

NANYANG JUNIOR COLLEGE JC 2 PRELIMINARY EXAMINATION Higher 2

CHEMISTRY

9729/01

Paper 1 Multiple Choice

24 September 2018 1 hour

Additional Materials: Multiple Choice Answer Sheet Data Booklet

READ THESE INSTRUCTIONS FIRST

Write in soft pencil.

Do not use staples, paper clips, glue or correction fluid. Write your name, class and tutor's name on the Answer Sheet in the spaces provided unless this has been done for you.

There are **thirty** questions on this paper. Answer **all** questions. For each question there are four possible answers **A**, **B**, **C** and **D**.

Choose the **one** you consider correct and record your choice in **soft pencil** on the separate Answer Sheet.

Read the instructions on the Answer Sheet very carefully.

Each correct answer will score one mark. A mark will not be deducted for a wrong answer. Any rough working should be done in this booklet. The use of an approved scientific calculator is expected, where appropriate.

This document consists of **16** printed pages and **0** blank page.

For each question there are four possible answers, **A**, **B**, **C** and **D**. Choose the **one** you consider to be correct.

- 1 Which of the following contains the smallest number of stated atoms at r.t.p.?
 - **A** oxygen atoms in 48 dm³ of NO_2
 - **B** sulfur atoms in 48 dm³ of SO₂
 - **C** carbon atoms in 44 g of CO₂
 - **D** nitrogen atoms in 44 g of N_2O
- **2** An isolated gaseous species has paired electrons in at least one of its 3d orbitals and a fully filled 4s subshell.

What could be the identity of the species?

A Cu B Fe^{3+} C Ni^{2+} D Sr^{2+}

- 3 Which one of the following species is **not** planar?
 - A BrF₃
 - **B** ICl_4^-
 - \mathbf{C} PCl₄⁺
 - **D** XeF₄
- **4** Hydrazine, N₂H₄, and hydrogen peroxide, H₂O₂, are both used as rocket propellants because they can produce large volumes of hot gases from a small volume of liquid.

Which of the following statements about these two compounds is correct?

- 1 The bond angle in N_2H_4 is larger than that in H_2O_2 .
- 2 The N–H bond is shorter than the O–H bond.
- 3 Hydrazine forms stronger intermolecular hydrogen bonds than hydrogen peroxide.
- 4 There are σ bonds formed by sp³ s orbital overlap in both molecules.
- **A** 1 and 4
- **B** 2 and 3
- **C** 1, 2 and 4
- **D** 1, 3 and 4

- **5** Which factor is the most significant in explaining the non-ideal behaviour of the gases present in the reaction chamber in the Haber process?
 - A strong bonds between the atoms in the molecule
 - **B** the presence of a catalyst
 - **C** the high temperature of 450 °C
 - **D** the high pressure of 150 atm
- **6** A student mixed 25.0 cm³ of 3.00 mol dm⁻³ hydrochloric acid with an equal volume of 6.00 mol dm⁻³ sodium hydroxide. The initial temperature of both solutions was 15.0 °C. The maximum temperature recorded was 24.5 °C. It was found that 15% of the heat produced during the experiment was lost to the surrounding.

Using these results, what is the enthalpy change of neutralisation?

- A -15.6 kJ mol⁻¹
- B –22.5 kJ mol⁻¹
- C -31.1 kJ mol⁻¹
- **D** –57.3 kJ mol⁻¹
- 7 A student calculated the lattice energy for calcium oxide using the data in the table below, and relevant data from the *Data Booklet*.

energy change	value / kJ mol ^{_1}
standard enthalpy change of atomisation of calcium	+178
first electron affinity for oxygen	-141
second electron affinity for oxygen	+798
standard enthalpy change of formation of calcium oxide	-635

However, the value calculated by the student for the lattice energy was **more** exothermic than the correct value.

Which error could have been made in the calculation?

- A omitting the second electron affinity for oxygen
- **B** omitting the standard enthalpy change of formation of calcium oxide
- **C** using the 1st and 2nd ionisation energies of magnesium instead of calcium
- **D** using the standard enthalpy change of combustion of calcium rather than the standard enthalpy change of formation of calcium oxide

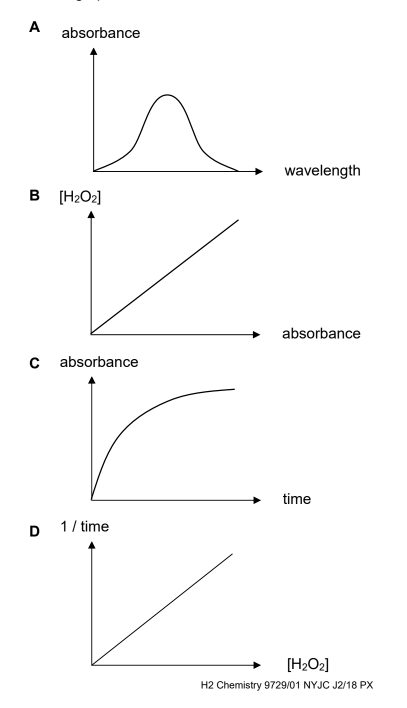
8 A student investigates the kinetics of the following reaction by using a spectrometer. The spectrometer is able to measure the concentration of iodine by measuring the absorbance of the solution.

$$H_2O_2(aq) + 2H^+(aq) + 2I^-(aq) \rightarrow I_2(aq) + 2H_2O(I)$$

The time taken for the absorbance of the reaction mixture to reach a fixed value is measured over a range of hydrogen peroxide concentrations.

Based on the data obtained, the student correctly drew a graph to prove that order of reaction is one with respect to H_2O_2 .

Which graph did the student draw?

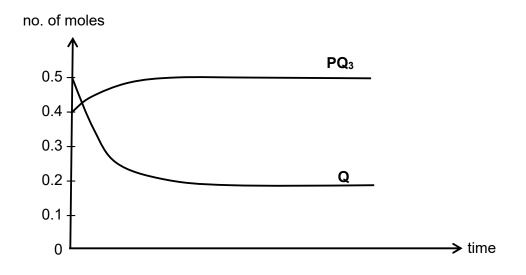


9 The decomposition of hydrogen peroxide follows first order kinetics. A certain solution of hydrogen peroxide undergoes complete decomposition to liberate 96 cm³ of oxygen gas. It is found that at 25 °C, 48 cm³ of oxygen was collected in 35 min.

How long will it take for 80 cm³ of the gas to be produced?

- A 87.5 min
- **B** 90.5 min
- **C** 97.5 min
- **D** 105 min
- **10** The system containing P, Q and PQ₃ is allowed to reach equilibrium in a 5 dm³ vessel at a temperature of 1000 K.

The diagram below shows the change in number of moles of PQ_3 and Q with time. The initial number of moles of P was 0.2.



What is the equilibrium constant K_c for the reaction?

$$\textbf{A} \quad \frac{0.5}{0.1 \times (0.2)^3} \qquad \textbf{B} \quad \frac{0.5}{0.2 \times (0.2)^3} \qquad \textbf{C} \quad \frac{0.5 \times 5^3}{0.1 \times (0.2)^3} \qquad \textbf{D} \quad \frac{0.5 \times 5^3}{0.2 \times (0.2)^3}$$

- **11** Which statement is correct about a reaction for which the equilibrium constant is independent of temperature?
 - 1 The rate constants for the forward and reverse reactions are both independent of temperature.
 - 2 Temperature has no effect on the position of equilibrium.
 - 3 The forward and reverse reactions have equal activation energies.
 - 4 The entropy change is zero.
 - **A** 1 and 2
 - **B** 2 and 3
 - **C** 3 and 4
 - **D** 2, 3 and 4
- **12** The two simplest carboxylic acids are formic acid and acetic acid. Formic acid is present in the venom of ant and bee stings while acetic acid is the major characterising component of vinegar. The pK_a values of the two acids is given in the table below.

Acid	Formula	р <i>К</i> а
Formic acid	НСООН	3.74
Acetic acid	CH₃COOH	4.76

Which solution has the lowest pH?

- A 100 cm^3 of 0.10 mol dm⁻³ acetic acid
- **B** 100 cm³ of 0.10 mol dm⁻³ sodium acetate
- **C** 100 cm³ of 1.0 mol dm⁻³ formic acid and 50 cm³ of 1.0 mol dm⁻³ aqueous sodium hydroxide
- ${\rm D}~$ 100 ${\rm cm}^3$ of 1.0 mol dm^{-3} formic acid and 75 ${\rm cm}^3$ of 1.0 mol dm^{-3} aqueous sodium hydroxide

13 Deuterium oxide, D₂O consists of an oxygen atom bonded to two atoms of the hydrogen isotope, deuterium ²₁D. It is known as 'heavy water' and is used for research into chemical reactions.

Like H₂O, pure D₂O is weakly ionised.

2D₂O Ý D₃O⁺ + OD⁻

For D₂O, we use the terms K_D instead of K_w and pD instead of pH. At 25 °C, pK_D = 14.8

Which statements about D₂O at 25 °C is correct?

- 1 pD of D₂O is 7.4
- 2 D₂O is not a neutral liquid.
- 3 D₂O dissociates to a smaller extent than H₂O.
- A 1 only
- B 3 only
- **C** 1 and 3
- **D** 2 and 3
- **14** W, X, Y and Z are four consecutive elements in Period 3 but not necessarily in the order presented.
 - Chloride of **W** dissolves in water and turns blue litmus red.
 - X is a good conductor of electricity but is insoluble in water.
 - Oxide of **Y** has the highest melting point.
 - **Z** has the highest first ionisation energy and largest ionic radius.

Which of the following is the correct sequence of the four elements in order of increasing atomic number?

- A X, Y, Z, W
- B X, Y, W, Z
- C Y, X, W, Z
- D Y, X, Z, W

15 M is a Group 2 metal. The carbonate of **M** decomposes when heated in a Bunsen flame to give carbon dioxide and a white solid residue as the only products. This white solid residue is sparingly soluble in water. Even when large amounts of the solid residue are added to water the pH of the saturated solution is less than that of limewater.

What could be the identity of M?

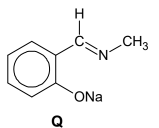
- A magnesium
- B calcium
- C strontium
- **D** barium
- **16** A white powder is a mixture of sodium chloride and sodium iodide. It is dissolved in water in a test-tube. Excess aqueous silver nitrate is added to the test-tube. A precipitate, X, is observed.

Excess concentrated ammonia is then added to the test-tube containing X. After the test-tube has been shaken, a precipitate, Y, is observed.

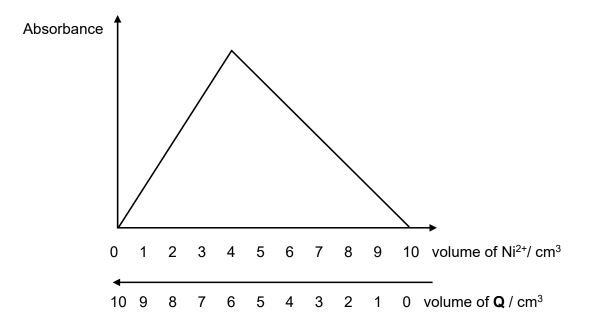
Which statement about X or Y is correct?

- **A** X is a pure white colour.
- **B** X is pure silver iodide.
- **C** Y is pure silver chloride.
- **D** Y is yellow.
- **17** Which of the following statements is correct?
 - 1 Enthalpy change of vaporisation of halogens increases down the group.
 - 2 Bond energy of hydrogen halides increases down the group.
 - 3 Oxidising power of halogens increases down the group.
 - 4 Thermal stability of hydrogen halides increases down the group.
 - A 1 only
 - **B** 1 and 3
 - **C** 2 and 4
 - **D** 3 and 4

18 The complex of nickel with ligand **Q** (shown below) is thermochromic, being coloured red at room temperature but changing to yellow-green when heated to 170 °C.



