Sections 1 & 2.

A separate Question/Answer booklet for Section 3.

A separate Multiple Choice Answer sheet for Section 1.

A Chemistry Data Sheet.

TO BE PROVIDED BY THE CANDIDATE

*Standard Items*: Pens, pencils, eraser, ruler

*Special Items*: A calculator satisfying the conditions set by the Curriculum Council, and a '2B' pencil for the separate Multiple Choice Answer sheet.

IMPORTANT NOTE TO CANDIDATES

No other items may be taken into the examination room.

It is your responsibility to ensure that you do not have any unauthorised notes or other items of a non-personal nature in the examination room. Please check carefully, and if you have any unauthorised material with you, hand it to the supervisor **BEFORE** reading any further.

**INSTRUCTIONS TO CANDIDATES**

This paper consists of **THREE SECTIONS** as follows:

**SECTION 1** contains **25 questions worth 2 marks each.** It is the multiple choice section.

Answer **ALL** questions in Section 1 on the Separate Multiple Choice Answer Sheet. Use a **'2B' PENCIL**. **DO NOT USE A BALL POINT OR INK PEN**. If you consider that two or more of the alternative answers are correct then select the BEST alternative. Marks will **NOT** be deducted for incorrect answers. This part is worth 50 marks and should take about 40 minutes.

**Do not use pencil for Sections 2 & 3.**

**SECTION 2** contains **9 short answer questions**. You should answer **ALL** the questions. The answers are to be written in the spaces provided in this Examination booklet. This part is worth 74 marks and should take about 65 minutes.

**SECTION 3** contains **5 extended response questions**. You should answer **ALL** the questions in detail in the **separate question/answer booklet provided.** This part is worth 89 marks and should take about 75 minutes. Express your answers in Section 3 to the appropriate number of significant figures,

At the end of the examination make sure that your **name** is on this Examination paper, the separate Question/Answer Booklet for Section 3 and your Multiple-Choice Answer Sheet.

*SPECIAL INSTRUCTIONS*

*Chemical Equations*

For full marks, chemical equations should refer only to those species consumed in the reaction and any new species produced. These species may be **ions** [for example Ag+(aq)], **molecules** [for example NH3(g), NH3(aq), CH3COOH(l), CH3COOH(aq)] or **solids** [for example BaSO4(s), Cu(s), Na2CO3(s)].

**Structure of this paper**

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Section | Number of questions available | Number of questions to  be answered | Suggested  working time | Marks  available | Percentage  of exam | **Score** |
| Section One:  Multiple-choice | 25 | 25 | 40 min | 50 | 23 |  |
| Section Two:  Short answer | 9 | 9 | 65 min | 74 | 35 |  |
| Section Three:  Extended answer | 5 | 5 | 75 min | 89 | 42 |  |
|  |  |  | **Total** | **213** | **100** |  |
|  |  |  |  |  |  | **%** |

**Section One: Multiple-choice 25% (50 marks)**

This section has **25** questions. Answer **all** questions on the separate Multiple-choice Answer Sheet provided. For each question, shade the box to indicate your answer. Use a **'2B' PENCIL**. **DO NOT USE A BALL POINT OR INK PEN**. If you consider that two or more of the alternative answers are correct then select the BEST alternative. Marks will not be deducted for incorrect answers. No marks will be given if more than one answer is completed for any question.

Suggested working time: 40 minutes.

**Questions 1, 2 and 3 relate to the following equilibrium system.**

Samples of NO3(g) and NO2(g) were injected into an empty flask and equilibrium was established according to the following equation.

NO3(g) + NO2(g) ⇌ N2O5(s) + heat

*colourless brown white*

1. While the system was establishing equilibrium, which of the following would **not** be observed?

1. The brown gas colour fades.
2. A white solid appears.
3. **The pressure in the flask increases.**
4. The temperature in the flask increases.

2. Which of the following conditions would increase the value of K for this equilibrium?

1. **Decreasing the temperature of the system.**
2. Increasing the temperature of the system.
3. Decreasing the volume of the system.
4. Increasing the volume of the system.

3. Which of the following conditions would favour a high equilibrium yield of N2O5(s)?

1. Increased concentrations of NO3(g) and NO2(g)
2. Increased volume of system
3. Increased subdivision of N2O5(s)
4. Increased temperature of system
5. **(i) only.**
6. (ii) only.
7. (i) and (iii) only.
8. (ii) and (iv) only.

**Questions 4 and 5 refer to the equilibria associated with calcification.**

Ocean acidification is a complex process which changes the chemistry of seawater and affects various marine organisms. Calcifying species use the Ca2+(aq) and CO32-(aq) in seawater to produce CaCO3(s), which then forms structures such as the shells or exoskeletons of marine organisms.

Two of the chemical equilibria associated with the process of calcification are shown below.

*Equilibrium 1:* CO2(g) + H2O(l) + CO32-(aq) ⇌ 2 HCO3-(aq)

*Equilibrium 2:* Ca2+(aq) + CO32-(aq) ⇌ CaCO3(s)

4. State the effect of an increased CO2(g) concentration on the rate of each forward reaction.

**Equilibrium 1 Equilibrium 2**

1. **increased rate decreased rate**
2. increased rate increased rate
3. decreased rate increased rate
4. decreased rate decreased rate

5. State the effect of an increased CO2(g) concentration on the equilibrium position of each reaction.

**Equilibrium 1 Equilibrium 2**

1. reverse favoured forward favoured
2. reverse favoured reverse favoured
3. **forward favoured reverse favoured**
4. forward favoured forward favoured

6. The endothermic autoionisation of water can be represented by the equation below.

H2O(l) + H2O(l) + heat ⇌ H3O+(aq) + OH-(aq)

Distilled water at a temperature of 15 °C would have

1. a concentration of hydronium ions greater than 1.0 x 10-7 mol L-1.
2. a concentration of hydroxide ions greater than 1.0 x 10-7 mol L-1.
3. a Kw value greater than 1.0 x 10-14.
4. **a pH greater than 7.**

7. Consider the following graph, which provides information regarding the solubility of cerium(III) sulfate, Ce2(SO4)3(s), at various temperatures.

An 8 g sample of Ce2(SO4)3(s) was added to a beaker containing 100 g of water at 20 °C. Equilibrium was established according to the following equation.