The following graph was obtained when the colour intensities of mixtures of a 4.0×10^{-3} mol dm⁻³ solution of **Q** and a 3.0×10^{-3} mol dm⁻³ solution of nickel(II) chloride were measured using a colorimeter at room temperature.



Which of the following statements is correct for the nickel(II) complex ion?

- **A** The complex ion absorbs red light at room temperature.
- **B** The co-ordination number of nickel(II) complex is 4.
- **C** The overall charge of the nickel(II) complex is 2+.
- **D** The shape of the nickel(II) complex ion is linear.

19 If a molecule contains two non-identical chiral carbon atoms, four optical isomers exist.

How many isomers are there with

- molecular formula C₇H₁₄O and
- a five-membered ring **and**
- a tertiary alcohol group?

A 7 **B** 8 **C** 9 **D** 10

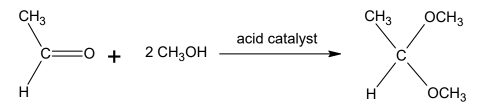
20 Which of the following trends are incorrect?

- 1 The boiling points of alcohols with the same molecular formula increases from primary to secondary to tertiary.
- 2 The pH values of the aqueous solutions increases from CH₃CH₂COCI to CH₃CH₂COOH to CH₃CH₂CONH₂ to CH₃CHCICH₂NH₂.
- 3 The ease of hydrolysis of the chlorine atoms decreases from chlorobenzene to chloroethane to ethanoyl chloride.
- 4 The pK_b values increases from ethanoate ion to phenoxide ion to ethoxide ion.
- A 2 only
- **B** 1 and 3 only
- C 2 and 4 only
- **D** 1, 3 and 4 only

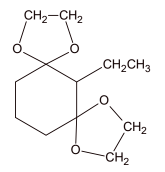
21 Alkenes undergo catalytic oxidation to produce aldehydes and ketones, for example

$$H_2C$$
 \longrightarrow CH_2 $+$ $1/2 O_2$ $\xrightarrow{catalyst}$ CH_3CHO

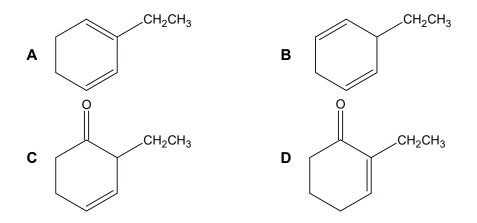
Acetals (molecules that contain 2 –OR groups bonded to the same carbon) can be formed from aldehydes and ketones in an acid catalysed process, for example



Compound **W** undergoes catalytic oxidation, followed by an acid catalysed reaction with HOCH₂CH₂OH to produce the following as the only product.



Which of the following best represents the structure of compound W?



22 The molecular formula of compound **X** is $C_5H_{12}O$.

Compound X:

- · reacts with alkaline aqueous iodine
- can be dehydrated to form two alkenes only.

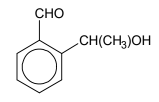
What could be the identity of compound X?

- **A** $CH_3CH_2CH(CH_3)CH_2OH$
- **B** $(CH_3)_2C(OH)CH_2CH_3$
- C (CH₃)₂CHCH(OH)CH₃
- **D** $CH_3CH_2CH_2CH(OH)CH_3$
- **23** $H_2NNHC_6H_3(NO_2)_2$ is the structural formula of 2,4-DNPH.

Many but not all, organic reactions need to be heated before reaction occurs.

Which reaction cannot occur at a good rate at room temperature (20 °C)?

- $\textbf{A} \quad CH_3OH + PCI_5 \rightarrow CH_3CI + POCI_3 + HCI$
- $\textbf{B} \quad CH_3CH_2OH + KBr \rightarrow CH_3CH_2Br + KOH$
- $\textbf{C} \quad CH_3COCI + CH_3CH_2OH \rightarrow CH_3COOCH_2CH_3 + HCI$
- **D** $(CH_3)_2CO + H_2NNHC_6H_3(NO_2)_2 \rightarrow (CH_3)_2C=NNHC_6H_3(NO_2)_2 + H_2O$
- 24 Compound X is heated under reflux with an excess of acidified K₂Cr₂O₇ to form compound Y.

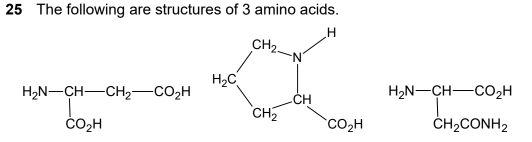


compound X

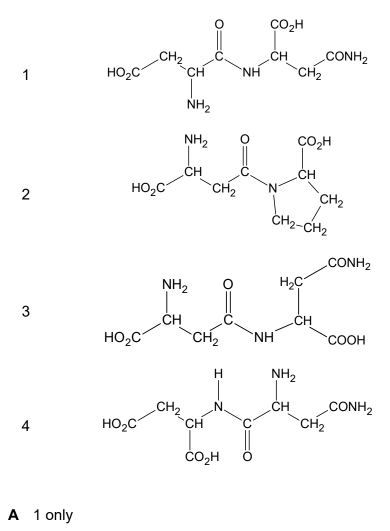
Both **X** and **Y** are separately warmed with Fehling's solution and the observations noted.

What are the observations?

- **A** Both **X** and **Y** give a red precipitate.
- **B** Only **X** gives a red precipitate.
- **C** Only **Y** gives a red precipitate.
- **D** Neither **X** nor **Y** gives a red precipitate.



Which of the following represents dipeptides formed from these amino acids?

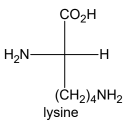


B 1 and 4 only

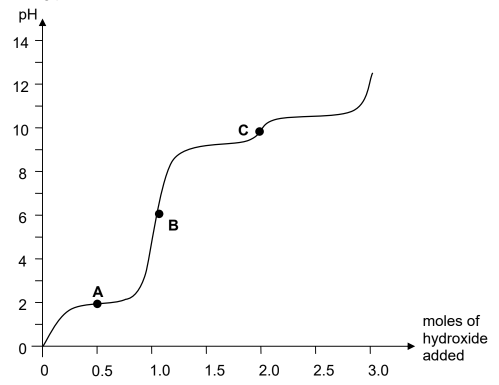
C 3 and 4 only

D All are correct

26 Lysine is an essential amino acid found in the body. It has three pK_a values associated with it: 2.2, 9.0 and 10.5



When one mole of protonated lysine was titrated against hydroxide ions, the following pH curve is obtained:



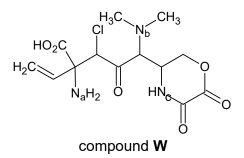
Which of the following statements are true with respect to the curve above?

- 1 The α -amino group has a p K_a value of 10.5.
- 2 Equal amounts of $H_3N^+CH(CO_2H)(CH_2)_4NH_3^+$ and $H_3N^+CH(CO_2H)(CH_2)_4NH_2$ are present at point **A**.
- 3 The major species present at point **C** has no net charge.
- 4 The major species present at point **B** will migrate towards the cathode of an electrolytic cell.
- A 1 and 3 only
- B 2 and 4 only
- C 3 and 4 only
- D All are correct

27 A carboxylic acid, **P**, has no possible chain isomers. It reacts with an alcohol, **Q**, that has only one other positional isomer.

What could be the ester formed from a reaction between P and Q?

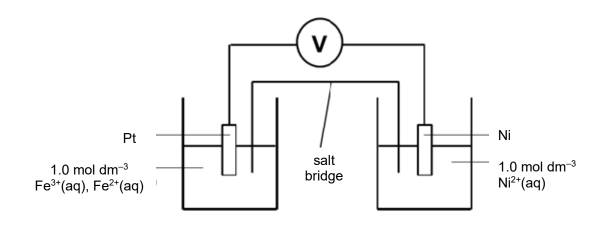
- A butyl propanoate
- B ethyl butanoate
- **C** pentyl ethanoate
- D propyl pentanoate
- 28 Which of the following statements about compound W is correct?



- A In the gaseous phase, N_b is a stronger Lewis base than N_a but in the aqueous phase, N_b is a weaker Bronsted-Lowry base than N_a .
- **B** When 96 dm³ of H₂ gas was reacted with one mole of compound **W**, followed by excess sodium metal, at room conditions, the gas volume expanded by 48 dm^3 .
- **C** The reduction of **W** by LiAlH₄ will cause the oxidation state of any carbon involved in the reduction to decrease from +3 to -1.
- **D W** will require 3 moles of NaOH(aq) for complete reaction if the reaction is to take place with heating.
- 29 Which of the following changes represent an oxidation process?
 - A CO₂ to C₂O₄²⁻
 - **B** CrO₄²⁻ to Cr₂O₇²⁻
 - \mathbf{C} Br₂ to BrF₃
 - $\boldsymbol{\mathsf{D}}\quad Cl_2 \text{ to } ICl_3$

30 The use of Data booklet is relevant to this question.

A Fe³⁺/Fe²⁺ half-cell was connected to a Ni²⁺/Ni half-cell as shown in the diagram below under standard conditions.



Which of the following statements is correct?

- 1 The solution in the Fe^{3+}/Fe^{2+} half-cell turns red-brown.
- 2 The cathode increases in size.
- 3 The electron flows from Fe^{3+}/Fe^{2+} half-cell to the Ni²⁺/Ni half-cell.
- 4 The standard cell potential is +1.02 V.
- A 4 only
- **B** 1 and 3
- **C** 2 and 3
- **D** 1, 3 and 4

Nanyang JC J2 Preliminary Exam 2018 H2 Chemistry 9729/01 Paper 1 MCQ Answers and Comments

Qn	Ans										
1	С	6	С	11	В	16	D	21	D	26	С
2	D	7	С	12	Α	17	Α	22	С	27	Α
3	С	8	D	13	С	18	В	23	В	28	Α
4	Α	9	В	14	С	19	С	24	D	29	С
5	D	10	С	15	Α	20	D	25	В	30	Α

1 C

A: 2 mol of NO₂ gives 4 mol of O atoms B: 2 mol of SO₂ gives 2 mol of S atoms C: 1 mol of CO₂ gives 1 mol of C atoms D: 1 mol of N₂O gives 2 mol of N atoms

2 D

Cu: $1s^2 2s^2 2p^6 3s^2 3p^6 3d^{10}4s^1$ Fe³⁺: $1s^2 2s^2 2p^6 3s^2 3p^6 3d^5$ Ni²⁺: $1s^2 2s^2 2p^6 3s^2 3p^6 3d^8$ Sr²⁺: $1s^2 2s^2 2p^6 3s^2 3p^6 3d^{10} 4s^2 4p^6$

3 C

- A BrF₃: T-shaped (3bp, 2 lp)
- **B** ICl₄ : square planar (4bp, 2 lp)
- **C** PCl_4^+ : tetrahedral (4 bp, 0 lp)
- **D** XeF₄: square planar (4bp, 2 lp)

4 A

(1 and 4 are correct)

- 1 True, bond angle in N_2H_4 is 107^o while bond angle in H_2O_2 is 105^o
- 2 False, the N–H bond is *longer* than the O–H bond as the O–H covalent bond in more polar. Students can also check the DB to compare the 2 bond energies.
- 3 False, hydrazine forms *weaker* intermolecular hydrogen bonds than hydrogen peroxide as N is less electronegative than O.
- 4 True, both N and O are sp³ hybridised, and form σ bonds by overlapping with the s orbital of H.
- 5 D

Gases deviate from non-ideal behaviour at low temperature or high pressure hence option D is true.