Ce2(SO4)3(s) ⇌ 2 Ce2+(aq) + 3 SO42-(aq)

If the temperature of the system is decreased, which of the following is **not** correct?

1. The conductivity of the solution would increase.
2. **The rate of dissociation would increase.**
3. The forward reaction would be favoured.
4. The value of K would increase.

8. Consider the following equilibrium system.

6 NOCl(g) + 237 kJ ⇌ 6 NO(g) + 3 Cl2(g)

A chemist wanted to;

* + favour the reverse reaction
  + decrease the rate of the reverse reaction
  + maintain a constant K value

Which change should they impose on the system to satisfy all 3 criteria?

1. Decrease the temperature of the system.
2. Decrease the volume of the system.
3. **Remove some NOCl(g) from the system.**
4. Add some NO(g) to the system.

9. ‘Black powder’ is typically composed of solid potassium nitrate and charcoal dust (C7H4O). When this powder is ignited, the products formed are a mixture of solid potassium carbonate, carbon dioxide gas, water vapour and nitrogen gas.

Identify the element oxidised and reduced in this reaction.

**Oxidised Reduced**

1. N K
2. **C N**
3. O N
4. C H

10. Compare the four halogens in their elemental form; F2(g), Cl2(g), Br2(l) and I2(s). Which of the following statements is **correct**?

1. F2(g) exists in the highest oxidation state.
2. Cl2(g) is the strongest reducing agent (reductant).
3. Br2(l) has the lowest tendency to gain electrons.
4. **I2(s) has the lowest reduction potential.**

11. ‘Disproportionation’ refers to a redox reaction, where atoms of a particular element within a single species are simultaneously oxidised and reduced. Which of the following would **not**  be regarded as a disproportionation reaction?

1. Cl2(g) + 2 OH-(aq) → ClO-(aq) + Cl-(aq) + H2O(l)
2. 2 H2O2(aq) → 2 H2O(l) + O2(g)
3. **NH4NO3(s) → N2O(g) + 2 H2O(g)**
4. 2 CuCl(s) → CuCl2(s) + Cu(s)

12. When zinc metal is placed into a solution of 1 mol L-1 nitric acid, nitrogen dioxide gas is formed and the zinc metal dissolves.

Which redox equation **best** represents the reaction occurring?

(a) Zn(s) + 2 HNO3(aq) + 2 H+(aq) → Zn2+(aq) + 2 NO2(g) + 2 H2O(l)

(b) Zn(s) + NO3-(aq) + 2 H+(aq) → Zn2+(aq) + NO2(g) + 2 H2O(l)

(c) **Zn(s) + 2 NO3-(aq) + 4 H+(aq) → Zn2+(aq) + 2 NO2(g) + 2 H2O(l)**

(d) Zn(s) + HNO3(aq) + 3 H+(aq) → Zn2+(aq) + NO2(g) + 2 H2O(l)

**Questions 13, 14 and 15 refer to the following information.**

An electrochemical cell is set up with an unknown metal half–cell joined by a salt bridge to a hydrogen half–cell with a platinum electrode. Under standard conditions a voltage of between 0.7 and 0.8 V is produced.

13. Which one of the following metals is likely to be deposited at the cathode?

(a) **silver**

(b) platinum

(c) zinc

(d) iron

14. Which one of the following best describes what will be observed at the platinum anode?

(a) Hydrogen ions will build up.

(b) **Bubbles of gas.**

(c) A build-up of a silver metal.

(d) The anode will dissolve.

15. Which two of the following statements best explain the function of the salt bridge?

The salt bridge

(i) contains a non-reactive electrolyte solution.

(ii) prevents direct contact between the oxidising and reducing agents.

(iii) ions flow through the salt bridge towards the electrodes.

(iv) prevents the build-up of charge on the electrodes.

(a) i and ii only

(b) ii and iii only

(c) i and iv only

(d) **ii and iv only**

**Questions 16, 17 and 18 refer to metal corrosion.**

Silver metal is particularly susceptible to a form of corrosion known as tarnishing. The chemical equation for this corrosion process is shown below.

*Corrosion of silver:* 4 Ag(s) + 2 H2S(g) + O2(g) → 2 Ag2S(s) + 2 H2O(l)

The corrosion of iron is a particularly damaging and expensive problem, due to the extensive use of iron in modern life. The chemical equation for this corrosion process is shown below.

*Corrosion of iron:* 4 Fe(s) + 3 O2(g) + 2 H2O(l) → 2 Fe2O3.H2O(s)

16. Select the option which correctly identifies the change in oxidation number of each element involved in the process of silver corrosion.

**Ag H S O**

1. **(0) to (+1) no change no change (0) to (-2)**
2. no change (0) to (+1) no change (0) to (-2)
3. (0) to (+2) no change (-2) to (-4) no change
4. (0) to (+1) no change (-2) to (-4) no change

17. Select the statement that is **not** common to both the corrosion of silver **and** the corrosion of iron.

1. The metal is the anodic site.
2. Oxygen gas is the oxidising agent.
3. The metallic element forms a compound.
4. **A thin layer of metallic oxide forms on the metals.**

18. Which one of the following could slow the corrosion of a small strip of iron?

(a) attaching a piece of tin to the iron

(b) attaching a piece of copper to the iron

(c) **attaching a piece of zinc to the iron**

(d) storing the iron under water

**Questions 19 and 20 refer to the following information.**

The reaction between magnesium metal and water occurs very slowly in cold water.

The balanced chemical equation is shown below. The reaction is much faster with steam (water vapour > 100 °C).

Mg(s) + 2 H2O(ℓ) → Mg(OH)2(s) + H2(g)

19. Which one of the following is the E° value for this reaction?

(a) +3.19 V

(b) –1.53 V

(c) **+1.53 V**

(d) +0.35 V

20. The reaction is exothermic but in order for magnesium to react with steam the magnesium is first ignited in a hot flame. Which one of these is the likely reason for the magnesium being ignited first?

In the reaction between magnesium and steam

(a) the enthalpy of reactants is higher than products.

(b) the activation energy is low.

(c) **the activation energy is high.**

(d) heat must be gained from the surroundings.

21. Four beakers (A, B, C and D) were placed on a laboratory bench, each containing distilled water and several drops of universal indicator. A small sample of a different salt was then dissolved into each beaker, according to the table below.

|  |  |  |  |
| --- | --- | --- | --- |
| **Beaker A** | **Beaker B** | **Beaker C** | **Beaker D** |
| + NH4NO3(s) | + MgF2(s) | + KCH3COO(s) | + Na3PO4(s) |

The colour of the solution in beaker

1. A would change from green to blue.
2. B would change from green to yellow.
3. C would change from green to orange.
4. **D would change from green to purple.**

22. Consider the equations below.

H2CO3(aq) + H2O(ℓ) ⇌ HCO3–(aq) + H3O+(aq)

HCO3–(aq) + H2O(ℓ) ⇌ CO32–(aq) + H3O+(aq)

Which one of the following species is the conjugate base of HCO3–?

(a) H2O

(b) **CO32–**

(c) H3O+

(d) H2CO3

23. Which two of the following pairs of solutions could form buffers?

(i) HCℓ/Cℓ–

(ii) NH3/HCℓ

(iii) H2SO4/HSO4–

(iv) CH3COOH/CH3COO–

(a) i and ii only

(b) ii and iii only

(c) iii and iv only

(d) **ii and iv only**

24. Which of the following pieces of equipment should be rinsed with the relevant chemical solution, rather than just distilled water, before using in an acid – base titration?

(i) burette

(ii) conical flask

(iii) pipette

(iv) volumetric flask

(a) i only

(b) **i and iii only**

(c) i, iii and iv only

(d) i, ii and iv only

25. The overall equation for the discharge reaction of a lead acid car battery is shown below.

Pb(s) + PbO2(s) + 4 H+(aq) + 2 SO42–(aq) → 2 PbSO4(s) + 2 H2O(ℓ)

Which species is the oxidising agent?

(a) Pb

(b) **PbO2**

(c) PbSO4

(d) H+

End of Section One

**Section Two: Short answer 35% (74 marks)**

This section has **9** questions. Answer **all** questions. Write your answers in the spaces provided.

When calculating numerical answers, show your working or reasoning clearly. Express numerical answers to the appropriate number of significant figures and include appropriate units where applicable.

Spare pages are included at the end of this booklet. They can be used for planning your responses and/or as additional space if required to continue an answer.

* Planning: If you use the spare pages for planning, indicate this clearly at the top of the page.
* Continuing an answer: If you need to use the space to continue an answer, indicate in the original answer space where the answer is continued, i.e. give the page number. Fill in the number of the question(s) that you are continuing to answer at the top of the page.

Suggested working time: 65 minutes.

**Question 26 (6 marks)**

Consider the following chemical experiments. Describe the reactants and also the contents of the flask at the end of the experiment.

(a) A small piece of copper metal is placed in an excess of iron(II) sulfate solution. (2 marks)

**A shiny salmon pink solid was added to a clear pale-green solution. The contents of the flask remained the same at the end of the experiment.**

(b) A **large** piece of cobalt metal is placed in a small volume of 2.00 mol L–1 hydrochloric acid.

(2 marks)

**A shiny grey/silver metallic solid was added to a clear colourless solution. The solid dissolved producing a pink solution. Some of the shiny solid remained.**

(c) Solutions of silver nitrate and sodium chloride are mixed. (2 marks)

**A clear colourless solution was added to a clear colourless solution to produce a white solid which settles beneath a clear colourless solution.**

**Question 27 (10 marks)**

In 1826 when many of the elements of the periodic table were yet to be discovered, a scientist added aqueous chlorine (Cℓ2(aq)) to the solid residue left after evaporating sea water. This yielded a pungent smelling orange solution. After further testing he concluded it was a new element.

Part of the material safety data sheet for chlorine is reproduced below

**SAFETY DATA SHEET–Chlorine**

**Hazards identification :**

OXIDIZING GASES - Category 1

GASES UNDER PRESSURE - Compressed gas

ACUTE TOXICITY (inhalation) - Category 2

SKIN CORROSION/IRRITATION - Category 1

SERIOUS EYE DAMAGE/ EYE IRRITATION - Category 1

AQUATIC HAZARD (ACUTE) - Category 1

AQUATIC HAZARD (LONG-TERM) - Category 1

**Signal word :**

Danger

**Hazard statements :**

May cause or intensify fire; oxidizer.

Contains gas under pressure; may explode if heated.

Fatal if inhaled.

Causes severe skin burns and eye damage.

Very toxic to aquatic life with long lasting effects.

(a) Write a balanced redox equation for the reaction between chlorine and the likely species present in the salt residue which produced the new element. (2 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| Cℓ2(aq) + 2 Br–(aq) → Br2(aq) + 2 Cℓ –(aq)  correct species  balanced equation  (State symbols not required) | **1**  **1** |
| **Total** | **2** |

At the time the experiment was done the chemicals in the salt residue were unknown as were the nature of any of the products of the reaction with chlorine.

(b) State two safety precautions you would take if you were to repeat the experiment.

(2 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| Any two   * Wear gloves * Wear a face mask * Carry out experiment in a fume hood * Wear protective clothing | **1–2** |
| **Total** | **2** |

**Question 27** (continued)

Chlorine gas is a stronger oxidising agent than solid iodine.

(c) Explain this statement with the aid of suitable half equations and standard reduction potentials. (3 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| Chlorine has a higher tendency to be reduced (+1.36 V) than iodine (+0.54 V) (therefore chlorine is a stronger oxidising agent) | **1** |
| Includes half equations and reduction potentials  Cℓ2(aq) + 2e– → 2 Cℓ –(aq) E° = +1.36 V  I2(aq) + 2e– → 2 I–(aq) E° = +0.54 | **1** |
| refers to half equations and reduction potentials | **1** |
| **Total** | **3** |

(d) Write three observations for the reaction when excess aqueous chlorine (Cℓ2(aq)) is slowly added to a potassium iodide solution (KI(aq)).