Explanation for option D: At high pressure, volume of container decreases hence volume of gas particles is more significant compared to volume of container, therefore gas behave less ideally.

Note for option A, it should be the significant <u>intermolecular</u> hydrogen bonds between molecules that result in deviation from ideal gas behaviour (not the <u>intramolecular</u> covalent bonds between atoms).

6 C

amount of water formed

$$= \frac{25.0}{1000} \times 3.00 = 0.07500 \text{ mol}$$

heat produced

= $(25.0 + 25.0) \times 4.18 \times 9.5 \times \frac{100}{85}$ = 2335 J mol^{-1} $\Delta H_n = -\frac{2335 \times 10^{-3}}{0.07500}$ = -31.14= $-31.1 \text{ kJ mol}^{-1}$ **7 C**

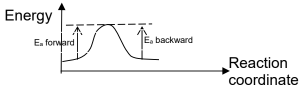
 $\Delta H^{\Theta}_{lattice energy} CaO$

= $-(\Delta H^{e}_{atomisation} \text{ Ca} + 1^{st} \text{ and } 2^{nd} \text{ ionisation}$ energies of calcium + $\Delta H^{e}_{atomisation} \text{ O}$ + first and second electron affinity for oxygen) + $\Delta H^{e}_{formation} \text{ CaO}$ Option A: Less exothermic Option B: Less exothermic Option C: More exothermic since 1st and 2nd POE if ΔH is zero. ionisation energies of magnesium are more endothermic than calcium Option D: No change since standard enthalpy change of combustion of calcium is equal to the standard enthalpy change of Energy formation of calcium oxide. E₂ for 8 D 1/ time presents rate of reaction Since first order wrt H₂O₂, hence $[H_2O_2] \alpha 1 / time$ 9 В pressure is provided. $\frac{96-80}{96} = \left(\frac{1}{2}\right)^{t_{35}}$ 40 . t = 90.5 min 10 C pH > 7. $[Q]_{eqm} = 0.2 / 5 \text{ mol } dm^{-3}$ $[PQ_3]_{eqm} = 0.5 / 5 \text{ mol dm}^{-3}$ Initial P = 0.2 mol = 3.74Since P and PQ₃ have the same mole ratio in the equation, an acidic buffer. $|change in P| = |change in PQ_3| = 0.1$ (read off from graph) 13 C Alternatively, you can also find change in P 1: $K_D = 10^{-14.8}$ = 1/3 change in Q. Hence $[P]_{eqm} = (0.2 - 0.1) = 0.1 / 5 \text{ mol dm}^{-3}$ pD = 7.4 $K_{c} = \frac{[PQ_{3}]}{[Q]^{3}[P]} = \frac{(\frac{0.5}{5})}{(\frac{0.2}{5})^{3}(\frac{0.1}{5})} = \frac{0.5 \times 5^{3}}{0.2^{3} \times 0.1}$ 11 B 14 C If equilibrium constant is independent of temperature, it means when temperature changes, neither forward nor backward reaction is favoured and this is only possible if ΔH is zero. 1. All rate constants are affected by Arrhenius

temperature according to equation: $k = A^{-Ea/RT}$.

2. Temperature would have no effect on

3. When ΔH is zero, activation energies for forward and backward reaction are equal as shown in the energy profile diagram below:



We cannot deduce any information about ΔS unless some information about how value of equilibrium constant changes with

12 A
A pH of 0.010 mol dm^{-3} acetic acid = - lg
$(\sqrt{10^{-4.76}(0.10)}) = 2.88$
B The solution contains a basic salt hence

C A buffer at maximum buffering capacity of

formic acid and its salt is formed. pH = pKa

D As more hydroxide is added compared to option C, pH of D > C but pH D < 7 as it is

 $[D^+] = \sqrt{10^{-14.8}} = 3.98 \times 10^{-8}$

2: Since $[H^+]=[OH^-]$, D₂O is netural.

3: $K_D = 10^{-14.8} < K_w = 10^{-14}$ hence D_2O dissociates to a smaller extent than H₂O.

NYJC 2018 H2 Chemistry 9729 P1 worked solns.doc 20-Aug-2018 10:37 AM

Chloride of **W** dissolves in water and turns blue litmus red: **W** is not Na (NaCl is neutral)

X is a good conductor of electricity but insoluble in water. **X** can be Mg or Al (However, since **Y** is Mg, **X** must be Al)

Oxide of **Y** has the highest melting point: MgO (**Y** is Mg)

Z has the highest first ionisation energy and largest ionic radius. (**Z** is P)

Since the four elements in Period 3 must be consecutive, **W** must be Si.

Hence the order is Y, X, W, Z.

(Mg, Al, Si, P)

15 A

 $\text{MCO}_3 \rightarrow \text{MO} + \text{CO}_2$

Down the group, solubility of oxides increases and pH of the solution increases. MgO + H₂O \checkmark Mg(OH)₂

BaO + H₂O \rightarrow Ba(OH)₂

MgO dissolves sparingly in water to form weak base Mg(OH)₂ whereas BaO dissolves readily in water to form strong base, Ba(OH)₂.

Since pH of the saturated solution is less than that of limewater, $Ca(OH)_2$, **M** must be Mg.

16 D

Only AgCl dissolves in conc NH_3 , AgI remains insoluble in conc NH_3 . Hence Y is AgI(s) – yellow ppt.

17 A

Enthalpy change of vaporisation is inversely proportional to boiling point.

Halogens have simple molecular structure with weak instantaneous dipole-induced dipole forces (id-id) between molecules. As the number of electrons or size of electron cloud increases, the electron cloud gets more polarised, id-id forces increases down the group. More energy required to overcome id-id forces. Boiling point increases and hence ΔH_{vap} decreases down the group.

Thermal stability of HX <u>decreases</u> down group 17 due to <u>decreasing</u> bond strength of H–X (<u>increasing</u> H–X bond length)

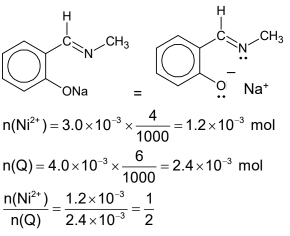
Electrode reaction	E ⁺ / V
F₂ + 2e⁻ ⊻́ 2F⁻	+2.87
Cl₂ + 2e⁻ ⊻́ 2Cl⁻	+1.36
Br₂ + 2e⁻ ⊻́ 2Br⁻	+1.07
I₂ + 2e⁻ ⊻́ 2I⁻	+0.54

 $E^{\bullet}(X_2|X^-)$ becomes less positive down the group, oxidising power of elements <u>decreases</u> down the group.

18 B

A The red complex ion absorbs <u>complementary colour (green)</u> at room temperature.

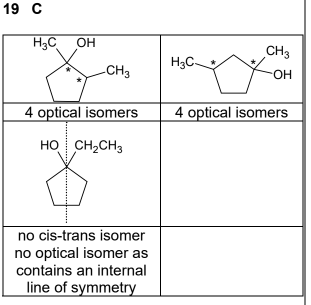
В



Since \mathbf{Q} is a bidentate ligand (lone pair on N and lone pair on O⁻ for dative bonding), the co-ordination number of nickel(II) complex is 4.

C The overall charge of the nickel(II) complex is 0 as there are 2 ligands of **Q** to one nickel(II) ion.

D The shape of the nickel(II) complex ion is not linear as the co-ordination number is not 2.



20 D (1,3 and 4) **Option 1 (incorrect)**

The surface area decreases from primary (elongated) to secondary to tertiary alcohol (spherical). Hence strength of id-id forces of attraction decreases from primary to secondary to tertiary.

Option 2 (correct)

CH₃CH₂COCI hydrolyses in water to produce a strong acid, HCI. Hence the pH of its aqueous solution is the lowest. CH₃CH₂CONH₂ contains an amide functional group which is neutral. $CH_3CH(CI)CH_2NH_2$ contains an amine functional group which is basic and hence its pH value is the highest.

Option 3 (incorrect)

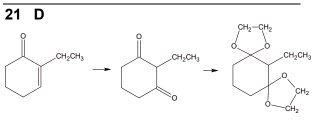
The ease of hydrolysis of the chlorine atoms decreases increases from chlorobenzene to chloroethane to ethanoyl chloride.

Option 4 (incorrect)

Acidity decreases from ethanoic acid to phenol to ethanol.

Ethanoic acid is the strongest among the three, hence its conjugate base, ethanoate ion will be the weakest, with the largest pK_b value.

Ethanol is the weakest among the three, hence its conjugate base, ethoxide ion will be the strongest, with the smallest pK_b value.

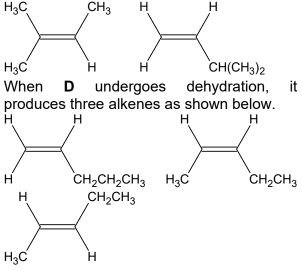


22 C

A and **B** do not contain methyl alcohol, hence will not be able to react with alkaline aqueous iodine.

C, $(CH_3)_2CHCH(OH)CH_3$ and **D**, $(CH_3CH_2CH_2CH(OH)CH_3)$ contain methyl alcohol, hence will react with alkaline aqueous iodine.

When **C** undergoes dehydration, it produces two alkenes only as shown below.



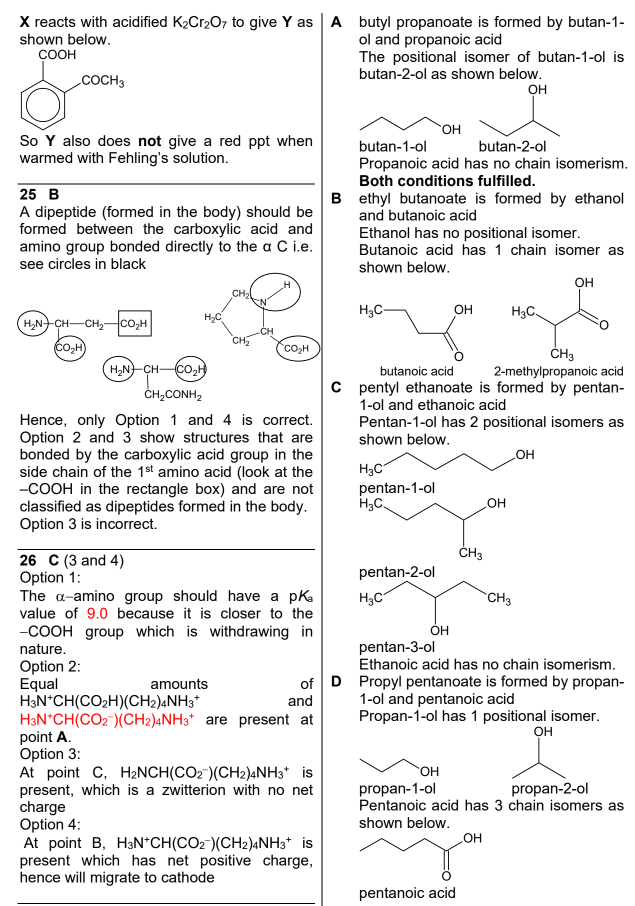
23 B

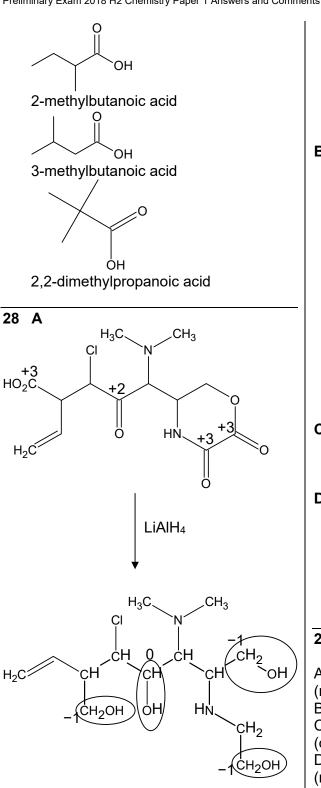
All can occur at a good rate at room temperature (20 °C) except

 $CH_3CH_2OH + KBr \rightarrow CH_3CH_2Br + KOH$ as conc. H_2SO_4 needs to be added to react with KBr with heating to produce HBr which will then reacts with CH_3CH_2OH to produce CH_3CH_2OH .