(3 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| Initial pale yellow solution  added to a colourless solution  Which changes to a brown solution | **1**  **1**  **1** |
| **Total** | **3** |

**Question 28 (8 marks)**

Calculate the final pH of a solution made by dissolving 1.54g NaOH(s) in 100.0 mL of 0.100 mol L–1 H2SO4(aq). Assume no volume change.

|  |  |
| --- | --- |
| **Description** | **Marks** |
| n(NaOH) = m/M = 1.54/39.998 = 0.0385 | **1** |
| n(NaOH) = n(OH–) = 0.0385 mol | **1** |
| n(H2SO4) = 0.100 x 0.100 = 0.0100 mol | **1** |
| n(H+) = 2 x n(H2SO4) = 0.0100 x 2 = 0.0200 mol | **1** |
| n(OH–) – n(H+) = 0.0385 – 0.0200 = 0.0185 mol | **1** |
| [OH–] = n/v = 0.0185/0.100 = 0.185 mol L–1 | **1** |
| [H+] = 10–14/0.185 = 5.405 x 10–14 | **1** |
| pH = –log[5.405 x 10–14] = 13.3 | **1** |
| **Total** | **8** |

Alternative calculation using pOH

|  |  |
| --- | --- |
| **Description** | **Marks** |
| n(NaOH) = m/M = 1.54/39.998 = 0.0385 | **1** |
| n(NaOH) = n(OH–) = 0.0385 mol | **1** |
| n(H2SO4) = 0.100 x 0.100 = 0.0100 mol | **1** |
| n(H+) = 2 x n(H2SO4) = 0.0100 x 2 = 0.0200 mol | **1** |
| n(OH–) – n(H+) = 0.0385 – 0.0200 = 0.0185 mol | **1** |
| [OH–] = n/v = 0.0185/0.100 = 0.185 mol L–1 | **1** |
| pOH = – log[OH–] = 0.7328 | **1** |
| pH = 14 – pOH = 13.3 | **1** |
| **Total** | **8** |

**Question 29 (6 marks)**

Permanganic acid (HMnO4) can be found as an unstable red-violet solution which decomposes at temperatures above 40 °C.

It can be made by mixing a solution containing manganese(II) ions with powdered lead dioxide, PbO2, in the presence of sulfuric acid. This suspension is then stirred. The reaction produces permanganic acid and a precipitate of lead(II) sulfate.

Write oxidation and reduction half-equations and an overall redox equation for this reaction.

You may assume the sulfuric acid is present entirely as H+(aq) and SO42-(aq).

|  |  |
| --- | --- |
| Oxidation  half-equation  **(2)** | **Mn2+ + 4 H2O → MnO4- + 8 H+ + 5 e- (x2)**  **OR**  **Mn2+ + 4 H2O → HMnO4 + 7 H+ + 5 e- (x2)** |
| Reduction  half-equation  **(2)** | **PbO2 + 4 H+ + SO42- + 2 e- → PbSO4 + 2 H2O (x5)** |
| Overall redox equation  **(2)** | **2 Mn2+ + 4 H+ + 5 PbO2 + 5 SO42- → 2 MnO4- + 5 PbSO4 + 2 H2O**  **OR**  **2 Mn2+ + 6 H+ + 5 PbO2 + 5 SO42- → 2 HMnO4 + 5 PbSO4 + 2 H2O** |

**Question 30 (7 marks)**

Potassium hydrogen iodate, KH(IO3)2, can be used as a primary standard in acid-base titrations, providing the percentage purity of the solid is known.

An 8.435 g sample of **impure** KH(IO3)2(s) was dissolved in distilled water, transferred to a volumetric flask and made up to 500.0 mL. Then 25.00 mL aliquots of this solution were titrated against standardised 0.05084 mol L-1 NaOH(aq). An average titre of 20.70 mL was required to reach the equivalence point.

The chemical equation for the titration reaction that occurred is given below.

KH(IO3)2(aq) + NaOH(aq) → KNa(IO3)2(aq) + H2O(l)

Calculate the percentage purity of the KH(IO3)2(s) that was used in this experiment.

**n(NaOH) = cV**

**= 0.05084 x 0.02070**

**= 0.001052388 mol (1)**

**n(KH(IO3)2) in 25 mL) = 0.001052388 mol (1)**

**n(KH(IO3)2) in 500 mL) = 0.001052388 x 500 / 25**

**= 0.02104776 mol (1)**

**= n(KH(IO3)2 in powder sample) (1)**

**m(KH(IO3)2) = nM**

**= 0.02104776 x 389.908 (1 for M)**

**= 8.20669 g (1)**

**% purity = 8.20669 / 8.435 x 100**

**= 97.293 % (1)**

**= 97.29% (4 SF)**

**Question 31 (8 marks)**

Consider the following reaction that has been allowed to establish equilibrium.

2 OH-(aq) + Cl2(g) ⇌ Cl-(aq) + ClO-(aq) + H2O(l)

The following table considers the effects of imposing different changes upon this system.

(a) Complete the following table by predicting which direction would be favoured, as well as the resulting effect on the concentration of ClO-(aq) when compared with the original equilibrium. Consider each of the imposed changes in isolation.

|  |  |  |
| --- | --- | --- |
| **Imposed change** | **Favoured direction**  (forward, reverse, no shift) | **Effect on final concentration of ClO-(aq)**  (increase, decrease, no change) |
| A small amount of NaClO(s) was added | **reverse** | **increase** |
| The volume of the canister was decreased | **forward** | **increase** |
| Distilled water was added | **no shift** | **decrease** |
| Several drops of 2 mol L-1 Mg(NO3)2(aq) was added | **reverse** | **decrease** |

**Question 32 (10 marks)**

A water tank contained 25.0 kL of contaminated water with a pH of 10.1. A chemist decided to add some solid acetylsalicylic acid crystals to the water tank, to lower the pH. Acetylsalicylic acid is a monoprotic acid with the formula C9H8O4 (M = 180.158 g mol-1).

Calculate the mass of acetylsalicylic acid crystals that should be added to the tank and dissolved in the water, to lower the pH to 7.8.

**At pH 10.1;**

**[H+] = 10-10.1 = 7.94328 x 10-11 mol L-1 (1)**

**[OH-] = 10-14 / 10-10.1 = 1.258925 x 10-4 mol L-1 (1)**

**n(OH-) = cV**

**= 1.258925 x 10-4 x 25 000**

**= 3.1473125 mol (1)**

**At pH 7.8;**

**[H+] = 10-7.8 = 1.584893 x 10-8 mol L-1 (1)**

**[OH-] = 10-14 / 10-7.8 = 6.309573 x 10-7 mol L-1 (1)**

**n(OH-) = cV**

**= 6.309573 x 10-7 x 25 000**

**= 0.015773933 mol (1)**

**n(OH- at pH 10.1) – n(OH- at pH 7.8) = 3.1473125 - 0.015773933**

**= 3.1315386 mol (1)**

**Therefore want to add 3.1315386 mol of H+ (1)**

**Since monoprotic, need n(C9H8O4) = 3.1315386 mol (1)**

**m(C9H8O4) = nM**

**= 3.1315386 x 180.158**

**= 564.17 g**

**= 564 g (1)**

**Question 33 (9 marks)**

A chemistry class was given four (4) jars labelled A, B, C and D.

Each jar contained small pieces of a silver-coloured metal. The teacher informed the students that the metals were tin, cobalt, cadmium and nickel.

The students were asked to design an investigation that would determine the identity of the metal in each jar.

The students set up 4 test tubes, each containing 5 mL of 1.0 mol L-1 lead(II) nitrate solution, Pb(NO3)2(aq). A piece of each different metal was then added to each test tube. The students’ observations are recorded in the table below.

|  |  |
| --- | --- |
|  | **Observations** |
| Pb(NO3)2(aq) + metal A | A silver metal dissolves in a colourless solution.  A new grey solid and a green solution are produced. |
| Pb(NO3)2(aq) + metal B | A silver metal dissolves in a colourless solution.  A new grey solid is produced. Solution remains colourless. |
| Pb(NO3)2(aq) + metal C | A silver metal dissolves in a colourless solution.  A new grey solid is produced. Solution remains colourless. |
| Pb(NO3)2(aq) + metal D | A silver metal dissolves in a colourless solution.  A new grey solid and a pink solution are produced. |

(a) Identify metals A and D. (2 marks)

|  |  |
| --- | --- |
| A | **Nickel (1)** |
| D | **Cobalt (1)** |

To distinguish between metals B and C, one student suggested adding a piece of each to separate solutions of aqueous nickel bromide, NiBr2(aq).

(b) Justify how this would allow metals B and C to be distinguished. Include observations for both samples and a relevant chemical equation in your answer.

(5 marks)

1. **tin metal would not react with the NiBr2(aq) whereas cadmium metal would react with NiBr2(aq)**
2. **reaction with cadmium has EMF > 0 / reaction is spontaneous OR reaction with tin has EMF < 0 / is non-spontaneous**
3. **observations with tin metal – solution would remain green**
4. **observations with cadmium metal – green solution would fade as silver metal dissolves, new silver / grey metal / precipitate produced**
5. **Cd(s) + Ni2+(aq) → Ni(s) + Cd2+(aq)**

(c) Using the results of this investigation, how would the students be able to identify which of the metals (A, B, C or D) was the strongest reducing agent (reductant)?

(2 marks)

1. **whichever metal of B or C that reacted with NiBr2(aq) would be the strongest reductant**
2. **this is because it would be the metal most easily oxidised / the metal with the highest oxidation potential**

**Question 34 (10 marks)**

On a table of standard reduction potentials, the ‘hydrogen half-cell’ or ‘standard hydrogen electrode’ is designated an EMF of 0 V. The standard hydrogen half-cell is set up using a platinum electrode, hydrogen gas (H2) and a solution containing hydrogen ions (H+).

(a) State the values that would be used when setting up the hydrogen half-cell under standard conditions. (2 marks)

|  |  |
| --- | --- |
| Partial pressure of H2(g) | **100 kPa** |
| Concentration of H+(aq) | **1 mol L-1** |

A standard hydrogen half-cell was joined by wires and a salt bridge to a second half-cell. Two different experiments were performed;

* In Experiment 1, the Ag+(aq)/Ag(s) half-cell was used.
* In Experiment 2, the Zn2+(aq)/Zn(s) half-cell was used.

A diagram of the experimental set up is shown below. Assume all half-cells were set up under standard conditions.

V

H+(aq)

Pt(s)

H2(g)

EITHER

Ag+(aq)/Ag(s)

OR

Zn2+(aq)/Zn(s)

(b) In which experiment would the greater EMF have been produced? State the value of this EMF. (2 marks)

1. **Experiment 1 / silver half-cell**
2. **EMF = +0.80 V**

(c) Discuss how the role of the hydrogen half-cell differs in Experiment 1 and 2. Include a relevant half-equation for the hydrogen half-cell in each case.

(6 marks)

**In Experiment 1 the hydrogen half-cell; (2) for any 2 of the following**

* **is the anode**
* **is the site where oxidation occurs**
* **has a negative (-) polarity**
* **has a lower reduction potential than Ag/Ag+**

**H2(g) → 2 H+(aq) + 2 e- (1)**

**In Experiment 2 the hydrogen half-cell (2) for any 2 of the following**

* **is the cathode**
* **is the site where reduction occurs**
* **has a positive (+) polarity**
* **has a higher reduction potential than Zn/Zn2+**

**2 H+(aq) + 2 e- → H2(g) (1)**

END OF SECTION TWO

**Hale School**

**Semester One**

**Examination, 2020**



Write your name below:

**Solutions**

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

**Yr12 ATAR CHEMISTRY**

Circle your teacher’s initials:

JWZ PRB KF AD

\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_

**UNIT 3**

|  |  |
| --- | --- |
| **For Examiners only** | |
| Part 3 |  |

Section 3 Question and Answer Booklet

**Section Three: Extended answer 40% (89 marks)**

This section contains **five (5)** questions. You must answer **all** questions. Write your answers in the spaces provided below.

Where questions require an explanation and/or description, marks are awarded for the relevant chemical content and also for coherence and clarity of expression. Lists or dot points are unlikely to gain full marks.

**Final answers to calculations should be expressed to the appropriate number of significant figures.**

Spare pages are included at the end of this booklet. They can be used for planning your responses and/or as additional space if required to continue an answer.

* Planning: If you use the spare pages for planning, indicate this clearly at the top of the page.
* Continuing an answer: If you need to use the space to continue an answer, indicate in the original answer space where the answer is continued, i.e. give the page number. Fill in the number of the question(s) that you are continuing to answer at the top of the page.