24 D

X contains an aromatic aldehyde that does **not** give a red ppt when warmed with Fehling's solution.





A (Correct) As a Lewis base, N_b has more electron-donating grps than N_a which increases the electron density on the N atom and hence the availability of the lone pair of electrons for donation to a Lewis acid. So N_b is a stronger Lewis base than N_a . As a Bronsted-Lowry base, N_b experiences steric hindrance from the presence of 3 bulky alkyl grps, hence protons from the aq soln will have difficulty going near the lone pairs to form dative bond. So N_b is a weaker Bronsted-Lowry base than N_a .

B (Incorrect) When 96 dm³ of H₂ gas was reacted with one mole of compound W, followed by excess sodium metal, at room conditions, the gas volume expanded contracted by 4824 dm³.

96 dm³ is equivalent to 4 mol of H₂ gas. Only the alkene and ketone function grp will be reduced by H₂(g) <u>using up 2 out of</u> the 4 mol of H₂(g) present.

Alkene will be reduced to alkane while ketone will be reduced to 2° alcohol. The 2° alcohol and –COOH grp present will then react with Na to produce 1 mol of $H_2(g)$.

The final volume of $H_2(g)$ is 72 dm³.

The gas volume contracted by 24 dm³.

- **C** Incorrect. The numbers written beside the C atom (as shown above) shows the oxidation state of the carbon before and after the reduction by LiAlH₄.
- **D** Functional groups in **W** that will react with NaOH with heating are –COOH, RCI, amide and ester. So will need 4 mol of NaOH.

Functional groups in W that will react with NaOH without heating is –COOH. So will need 1 mol of NaOH.

29 C

A: O.S. of C changes from +4 to +3 (reduction) B: O.S. of Cr remains unchanged at +6

C: O.S. of Br changes from 0 to +3 (oxidation)

D: O.S. of CI changes from 0 to -1 (reduction)

30 A

 $\begin{array}{ll} \mathsf{Fe}^{3+} + e \rightarrow \mathsf{Fe}^{2+} & +0.77 \mathsf{V} \rightarrow +\mathsf{ve} \rightarrow [\mathsf{R}] \\ \mathsf{Ni} + 2e \rightarrow \mathsf{Ni}^{2+} & -0.25 \mathsf{V} \rightarrow -\mathsf{ve} \rightarrow [\mathsf{O}] \end{array}$

 Fe^{3+} is yellow and Fe^{2+} is pale green.

Reduction occurs at the Fe^{3+}/Fe^{2+} half-cell.

 $E_{cell} = +0.77 - (-0.25) = +1.02 V.$

Electrons flow from the anode (Ni²⁺/Ni half-cell) to the cathode (Fe³⁺/Fe²⁺ half-cell).

Pt cathode size remains unchanged.



NANYANG JUNIOR COLLEGE JC 2 PRELIMINARY EXAMINATION Higher 2

CANDIDATE NAME

CLASS

TUTOR'S NAME

CHEMISTRY

Paper 2 Structured

Candidates answer on the Question Paper

Additional Materials: Data Booklet

READ THESE INSTRUCTIONS FIRST

Write your name and class on all the work you hand in. 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. A Data Booklet is provided.

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 Exa Use	aminer's
1	/24
2	/11
3	/9
4	/13
5	/18
Total	/75

9729/02

2 hours

11 September 2018

2

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

1(a) An unknown sample was found to contain the anions, Cl⁻, ClO₃⁻ and NO₃⁻. A student weighed a sample into a beaker and recorded the following data.

mass of beaker and sample / g	68.962
mass of empty beaker / g	67.620

The sample was dissolved and diluted in a 250 cm³ volumetric flask to obtain solution L.

In **experiment 1**, a 50 cm³ portion of solution **L** was reacted with excess silver nitrate solution. The AgCl precipitated was transferred onto a dry filter paper and was placed under an infra-red lamp. The dry AgCl precipitate was weighed and the following data was obtained.

Experiment 1

mass of dry filter paper and AgCl / g	0.737
mass of dry filter paper / g	0.620

In **experiment 2**, a gas was bubbled into another 50 cm³ portion of solution **L** to convert ClO_3^- to Cl^- before the addition of excess silver nitrate solution. The AgCl precipitated was also dried and weighed. The following data was obtained.

Experiment 2

mass of dry filter paper and AgCl / g	0.799
mass of dry filter paper / g	0.651

(i) Write the half equation for the reduction of CIO_3^- to CI^- .

.....[1]

(ii) Determine the mass of CI^- in 50 cm³ of solution L.

[1]

(iii) Determine the mass of Cl^- converted from ClO_3^- in experiment 2.

(iv)	Hence, determine th	e percentage mass	s of ClO_3^- in the unknown	sample
------	---------------------	-------------------	-------------------------------	--------

[3]

For Examiner's

- (v) The $E_{\mathbb{Q}}$ (ClO₃⁻/Cl⁻) has a value of +1.47 V. From the list of standard electrode potentials in the *Data Booklet*, identify a gas that would reduce ClO₃⁻ to Cl⁻. Explain your answer.
 -[2]
- (b) (i) An aqueous solution of HCl has a density of 1.15 g cm⁻³ and is 30% by mass of HCl.

Calculate the concentration in mol dm⁻³ of this solution of HCl.

[2]

(ii) Calculate the volume of this solution required to prepare 5 dm³ of 0.20 mol dm⁻³ HCl by dilution with water.

(c) The gelatin silver process is the photographic process used with black-and-white films. The following information pertains to the process of taking photographs and developing films.

Taking photographs

- A 35 mm cartridge of black-and-white print film contains a long strip of plastic that has layered coatings on each side.
- On the front side of the film, the layers are made of gelatin which contain grains of silver chloride crystals.
- When the shutter of the camera is opened for a fraction of a second to allow the film to be exposed to light, these crystals undergo decomposition thereby producing an image on the film.

Developing films

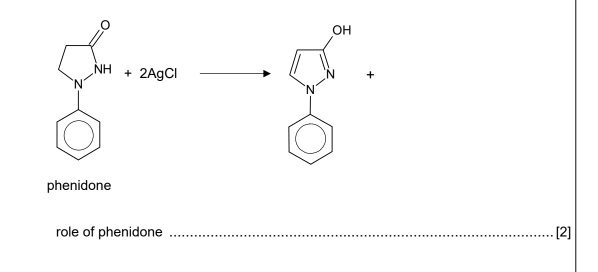
- After the photographs have been taken, the film is developed in a dark room under a light source that emits low energy light.
- Firstly, the film is soaked in water before adding phenidone. Phenidone makes the image more visible by reacting with the exposed silver chloride crystals to produce silver atoms and two other by-products.
- This reaction can only proceed at high pH.
- After some time, the reaction will then be quenched.
- Finally, the film will be soaked in ammonium thiosulfate, (NH₄)₂S₂O₃, which is used as a fixer to make the image permanent and light resistant. This is done through the reaction between the unexposed silver chloride crystals and the fixer.
- (i) Write the balanced equation for the decomposition of silver chloride crystals when it is exposed to light.

.....[1]

(ii) Suggest a suitable colour of the light source that is used in a dark room.

.....[1]

(iii) Complete the equation for the reaction between the developing agent, phenidone, and the exposed silver chloride crystals. Hence, state the role of phenidone in this reaction.



(iv)	Suggest a suitable reagent, other than excess cold water, that can be used to quench the development of the film. Explain.
	[2]
(v)	When the non-exposed silver chloride crystals react with the fixer, $(NH_4)_2S_2O_3$, a silver complex compound M is formed together with a chloride salt, N . Both M and N have the same cation. The silver-containing complex ion has a coordination number of 2 and is chlorine-free.
	Suggest the formulae of compounds M and N .
	compound M :[2]
(vi)	State the shape of the silver-containing ion in compound ${f M}$.
	[1]
(vii)	Complete the electronic configuration of silver in compound ${\bf M}$. Hence, deduce the colour of compound ${\bf M}$.
	compound M [Ar]3d ¹⁰ 4s ²
	[2]
(viii)	Explain why the fixer, $(NH_4)_2S_2O_3$, is able to make the image permanent and light resistant on the film.
	[1]
	[Total: 24]

- 2 Dinitrogen tetroxide, commonly referred to as nitrogen tetroxide, is the chemical compound N₂O₄. It is a useful reagent in chemical synthesis.
- (a) Colourless N_2O_4 readily dissociates to form brown NO_2 and the following equilibrium is reached fairly quickly in the gaseous phase.

 $N_2O_4(g)$ Ý $2NO_2(g)$ $\Delta H > 0$

(i) When 4.60 g of N_2O_4 is placed in an evacuated 1.48 dm³ flask at 27 °C, the equilibrium pressure is 1 atm.

Calculate the value of K_p at 27 °C.

(ii)	Describe and explain what you would observe when the stoppered flask is placed into a basin of boiling water.
	[1]

[3]

(b) N₂O₄ is also used in the large scale manufacture of nitric acid. It reacts with water to give both nitrous acid and nitric acid.

$$N_2O_4 + H_2O \rightarrow HNO_2 + HNO_3$$

The two acids have different acid strengths. To determine the acid strength of the two acids, two separate solutions containing $0.10 \text{ mol } \text{dm}^{-3}$ of each acid were prepared. The pH was found to be 2.17 and 1.00 for the solutions containing nitrous acid and nitric acid respectively.

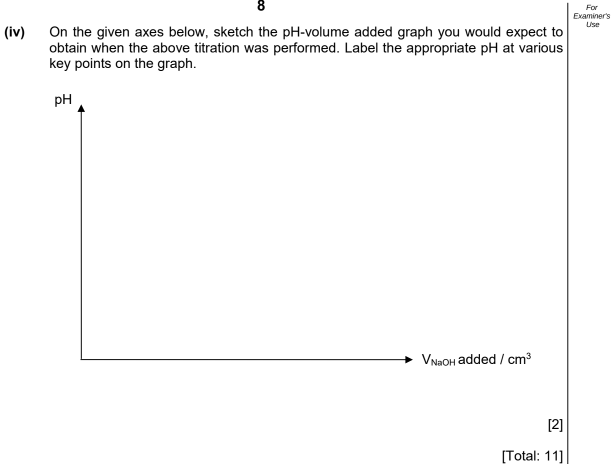
(i) Use the data provided to prove nitrous acid is a weak acid and hence, determine its K_a value.

[2]

(ii) Suggest a reason why nitrous acid is a weaker acid than nitric acid.

25.0 $\rm cm^3$ of the prepared 0.10 mol $\rm dm^{-3}$ nitrous acid was titrated with 0.10 mol $\rm dm^{-3}$ aqueous sodium hydroxide.

(iii) Using your value of K_a calculated in part (i), calculate the pH when 25.00 cm³ of aqueous sodium hydoxide has been added.