Suggested working time: 75 minutes.

**Question 35 (18 marks)**

Silver metal can be extracted from the ore ‘argentite’ which contains the compound Ag2S(s). In the first step of this extraction process, the ore containing Ag2S(s) is crushed, and then a solution of sodium cyanide, NaCN(aq), is added to leach the silver out of the ore. This forms the complex ion sodium argentocyanide, Na[Ag(CN)2](aq), as shown in the equation below.

Ag2S(s) + 4 NaCN(aq) ⇌ 2 Na[Ag(CN)2](aq) + Na2S(aq)

(a) Write the equilibrium constant expression for this reaction. (1 mark)

|  |
| --- |
| **K = [Na[Ag(CN)2]]2 [Na2S]**  **[NaCN]4**   1. **if completely correct** |

If 12.5 kL of 5.0 g L-1 NaCN(aq) was added to the leaching tank and poured over the crushed ore;

(b) Calculate the maximum mass of Ag2S(s) that could react. (4 marks)

**m(NaCN) = cV**

**= 5 x 12500**

**= 62500 g (1)**

**n(NaCN) = m / M**

**= 62500 / 49.01**

**= 1275.25 mol (1)**

**n(Ag2S) = ¼ x n(NaCN)**

**= ¼ x 1275.25**

**= 318.812 mol (1)**

**m(Ag2S) = nM**

**= 318.812 x 247.87**

**= 79024.05 g**

**= 79000 g (2 s.f.) (1)**

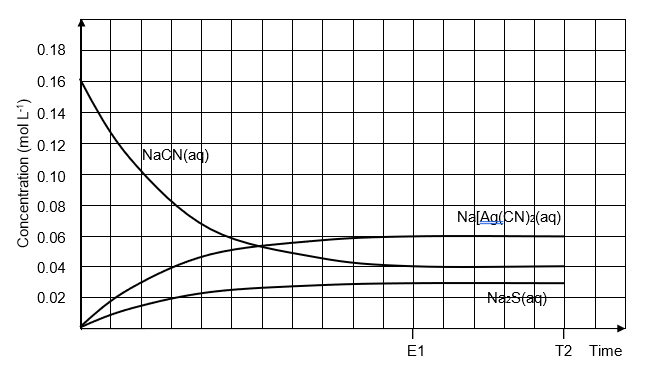
**= 79 kg**

(c) Explain, in terms of the collision theory, why the ore is crushed before mixing with the NaCN(aq) solution. (3 marks)

1. **Crushing the ore increases surface area / state of subdivision of Ag2S(s)**
2. **This will increase the frequency of collision between reactants**
3. **Therefore, a faster reaction rate occurs**

A particular leach tank contained an excess mass of crushed argentite ore containing Ag2S(s). A 0.16 mol L-1 solution of NaCN(aq) was poured over the ore, and the reactants were mixed. At Time E1, equilibrium was established. The concentration of the NaCN(aq) was measured again and found to be 0.04 mol L-1.

(d) On the axes below, sketch curves showing the concentration of all relevant species, from Time 0 until the establishment of equilibrium at Time E1. Continue your curves from Time E1 to Time T2. Label each curve. (5 marks)



1. **NaCN line correctly drawn**
2. **Na[Ag(CN)2] line correctly drawn**
3. **Na2S line correctly drawn**
4. **Labels on each line**
5. **Horizontal lines E1 to T2**

At Time T2, a temperature change is imposed on the system. The following rate graph was drawn to illustrate the effects of this imposed change.

reverse

forward

l l

T2 E2

Time

Reaction rate

(e) State whether the temperature was increased or decreased at Time T2. (1 mark)

1. **Decreased**

(f) Justify, using the information provided in the rate graph above, whether the forward reaction is endothermic or exothermic. (4 marks)

1. **the forward reaction rate is not decreased as much as the reverse reaction rate**
2. **this shows that the forward reaction has been favoured**
3. **a temperature decrease will favour the exothermic direction (in order to produce more heat)**
4. **therefore the forward reaction much be exothermic**

This page has been left blank intentionally

**Question 36 (18 marks)**

The ‘Hydroflex’ train was launched in the United Kingdom last year, as Britain’s first hydrogen-fuelled train. The technology is currently being trialled, but the government is hoping that hydrogen trains will be in common use by 2022, and that all diesel trains will be removed from use by 2040.

The diagram below provides some information about how the hydrogen train works.

Hydrogen is stored in **four fuel tanks**

➊

The **fuel cell** converts hydrogen and oxygen into water and electricity

➋

Energy can be stored in **two lithium-ion** batteries

➌

Electric motors move the train

➍

How a zero-emission train works

The fuel cell used is more specifically described as a ‘polymer electrolyte membrane’ (or PEM) fuel cell stack. These cells function at a temperature of around 50-100 °C. The electrolyte is a water-based acidic polymer membrane.

(a) Write oxidation and reduction half-equations and an overall equation to represent the reaction occurring in the PEM fuel cell. Calculate the EMF produced under standard conditions. (4 marks)

|  |  |
| --- | --- |
| Reduction  half-equation | **O­2(g) + 4 H+(aq) + 4 e- → 2 H2O(l)** |
| Oxidation  half-equation | **H2(g) → 2 H+(aq) + 2 e-** |
| Overall equation | **2 H2(g) + O2(g) → 2 H2O(l)** |
| EMF under standard conditions | **+1.23 V** |

The PEM is also often called a ‘proton-exchange membrane’ because one of its main functions is to conduct protons from one half-cell to another. The design of the PEM is instrumental in the function of the fuel cell. As well as conducting protons it has to;

* separate the two half-reactions
* prevent hydrogen and oxygen gas crossover
* allow the conduction of protons but not the conduction of electrons
* resist degradation from the surrounding environment.

(b) At which electrode are protons produced? Justify your answer. (2 marks)

1. **produced at the anode**
2. **protons are hydrogen ions / H+**

(c) Why must the PEM have the ability to conduct protons but **not** electrons? (2 marks)

1. **must conduct protons to act as the salt bridge / allow internal flow of charge / maintain electrical neutrality**
2. **must NOT conduct electrons as cell would short circuit / an unwanted direct reaction may occur**

As shown in the diagram on the previous page, the hydrogen train also contains lithium-ion batteries. These are similar in design to those used in electric vehicles, mobile phones and other portable devices.