8

3 The equation for the reaction between bromine and methanoic acid is as follows:

 $Br_2(aq) + HCOOH(aq) \rightarrow 2HBr(aq) + CO_2(g)$

It is hypothesised that the reaction is elementary. To prove this hypothesis, volumes of the two reactants were varied and the rate of the reaction is measured in terms of the rate at which the bromine concentration changes. When the total volume is kept constant, the following relationship is true.

rate of reaction $\alpha \frac{\text{volume of bromine used}}{\text{time for color of bromine to disappear}}$

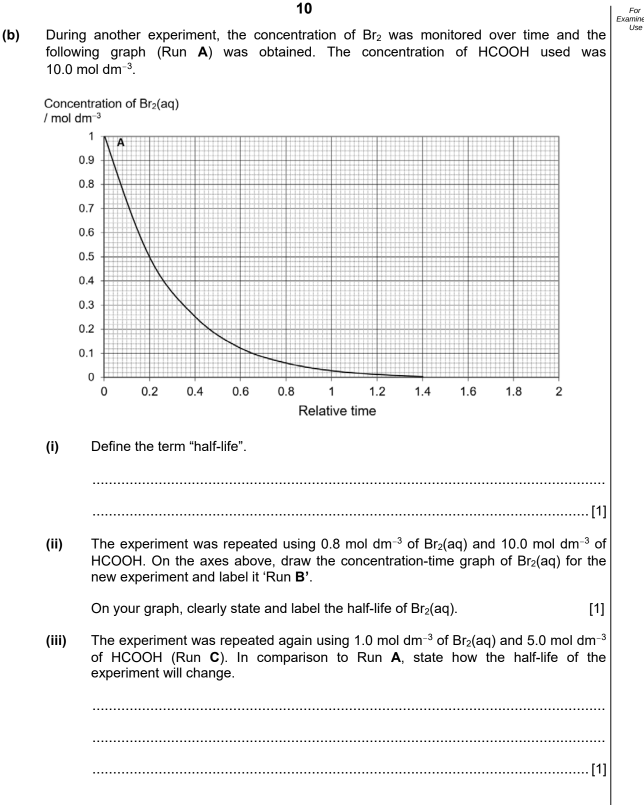
The temperature of the reaction mixture was maintained at 25 °C.

The following results were obtained in three repeated experiments:

Expt	Volume of 1.0 mol dm ⁻³ Br ₂ / cm ³	Volume of 10.0 mol dm ⁻³ HCOOH / cm ³	Volume of water added / cm ³	Relative time for colour of bromine to disappear
1	10	10	0	1.4
2	40	20	20	2.8
3	5	10	5	1.4

(a) By comparing the rates of reactions, explain how the results of the three experiments support the hypothesis that the reaction is elementary.

••••••	 	
		[3]
	 •••••••••••••••••••••••••••••••••••••••	[0]

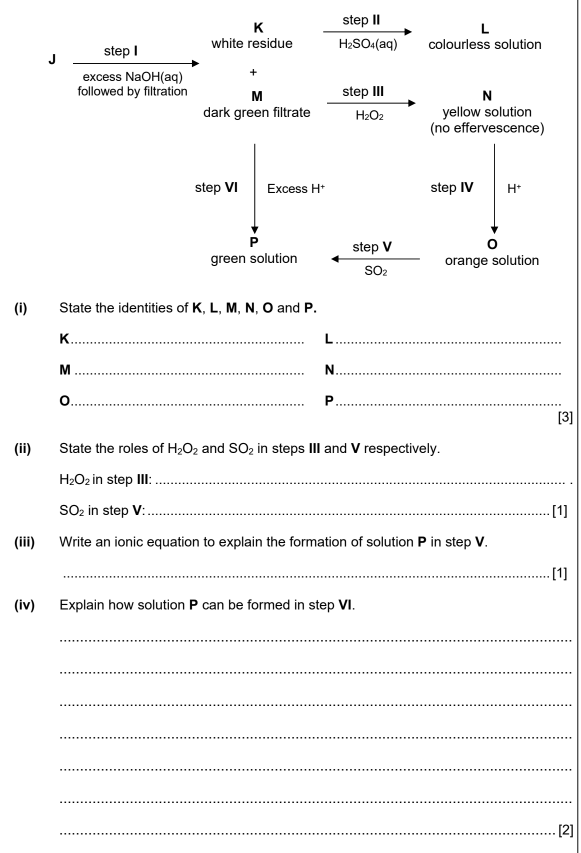


(c) By drawing a suitable illustration, estimate and explain the change in rate of reaction when temperature is increased to 35 °C.

 	 [3]
	[Total: 9]

For Examiner's

4(a) Solution **J** contains a Group 2 ion and a transition metal ion. The following reaction scheme shows how **J** reacts with some common reagents in the laboratory.



[Turn Over

(b) Among the elements of Group 14, those towards the top, carbon to germanium, have very different properties from those at the bottom, tin and lead. For example, the melting points show a marked change after germanium.

13

element	С	Si	Ge	Sn	Pb
mp / °C	>3550	1410	937	232	327

Carbon, silicon and germanium each form a solid with the same type of structure.

(i) Explain why the melting points of these elements decrease from carbon to germanium.[2] (ii) Explain how first ionisation energy changes from carbon to germanium.[2] Carbon and silicon each form a tetrachloride. CCl₄ has no reaction with water; SiCl₄ reacts violently with water. (iii) Write a balanced equation for the reaction of SiCl₄ with water.[1] (iv) Suggest an explanation for the inertness of CCl₄ to water.[1] [Total: 13]

For Examiner's

[Turn Over

Azo dyes are made in large quantites from benzene, C_6H_6 , via nitrobenzene, $C_6H_5NO_2$ (density = 1.20 g cm⁻³), and phenylamine, $C_6H_5NH_2$.

5

(a) The preparation of nitrobenzene requires benzene to be warmed under reflux at about 55 °C with a mixture of concentrated nitric and sulfuric acids. Some information about these substances is given below:

Benzene: immiscible with water; highly flammable; extremely toxic by ingestion or inhalation; known carcinogen.

Concentrated nitric acid: miscible with water; causes severe burns to eyes and skin; strong oxidising agent. The acid contains about 30 % water by volume.

Concentrated sulfuric acid: miscible with water; causes severe burns to eyes and skin; strong oxidising agent; dilution with water is very exothermic and can be dangerous.

(i) Nitric acid is placed in a suitable flask and sulfuric acid is added slowly with cooling of the flask. Explain why cooling is necessary.

.....[1]

(ii) Benzene is added slowly to the acid mixture, which is then warmed at 55 °C for 45 minutes under reflux with vigorous stirring of the reaction mixture.

Explain why the reflux condenser is necessary and also why the mixture is vigorously stirred.

.....[2]

(iii) State, with a reason, **one** other precaution (other than wearing protective wear) that would be necessary when carrying out the experiment.

.....[1]

(iv) The reaction mixture is then poured into a large excess of cold water, the liquid nitrobenzene layer is separated and *washed* with sodium carbonate solution. Explain why this washing is necessary.

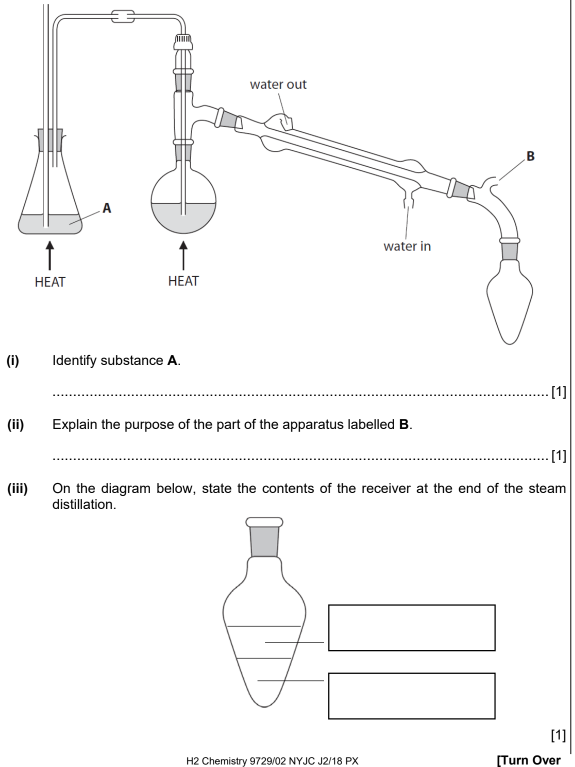
.....[1]

.....[1]

(v) The nitrobenzene layer is dried before being distilled to purify it. Identify a suitable drying agent.

(b) Steam distillation is a purification process to separate nitrobenzene from the reaction mixture. During the process of steam distillation, a current of steam is blown through a mixture containing the desired organic substance to be distilled. This caused the desired organic substance to vaporise. The vapour containing the desired organic substance can then be condensed and collected. This method is used predominantly to purify liquids that are not very volatile and are immiscible with water.

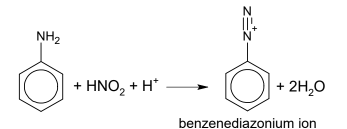
The diagram below shows a steam distillation apparatus used to extract nitrobenzene from the reaction mixture.



(c) The purified nitrobenzene is then reduced to phenylamine, $C_6H_5NH_2$.

The phenylamine is diazotised by reaction with nitrous acid at a temperature between 0 °C and 10 °C. Nitrous acid is generated in the reaction mixture from sodium nitrite and hydrochloric acid.

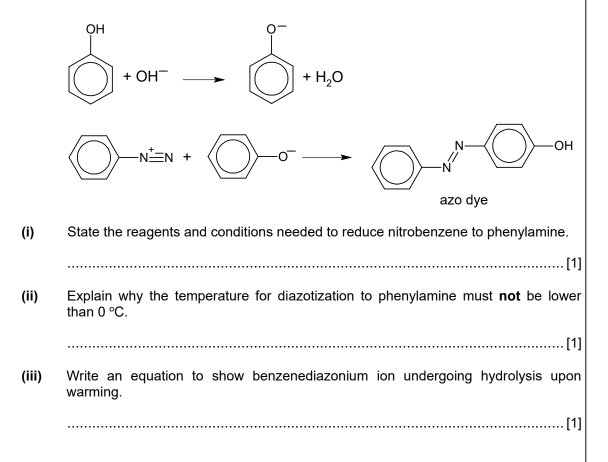
The ionic equation for the diazotisation of phenylamine to produce benzenediazonium ion is as shown below.



If the above reaction is warmed, benzenediazonium ion will undergo hydrolysis to give phenol. A gas will also be produced and the resulting mixture is acidic.

Reaction of the benzenediazonium compound with an alkaline solution of a phenol, C_6H_5OH , will produce a solid azo dye, which is purified by recrystallisation.

The equations for the reaction between benzenediazonium ion and phenol to produce the solid azo dye are shown below.



	17	Exa
Purifi 1. 2. 3. 4. 5.	cation by recrystallisation requires the following steps: The azo dye is dissolved in a minimum volume of hot solvent. The solution is filtered through a pre-heated funnel. The solution is cooled and filtered using a Buchner funnel. The solid is washed with a small amount of cold solvent. The solid is dried in a desiccator.	
(i)	Explain why a minimum volume of hot solvent is used in step 1.	
	[1]
(ii)	Explain why the funnel must be pre-heated.	
	[1]
(iii)	Suggest a reason why it is preferable to dry the solid in a desiccator rather than ir an oven.	ו
	[1]
This	question compares the acidity and basicity of some organic compounds.	
(i)	Explain why an aqueous solution of of N,N-dimethylphenylamine is more basic than an aqueous solution of phenylamine.	ו
	H ₃ C CH ₃	
	N,N-dimethylphenylamine	
	[1]
(ii)	Explain why an aqueous solution of azo dye is more acidic than an aqueous solution of phenol.	5
		•
	[2]
	[Total: 18]
	H2 Chemistry 9729/02 NYJC J2/18 PX [End of Paper	

[End of Paper]



NANYANG JUNIOR COLLEGE JC 2 PRELIMINARY EXAMINATION Higher 2

CANDIDATE NAME

Teachers' Mark Scheme

CLASS

TUTOR'S NAME

CHEMISTRY

Paper 2 Structured

Candidates answer on the Question Paper

Additional Materials: Data Booklet

READ THESE INSTRUCTIONS FIRST

Write your name and class on all the work you hand in. 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. A Data Booklet is provided.

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	/24	
2	/11	
3	/9	
4	/13	
5	/18	
Total	/75	

9729/02

2 hours

11 September 2018

2

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

1(a) An unknown sample was found to contain the anions, Cl⁻, ClO₃⁻ and NO₃⁻. A student weighed a sample into a beaker and recorded the following data.

mass of beaker and sample / g	68.962
mass of empty beaker / g	67.620

The sample was dissolved and diluted in a 250 cm³ volumetric flask to obtain solution L.

In **experiment 1**, a 50 cm³ portion of solution **L** was reacted with excess silver nitrate solution. The AgCl precipitated was transferred onto a dry filter paper and was placed under an infra-red lamp. The dry AgCl precipitate was weighed and the following data was obtained.

Experiment 1

mass of dry filter paper and AgCl / g	0.737
mass of dry filter paper / g	0.620

In **experiment 2**, a gas was bubbled into another 50 cm³ portion of solution **L** to convert CIO_3^- to CI^- before the addition of excess silver nitrate solution. The AgCl precipitated was also dried and weighed. The following data was obtained.

Experiment 2

mass of dry filter paper and AgCl / g	0.799
mass of dry filter paper / g	0.651

(i) Write the half equation for the reduction of CIO_3^- to CI^- .

 $CIO_3^- + 6H^+ + 6e^- \rightarrow CI^- + 3H_2O$ [1].....[1]

(ii) Determine the mass of CI^- in 50 cm³ of solution L.

mass of AgCl precipitate in expt 1 = 0.737 - 0.620 = 0.117 g mass of Cl⁻ present in 50 cm³ portion = $\frac{36.6}{36.5 + 107.9} \times 0.117 = 0.02896$ g ≈ 0.0290 g [1]

[1]

(iii) Determine the mass of Cl^- converted from ClO_3^- in experiment 2.

mass of AgCl precipitate in expt 2 = 0.799 - 0.651 = 0.148 g mass of Cl⁻ present in 50 cm³ portion = $\frac{36.6}{36.6 + 107.9} \times 0.148 = 0.03663$ g [1] mass of Cl⁻ from ClO₃⁻ in 50 cm³ = 0.03663 - 0.02896= 0.007674 g ≈ 0.00767 g [1]

H2 Chemistry 9729/02 NYJC J2/18 PX

(iv) Hence, determine the percentage mass of CIO_3^- in the unknown sample.

Since $n(CIO_3^-)$: $n(CI^-)$ is 1 : 1, mass of CIO_3^- present in 50 cm³ = 0.007674 x $\frac{\$5.5+\$(16)}{\$5.5}$ = 0.01805 g [1] mass of CIO_3^- present in 250 cm³ = 0.01805 x $\frac{260}{50}$ = 0.1805 g [1] % CIO_3^- present in unknown compound = $\frac{0.1805}{68.962-67.620}$ x 100% = 13.45% \approx 13.5% [1]

- [3]
- (v) The E_{\uparrow} (ClO₃⁻/Cl⁻) has a value of +1.47 V. From the list of standard electrode potentials in the *Data Booklet*, identify a gas that would reduce ClO₃⁻ to Cl⁻. Explain your answer.

 E_{\uparrow} (H⁺/H₂) = 0.00 V H₂ gas is an appropriate reducing agent. [1] (also accept E_{\uparrow} (NO₃⁻/NO₂) or E_{\uparrow} (SO₄²⁻/SO₂) as their E_{\uparrow} < +1.47V)

 $E_{\text{cell}} = (+1.47) - (0.00) = +1.47 \text{ V} > 0$

Since, $E_{cell} > 0$, the reaction is feasible. [1]

-[2]
- (b) (i) An aqueous solution of HCl has a density of 1.15 g cm⁻³ and is 30% by mass of HCl.

Calculate the concentration in mol dm⁻³ of this solution of HCl.

mass of HCl in 1 cm³ = $\frac{30}{100}$ x 1.15 = 0.3450 g [1] [HCl] = $\frac{0.8450}{1.0+36.6}$ x 1000 = 9.45 mol dm⁻³ [1]

[2]

(ii) Calculate the volume of this solution required to prepare 5 dm³ of 0.20 mol dm⁻³ HCl by dilution with water.

volume of HCl required = $\frac{6 \times 0.20}{9.462}$ = 0.106 dm⁻³ = 106 cm³ [1]

(c) The gelatin silver process is the photographic process used with black-and-white films. The following information pertains to the process of taking photographs and developing films.

Taking photographs

- A 35 mm cartridge of black-and-white print film contains a long strip of plastic that has layered coatings on each side.
- On the front side of the film, the layers are made of gelatin which contain grains of silver chloride crystals.
- When the shutter of the camera is opened for a fraction of a second to allow the film to be exposed to light, these crystals undergo decomposition thereby producing an image on the film.

Developing films

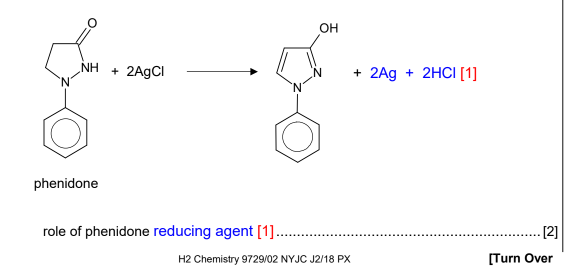
- After the photographs have been taken, the film is developed in a dark room under a light source that emits low energy light.
- Firstly, the film is soaked in water before adding phenidone. Phenidone makes the image more visible by reacting with the exposed silver chloride crystals to produce silver atoms and two other by-products.
- This reaction can only proceed at high pH.
- After some time, the reaction will then be quenched.
- Finally, the film will be soaked in ammonium thiosulfate, (NH₄)₂S₂O₃, which is used as a fixer to make the image permanent and light resistant. This is done through the reaction between the unexposed silver chloride crystals and the fixer.
- (i) Write the balanced equation for the decomposition of silver chloride crystals when it is exposed to light.



-[1]
- (ii) Suggest a suitable colour of the light source that is used in a dark room.

Red. (Also accept orange or yellow) [1][1]

(iii) Complete the equation for the reaction between the developing agent, phenidone, and the exposed silver chloride crystals. Hence, state the role of phenidone in this reaction.



(iv)	Suggest a suitable reagent, other than excess cold water, that can be used to quench the development of the film. Explain.
	Acetic acid / Citric acid / Any plausible acids. e.g. HCI, HNO ₃ , H ₂ SO ₄ [1]
	When acid is added, the pH will be lowered. Hence, the reaction will not proceed a lower pH. [1]
	[2
(v)	When the non-exposed silver chloride crystals react with the fixer, $(NH_4)_2S_2O_3$, a silver complex compound M is formed together with a chloride salt, N . Both M and N have the same cation. The silver-containing complex ion has a coordination number of 2 and is chlorine-free.
	Suggest the formulae of compounds M and N .
	compound M : (NH ₄) ₃ [Ag(S ₂ O ₃) ₂] [1] compound N : NH ₄ Cl [1][2]
(vi)	State the shape of the silver-containing ion in compound ${f M}$.
	Linear [1][1]
(vii)	Complete the electronic configuration of silver in compound ${f M}$. Hence, deduce the colour of compound ${f M}$.
	compound M [Ar]3d ¹⁰ 4s ² 4p ⁶ 4d ¹⁰ [1]
	Since the 4d orbitals are fully occupied, d–d transition cannot occur. [1]
	Hence, compound M is colourless. [1]
	[2]
(viii)	Explain why the fixer, $(NH_4)_2S_2O_3$, is able to make the image permanent and light resistant on the film.
	The fixer is able to remove any remaining unexposed AgCI. Hence, there is <u>no AgC</u> remaining on the film to undergo decomposition through the exposure of light to change the image on the film. [1]
	[1]

[Total: 24]

- 2 Dinitrogen tetroxide, commonly referred to as nitrogen tetroxide, is the chemical compound N₂O₄. It is a useful reagent in chemical synthesis.
- (a) Colourless N_2O_4 readily dissociates to form brown NO_2 and the following equilibrium is reached fairly quickly in the gaseous phase.

$$N_2O_4(g)$$
 Ý $2NO_2(g)$ $\Delta H > 0$

(i) When 4.60 g of N_2O_4 is placed in an evacuated 1.48 dm³ flask at 27 °C, the equilibrium pressure is 1 atm.

Calculate the value of K_p at 27 °C.

 Initial no of moles of N₂O₄ = 4.60 / (14.0 × 2 + 16.0 × 4) = 0.0500 mol ICE table/eqm amounts: N₂O₄ (g) ⇐ 2NO₂ (g)

Initial / mol 0.0500 0 Eqm / mol 0.0500 - x +2x

- PV = nRT (101325)(1.48 × 10⁻³) = n (8.31)(300) n(gases at eqm) = 0.06015 = 0.0602 mol
- Solve for x and find eqm amts
 0.0500 x + 2x = 0.06015
 x = 0.01015

$$N_2O_4(g) \rightleftharpoons 2NO_2(g)$$

• Eqm / mol 0.03985 0.0203
•
$$K_p = \frac{(P_{NO_2})^2}{P_{N_2O_4}} = \frac{(\frac{0.02030}{0.06015} \times 1)^2}{\frac{0.03985}{0.06015} \times 1} = 0.172 \text{ atm}$$

4 points 3 marks; 3 points: 2 marks; 2 points: 1 mark

Alternative solution in terms of partial pressure ICE table.

 $P(N_2O_4 \text{ initial}) = \frac{0.05(8.314)(300)}{1.48 \times 10^{-3}} = 84220 Pa = 0.08312 atm$

Initial / atm 0.08312 0 Eqm / atm 0.08312 - x +2x

0.08312 - x + 2x = 1x = 0.1687 atm

$$K_{p} = \frac{(P_{NO_{2}})^{2}}{P_{N_{2}O_{4}}} = \frac{(0.1687 \times 2)^{2}}{0.8312 - 0.1687} = 0.172 \text{ atm}$$

(ii) Describe and explain what you would observe when the stoppered flask is placed into a basin of boiling water.

<u>Temperature of system increases</u> and by Le Chatelier's Principle, the <u>forward</u> endothermic reaction is favoured. Position of equilibrium shifts to the right to absorb added heat and more NO_2 is formed. Hence the reaction mixture becomes <u>more</u> brown. [1]

.....[1]

[3]

(b) N₂O₄ is also used in the large scale manufacture of nitric acid. It reacts with water to give both nitrous acid and nitric acid.

$$N_2O_4 + H_2O \rightarrow HNO_2 + HNO_3$$

The two acids have different acid strengths. To determine the acid strength of the two acids, two separate solutions containing $0.10 \text{ mol } \text{dm}^{-3}$ of each acid were prepared. The pH was found to be 2.17 and 1.00 for the solutions containing nitrous acid and nitric acid respectively.

(i) Use the data provided to prove nitrous acid is a weak acid and hence, determine its K_a value.