The chemistry of lithium-ion cells is quite complex, but in general, relies on the transfer of lithium ions between electrodes. The relevant half-equations for a lithium-ion battery are given below.

*Cathode:* CoO2 + Li+ + e- → LiCoO2

*Anode:* LiC6 → C6 + Li+ + e-

(d) Justify why fuel cells and lithium-ion batteries can both be classified as ‘galvanic’ cells.

(2 marks)

1. **both produce electrical energy from chemical energy**
2. **both involve a spontaneous redox reaction / have a positive EMF**

(e) Classify the lithium-ion battery as a ‘primary’ or ‘secondary’ cell. Justify your choice using the information provided. (2 marks)

1. **secondary**
2. **diagram states that lithium-ion cells store energy produced by fuel cells**

**which means they must be able to be charged and discharged**

One of the main advantages of replacing current diesel-powered trains with hydrogen trains is to reduce carbon dioxide emissions. The increase in atmospheric CO2(g) is a primary cause of global warming, but can also lead to other negative environmental consequences such as ocean acidification.

(f) Describe, using relevant chemical equations, how an increase in atmospheric CO2(g) has led to a decrease in the pH of our oceans.

(6 marks)

1. **Atmospheric carbon dioxide dissolves in water to produce carbonic acid**
2. **CO2(g) → CO2(aq)**

**CO2(aq) + H2O(l) → H2CO3(aq)**

**OR**

**CO2(g) + H2O(l) → H2CO3(aq)**

1. **The carbonic acid ionises to produce hydronium ions**
2. **H2CO3(aq) + H2O(l) → H3O+(aq) + HCO3-(aq)**
3. **Since pH = -log[H3O+] / since this results in an increased concentration of H3O+(aq)**
4. **This leads to a decrease in pH**

This page has been left blank intentionally

**Question 37 (18 marks)**

A group of chemistry students were comparing and contrasting two different acid-base titrations.

They set up ‘Titration A’ and ‘Titration B’ as follows;

|  |  |  |
| --- | --- | --- |
|  | **Titration A** | **Titration B** |
| Substance in conical flask | 20.00 mL of 0.10 mol L-1 NaOH(aq) | 20.00 mL of 0.10 mol L-1 NaOH(aq) |
| Substance in burette | 0.10 mol L-1 HCl(aq) | 0.10 mol L-1 CH3COOH(aq) |
| Sketch of titration curve | pH  volume of acid added (mL) | pH  \*  volume of acid added (mL) |

(a) In the table above, label the equivalence point on **both** titration curves. (2 marks)

**(1) For each graph, see arrows (should be located midpoint of vertical drop)**

(b) State the pH at the equivalence point in Titration A. Justify your answer. (3 marks)

pH

1. **7**
2. **The salt produced (NaCl) is neutral and does not hydrolyse**
3. **Therefore, the concentration of H3O+(aq) is equal to the concentration of OH-(aq)**

(c) In Titration A, state the volume of HCl(aq) that would have been added from the burette to reach the equivalence point. (Note that a calculation is not required.) (1 mark)

1. **20 mL**

(d) Would the volume of CH3COOH(aq) required to reach the equivalence point in Titration B be higher, lower or equal to the volume of HCl(aq) required in Titration A? Circle your choice below, and briefly justify your answer. (2 marks)

higher lower equal

1. **Equal volumes and concentrations used and both acids are monoprotic (therefore will require the same moles of base to neutralise)**

The students decided to investigate Titration A in more detail. Once they had successfully reached the equivalence point, they added 5 extra drops of HCl(aq).

(e) Calculate the pH of the resulting solution within the conical flask. Assume the volume of 1 drop is equal to 0.05 mL. (5 marks)

**V(HCl) = 5 x 0.05 x 10-3**

**= 0.00025 L (1)**

**n(HCl) = cV**

**= 0.1 x 0.00025**

**= 2.5 x 10-5 mol**

**= n(H+) (1)**

**c(H+) = n / V**

**= 2.5 x 10-5 / 0.04025 (1 for V)**

**= 6.211 x 10-4 mol L-1 (1)**

**pH = - log [H3O+]**

**= - log (6.211 x 10-4)**

**= 3.207 (1)**

**= 3.2**

As can be seen in the curve for Titration B, if extra CH3COOH(aq) is added after the equivalence point has been reached, the change in pH is much less pronounced. In fact, the addition of 5 extra drops of CH3COOH(aq) lowers the pH by much less than 2 units.

(f) Complete the table below, by listing the next two **compounds** with highest concentration (after water) that would be present in the conical flask for Titration B, when 5 drops of CH3COOH(aq) has been added past the equivalence point. (2 marks)

|  |  |
| --- | --- |
| **Titration A** | **Titration B** |
| 1. H2O(l) 2. NaCl(aq) 3. HCl(aq) | 1. H2O(l) 2. **NaCH3COO(aq)**      1. **CH3COOH(aq)** |

\*

Further addition of CH3COOH(aq) past the equivalence point (the region indicated by on the titration curve) continues to cause a much smaller change in pH, compared to Titration A.

(g) Justify why, upon addition of excess acid after equivalence, the change in pH in Titration B occurs to a much lesser degree than in Titration A. (3 marks)

1. **A buffer solution is formed**
2. **Due to the presence of the weak conjugate acid-base pair CH3COOH / CH3COO-**
3. **Therefore, the pH change upon addition of excess acid is not as great as in Titration A where no buffer solution is formed**

**Question 38 (18 marks)**

The Bosch reaction has applications in space travel and is utilised aboard space shuttles, since it is able to remove carbon dioxide from the air and generate clean water in the process. The reaction requires a metallic catalyst such as iron, cobalt or nickel. The equation for the Bosch reaction is given below.

CO2(g) + 2 H2(g) ⇌ 2 H2O(g) + C(s) + 90.13 kJ

Consider a reaction chamber, where the Bosch reaction has been allowed to establish equilibrium. The graph below shows the effect of imposing different changes on the system.