[H⁺] in solution of HNO₂ = $10^{-2.17} = 0.00676 \text{ moldm}^{-3}$ which is less than [HNO₂] = 0.10 moldm⁻³. Hence HNO₂ only dissociates partially and it is a weak acid. [1]

OR for the same concentration of acid, HNO_2 dissociates to produce a lower [H⁺] (as shown by the higher pH of the solution). Hence, it dissociates to a smaller extent and is a weak acid. \boxtimes higher pH so weaker acid.

(FYI. Not required in answer: $[H^+]$ in solution of $HNO_3 = 10^{-1.00} = 0.100$ mol dm⁻³ = $[HNO_3]$ hence Not required it is completely dissociated and HNO_3 is a strong acid.)

$$K_{a} = \frac{[NO_{2}^{-}][H^{+}]}{[HNO_{2}]} = \frac{(0.00676)^{2}}{0.10 - 0.00676} = 4.90 \times 10^{-4} \text{ mol dm}^{-3} \text{ [1]}$$

[Note if students use [H⁺] = $\sqrt{K_{a} \cdot c}$ or K_a = $\frac{(0.00676)^2}{0.10} = 4.57 \times 10^{-4}$]

[2]

(ii) Suggest a reason why nitrous acid is a weaker acid than nitric acid.

Nitric acid is a stronger acid as NO_3^- is a more stable conjugate base than NO_2^- as the negative charge is more effectively dispersed over a greater number of electronegative oxygen atoms (or vice versa). [1]

......[1]

For Examiner's Use

25.0 cm³ of the prepared 0.10 mol dm⁻³ nitrous acid was titrated with 0.10 mol dm⁻ aqueous sodium hydroxide.

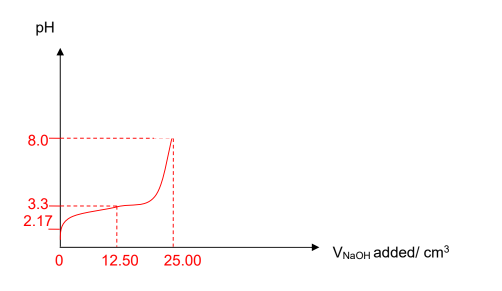
(iii) Using your value of K_a calculated in part (i), calculate the pH when 25.00 cm³ of aqueous sodium hydoxide has been added.

Equivalence volume = 25.00 cm³ At equivalence point, solution contains basic salt only.

$$\begin{split} \mathsf{K}_{\mathrm{b}} &= 10^{-14} / 4.90 \times 10^{-4} = 2.040 \times 10^{-11} \\ \text{[salt]} &= 25.0 \times 0.1 / 50.0 = 0.05 \text{ moldm}^{-3} \\ \text{[OH}^{-}] &= \sqrt{K_b.c} = \sqrt{2.040 \times 10^{-11} (0.05)} = 1.010 \times 10^{-6} \\ \mathsf{pH} &= 14 - \mathsf{pOH} = 14 - (-\mathsf{lg}(1.010 \times 10^{-6})) = 8.0 \\ \textbf{4 points [2], 2 points [1] ecf K_a value} \end{split}$$

[2]

(iv) On the given axes below, sketch the pH-volume added graph you would expect to obtain when the above titration was performed. Label the appropriate pH at various key points on the graph.



Label following values

- Initial pH (given in question)
- Maximum buffering capacity occurs at 12.50 cm³: $pH = pK_a$ (-lg(4.90x10⁻⁴) = 3.3)
- Equivalence point (calculated in iii)
- Correct shape (relatively flat at buffer region)

*don't penalize if students sketch graph beyond 25.00 cm³

4 points [2], 2-3 points [1]

[2]

3 The equation for the reaction between bromine and methanoic acid is as follows:

 $Br_2(aq) + HCOOH(aq) \rightarrow 2HBr(aq) + CO_2(q)$

It is hypothesised that the reaction is elementary. To prove this hypothesis, volumes of the two reactants were varied and the rate of the reaction is measured in terms of the rate at which the bromine concentration changes. When the total volume is kept constant, the following relationship is true.

rate of reaction $\alpha \frac{\text{volume of bromine used}}{\text{time for color of bromine to disappear}}$

The temperature of the reaction mixture was maintained at 25 °C.

The following results were obtained in three repeated experiments:

Expt	Volume of 1.0 mol dm ⁻³ Br ₂ / cm ³	Volume of 10.0 mol dm ⁻³ HCOOH / cm ³	Volume of water added / cm ³	Relative time for colour of bromine to disappear
1	10	10	0	1.4
2	40	20	20	2.8
3	5	10	5	1.4

(a) By comparing the rates of reactions, explain how the results of the three experiments support the hypothesis that the reaction is elementary.

Expt	Vol. of 1.0	Vol. of 10.0 mol	Vol. of	Relative time	Rate
	mol dm ⁻³ Br ₂	dm [–] 3 HCOOH /	water	for colour of	
	/ cm ³	cm ³	added /	bromine to	
			cm ³	disappear	
1	10	10	0	1.4	10/1.4=7.14
2	40	20	20	2.8	
2a	40/4=10	20/4=5	20/4=5	2.8	10/2.8=3.57
3	5	10	5	1.4	5/1.4=3.57

Using expt 1 and 3,

When vol of Br_2 halved i.e. 10/5, the rate halved i.e. 7.14/3.57. Hence rate is directly proportional to Br_2 , 1st order wrt Br_2 .

Using expt 1 and 2a,

When vol of HCOOH halved i.e. 10/5, the rate halved i.e. 7.14/3.57. Hence rate is directly proportional to HCOOH, 1st order wrt HCOOH.

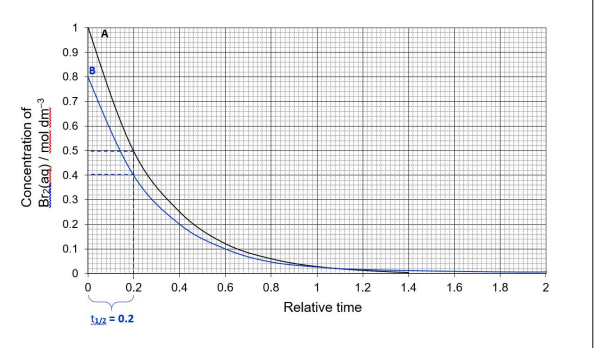
Since orders of reaction correspond to stoichiometric ratio of the overall equation, the reaction is elementary.

1 mark for finding order wrt Br₂

1 mark for finding order wrt HCOOH and deducing the hypothesis correctly

.....[3]

For Examiner's Use (b) During another experiment, the concentration of Br₂ was monitored over time and the following graph (Run A) was obtained. The concentration of HCOOH used was 10.0 mol dm⁻³.



Define the term "half-life". (i)

Time taken for concentration of reactant to reach half its original concentration.

.....[1]

The experiment was repeated using 0.8 mol dm⁻³ of Br₂(aq) and 10.0 mol dm⁻³ of (ii) HCOOH. On the axes above, draw the concentration-time graph of Br₂(aq) for the new experiment and label it 'Run B'.

On your graph, clearly state and label the half-life of Br₂(aq). [1]

Curve must show at least 2 constant half-lives. Only one half-life needs to be clearly labelled.

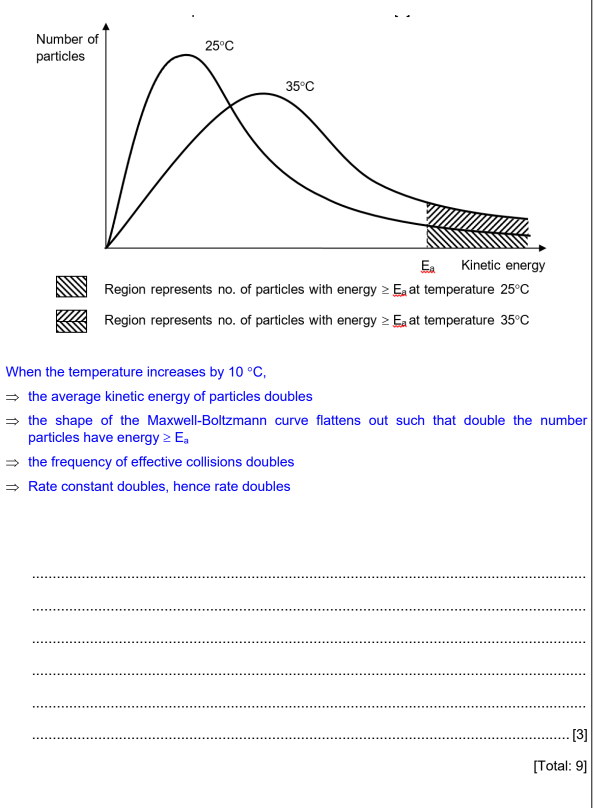
The experiment was repeated again using 1.0 mol dm⁻³ of Br₂(ag) and 5.0 mol dm⁻³ (iii) of HCOOH (Run C). In comparison to Run A, state how the half-life of the experiment will change.

The reaction is a pseudo-first order reaction in which $\mathbf{t}_{1/2} = \frac{112}{\text{k[HCOOH]}}$

Hence if [HCOOH] halves, $t_{1/2}$ will double from 0.2 to 0.4.

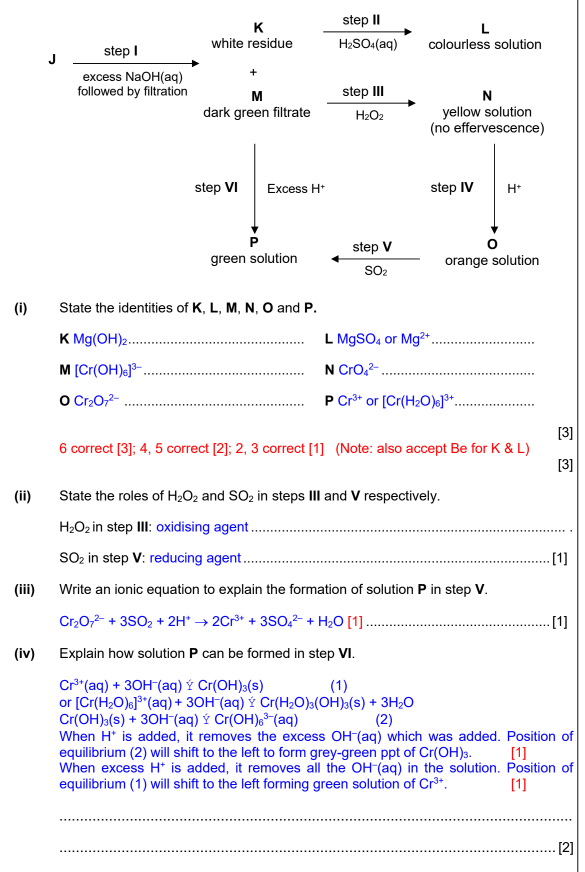
......[1]

(c) By drawing a suitable illustration, estimate and explain the change in rate of reaction when temperature is increased to 35 °C.



For Examiner's

4(a) Solution **J** contains a Group 2 ion and a transition metal ion. The following reaction scheme shows how **J** reacts with some common reagents in the laboratory.



H2 Chemistry 9729/02 NYJC J2/18 PX

[Turn Over

(b) Among the elements of Group 14, those towards the top, carbon to germanium, have very different properties from those at the bottom, tin and lead. For example, the melting points show a marked change after germanium.

element	С	Si	Ge	Sn	Pb
mp / °C	>3550	1410	937	232	327

Carbon, silicon and germanium each form a solid with the same type of structure.

(i) Explain why the melting points of these elements decrease from carbon to germanium.

They have giant covalent structures with strong covalent bonds between atoms in a 3-dimensional network. [1] From carbon to germanium, the <u>atomic radius increase</u> and <u>the bond length increase</u> (C–C bond length < Si–Si bond length < Ge–Ge bond length), hence the <u>covalent bond strength decrease</u> from carbon to silicon to germanium. [1] Since the melting of these elements require breaking the covalent bonds between the respective atoms, the melting point decreases since less energy is required to break the weaker covalent bonds.