E1 T1 E2 T2 E3

H2O(g)

CO2(g)

H2(g)

(a) State the change imposed on the system at Time T1. (1 mark)

1. **Volume decreased / halved**

(b) Explain, in terms of the collision theory, the effect of the imposed change at Time T1 on the position of equilibrium. (4 marks)

1. **Both forward and reverse reaction rates are increased**
2. **Same amount of gas particles are in a smaller space, therefore frequency of collision is increased**
3. **Since the ratio of gas particles is 3:2, the forward reaction rate is increased more than / increases relative to the reverse reaction rate**
4. **This results in a shift to the right / the products being favoured**

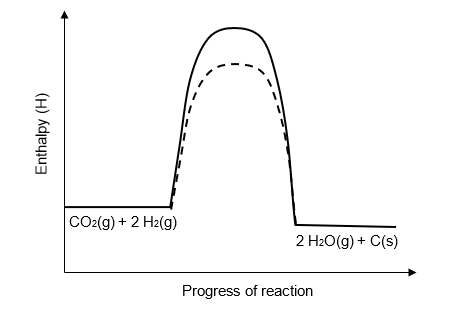
(c) State the change imposed on the system at Time T2, as well as the effect this change has on the position of equilibrium. (2 marks)

1. **Water vapour removed**
2. **Equilibrium will shift right / favour forward reaction**

(d) Complete the table below, regarding the changes that would be observed in reaction rate in response to the change imposed at Time T2. In each box of the table, write either ‘increased’, ‘decreased’ or ‘no change’. Your answers should be stated in comparison to the previous equilibrium that had been established at Time E2. (4 marks)

|  |  |  |
| --- | --- | --- |
|  | **Immediate change in rate**  (increase / decrease / no change) | **Final / overall change in rate**  (increase / decrease / no change) |
| Forward reaction | **none** | **decrease** |
| Reverse reaction | **decrease** | **decrease** |

The energy profile diagram for the Bosch reaction is shown below. The reaction has a very high activation energy and must be carried out at a temperature greater than 450 °C. It is only mildly exothermic.



(e) Comment on the likely reversibility of this reaction. (2 marks)

1. **Not likely to be reversible / reverse reaction is less likely to occur than forward reaction**
2. **Activation energy is high in the forward direction and high temperatures are required, therefore reverse reaction is unlikely to occur as reverse activation energy is even higher**

**(Marks may be awarded if students argue that the reverse reaction would potentially also occur, under conditions of high temperature, since the activation energy for the reverse is only slightly higher than forward)**

(f) Add to the energy profile diagram above, by indicating the effect of including metallic iron in the reaction system. (1 mark)

1. **See dashed line on energy profile diagram above**

(g) Explain, in terms of the collision theory, the role of the iron in this reaction. (4 marks)

1. **Provides an alternate reaction pathway**
2. **With a lower activation energy**
3. **Therefore, a greater proportion of particles can react / have collision energy greater than activation energy**
4. **Reaction rate is increased**

**(1m may be awarded for discussion of how orientation of collision can be improved by adsorption/desorption of reactants onto catalyst etc)**

**Question 39 (17 marks)**

Titratable acidity in wine is a commonly performed technique carried out during winemaking. Acid content in wine affects its colour, balance of flavour and taste. Several acids can be present in wine including tartaric, malic and citric acid but acid content in wine is reported as percentage tartaric acid (a diprotic acid).

The titration can be performed using previously standardised sodium hydroxide solution with phenolphthalein as the end point indicator. The equivalence point of this reaction occurs at pH 8.2.

(a) Explain, in terms of the equivalence point and end point, why phenolphthalein is a suitable indicator for this reaction. State the colour change observed. (4 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| phenolphthalein changes colour at a pH above 7 | **1** |
| indicating the end point of the titration | **1** |
| Colourless to pink | **1** |
| This shows the equivalence point has been reached (pH 8.2)) | **1** |
| **Total** | **4** |

(b) State three reasons why the sodium hydroxide solution must be standardised before use.

(3 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| Any three of the following   * sodium hydroxide solution will absorb (react with) carbon dioxide from the atmosphere   sodium hydroxide solid (used to make the solution)   * cannot be weighed accurately as it absorbs and dissolves in moisture from the air (deliquescent) (allow hygroscopic) * has a low molecular weight * purity not known * will absorb (react with) carbon dioxide from the atmosphere | **1–3** |
| **Total** | **3** |

Generally the titration is carried out using a pH meter, rather than phenolphthalein, as the result is “less prone to human error”.

(c) Explain why using a pH meter is a preferred method to minimise random error. (2 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| obtaining a consistent colour pink (to indicate the equivalence point) is subject to human/random error | **1** |
| compared to a pH meter which can measure the equivalence point at exactly 8.2 | **1** |
| **Total** | **2** |

A pH meter must be calibrated before use with known buffer solutions, usually pH 4.00 and 7.00. A pH 4.00 buffer can be made by dissolving 8.954 g of Na2HPO4.12H2O and 3.4023 g KH2PO4 in distilled water and making the volume up to exactly 1.000 L.

(d) Write the chemical reaction for this buffer using the HPO42– and H2PO4– ions.

(2 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| H2PO4–(aq) + H2O(ℓ) ⇌ H3O+(aq) + HPO42–(aq) |  |
| Correct species | **1** |
| Uses ⇌ | **1** |
| **Total** | **2** |

(e) Calculate the molar concentration of the Na2HPO4.12H2O in the buffer to the appropriate number of significant figures.

(4 marks)

|  |  |
| --- | --- |
| **Description** | **Marks** |
| M(Na2HPO4.12H2O) = 358.15 g mol–1 | **1** |
| n(Na2HPO4.12H2O) = 8.954 / 358.15 = 0.02500 | **1** |
| [Na2HPO4.12H2O] = 0.02500/ 1.000 = 0.02500 mol L–1 | **1** |
| Answer to 4 sf | **1** |
| **Total** | **4** |

(f) Define ‘buffering capacity’. (2 marks)

Buffer capacity is a measure of the effectiveness of a buffer solution to resist a change in pH when either a strong acid or strong base is added.

**End of questions**

Supplementary page

Question number:

Supplementary page

Question number:

Supplementary page

Question number:

**END OF QUESTIONS**

Spare answer page

Question number: \_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_\_