.....[2]

(ii) Explain how first ionisation energy changes from carbon to germanium.

Down the group from carbon to germanium,

- the <u>number of protons increases</u>, <u>nuclear charge increases</u>.
- As the <u>number of electron shells increases</u>, <u>shielding effect increases</u> <u>significantly</u>.
- The <u>outermost electron is further away from the nucleus</u>, hence attraction between the nucleus and outermost electron <u>decreases</u>

• The <u>first ionisation energy decreases</u> down a group.

4 points – [2]; 2,3 points – 1

.....[2]

Carbon and silicon each form a tetrachloride. CCI_4 has no reaction with water; $SiCI_4$ reacts violently with water.

(iii) Write a balanced equation for the reaction of SiCl₄ with water.

 $SiCl_4 + H_2O \rightarrow SiO_2 + 4HCI [1] [1]$

(iv) Suggest an explanation for the inertness of CCl₄ to water.

Water molecules could not form co-ordinate/dative bonds with the central carbon atom of CCl_4 because carbon is in period 2 and does not have <u>energetically accessible low lying orbitals</u> to accommodate lone pair of electrons from O atom in H₂O. [1]

.....

.....[1]

[Total: 13]

[Turn Over

14

- 5 Azo dyes are made in large quantites from benzene, C_6H_6 , via nitrobenzene, $C_6H_5NO_2$ (density = 1.20 g cm⁻³), and phenylamine, $C_6H_5NH_2$.
- (a) The preparation of nitrobenzene requires benzene to be warmed under reflux at about 55 °C with a mixture of concentrated nitric and sulfuric acids. Some information about these substances is given below:

Benzene: immiscible with water; highly flammable; extremely toxic by ingestion or inhalation; known carcinogen.

Concentrated nitric acid: miscible with water; causes severe burns to eyes and skin; strong oxidising agent. The acid contains about 30 % water by volume.

Concentrated sulfuric acid: miscible with water; causes severe burns to eyes and skin; strong oxidising agent; dilution with water is very exothermic and can be dangerous.

(i) Nitric acid is placed in a suitable flask and sulfuric acid is added slowly with cooling of the flask. Explain why cooling is necessary.

To avoid the temperature rising too much OR the reaction between sulfuric acid and water is exothermic as the sulfuric acid is diluted by the water in nitric acid

.....

.....[1]

(ii) Benzene is added slowly to the acid mixture, which is then warmed at 55 °C for 45 minutes under reflux with vigorous stirring of the reaction mixture.

Explain why the reflux condenser is necessary and also why the mixture is vigorously stirred.

Prevents escape of benzene / volatile liquids [1] reactants are immisicible/do not mix/form separate layers so they need to be stirred to make <u>reaction rate acceptable or increase frequency of effective collisions or</u> <u>increase the surface area of contact between the two immiscible layers or to enable</u> <u>the reactant molecules to collide with the correct orientation [1]</u>

(iii) State, with a reason, **one** other precaution (other than wearing protective wear) that would be necessary when carrying out the experiment.

EITHER benzene is toxic so use fume cupboard OR benzene/nitrobenzene is flammable so use heating mantle/water bath [1] (iv) The reaction mixture is then poured into a large excess of cold water, the liquid nitrobenzene layer is separated and *washed* with sodium carbonate solution. Explain why this washing is necessary.

sodium carbonate removes/neutralises (residual) acid [1]

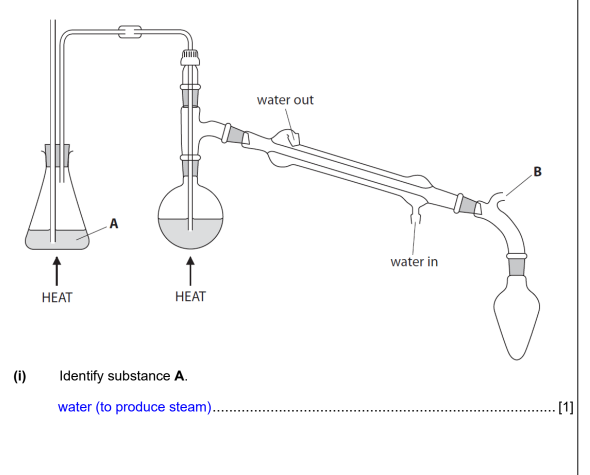
.....[1]

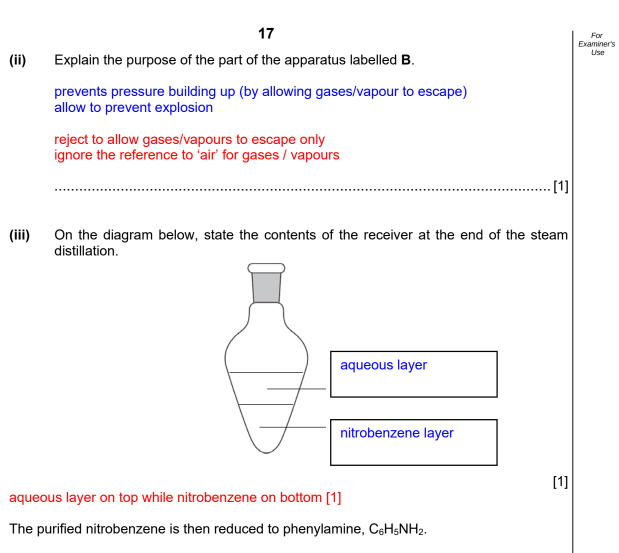
(v) The nitrobenzene layer is dried before being distilled to purify it. Identify a suitable drying agent.

(anhydrous) sodium sulfate / magnesium sulfate OR (anhydrous) calcium chloride OR silica gel

-[1]
- (b) Steam distillation is a purification process to separate nitrobenzene from the reaction mixture. During the process of steam distillation, a current of steam is blown through a mixture containing the desired organic substance to be distilled. This caused the desired organic substance to vaporise. The vapour containing the desired organic substance can then be condensed and collected. This method is used predominantly to purify liquids that are not very volatile and are immiscible with water.

The diagram below shows a steam distillation apparatus used to extract nitrobenzene from the reaction mixture.

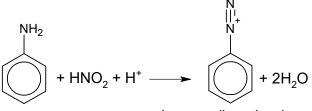




The phenylamine is diazotised by reaction with nitrous acid at a temperature between 0 $^{\circ}$ C and 10 $^{\circ}$ C. Nitrous acid is generated in the reaction mixture from sodium nitrite and hydrochloric acid.

(C)

The ionic equation for the diazotisation of phenylamine to produce benzenediazonium ion is as shown below.

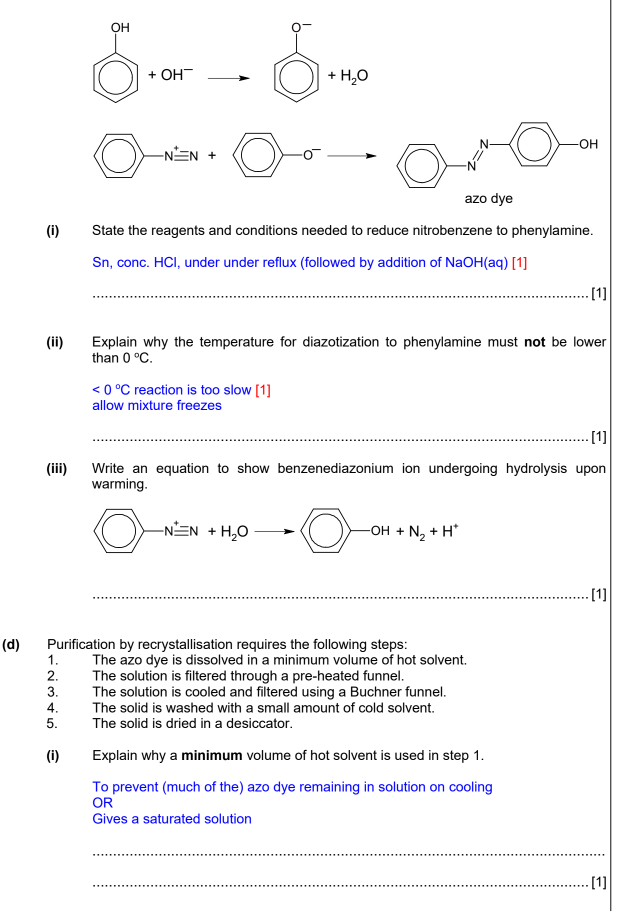


benzenediazonium ion

If the above reaction is warmed, benzenediazonium ion will undergo hydrolysis to give phenol. A gas will also be produced and the resulting mixture is acidic.

Reaction of the benzenediazonium compound with an alkaline solution of a phenol, C_6H_5OH , will produce a solid azo dye, which is purified by recrystallisation.

The equations for the reaction between benzenediazonium ion and phenol to produce the solid azo dye are shown below.



[Turn Over

For Examiner's Use

18

	19	Exa
(iii)	Explain why the funnel must be pre-heated.	
	To prevent crystallization (of the azo dye) [1]	
	[1]	
(iv)	Suggest a reason why it is preferable to dry the solid in a desiccator rather than in an oven.	
	Decomposition could occur if the compound were to be heated OR	
	Compound might melt	
	[1]	
This c	question compares the acidity and basicity of some organic compounds.	
(i)	Explain why an aqueous solution of N,N-dimethylphenylamine is more basic than an aqueous solution of phenylamine.	
	H ₃ C CH ₃	
	\downarrow	
	N,N-dimethylphenylamine	
	There are <u>2 electron-donating $-CH_3$ groups</u> [pt 1] attached to the N atom in N,N-dimethylphenylamine. Hence the <u>electron density on the N atom in N,N-dimethylphenylamine is higher</u> [pt 2] than that in phenylamine. The <u>lone pair of electrons on the N atom in N,N-dimethylphenylamine is more available</u> [pt 3] for dative bonding with a H ⁺ .	
	[1]	
(ii)	Explain why an aqueous solution of azo dye is more acidic than an aqueous solution of phenol.	
	This group, $\underline{C}_{6}\underline{H}_{5}\underline{N}=\underline{N}$ is an <u>electron-withdrawing</u> [pt 4] group which <u>further</u> <u>disperses the negative charge on the O atom</u> [pt 5] in the conjugate base of azo dye. The <u>conjugate base formed by the azo dye is further stabilized</u> [pt 6]. <u>The azo dye</u> <u>donates proton more readily</u> [pt 7].	
	[2]	
	7 pts – 3 marks 5 – 6 pts – 2 marks 3 – 4 pts – 1 mark	

H2 Chemistry 9729/02 NYJC J2/18 PX

[End of Paper]



NANYANG JUNIOR COLLEGE JC 2 PRELIMINARY EXAMINATION Higher 2

CHEMISTRY

Paper 3 Free Response

9<mark>647<u>29</u>/03</mark>

21_18_September 20186 2 hours

Formatted: Not Highlight

Additional Materials:

Writing Paper Data Booklet

READ THESE INSTRUCTIONS FIRST

Write your name and class on all the work you hand in. Write in dark blue or black pen on both sides of the paper. You may use a soft pencil for any diagrams, graphs or rough working. Do not use staples, paper clips, highlighters, glue or correction fluid.

Section A Answer all questions.

Section B Answer one question.

<u>A Data Booklet is provided.</u> <u>The use of an approved scientific calculator is expected, where appropriate.</u> Answer any **four** questions. A Data Booklet is provided. You are reminded of the need for good English and clear presentation in your answers.

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.

[Turn over

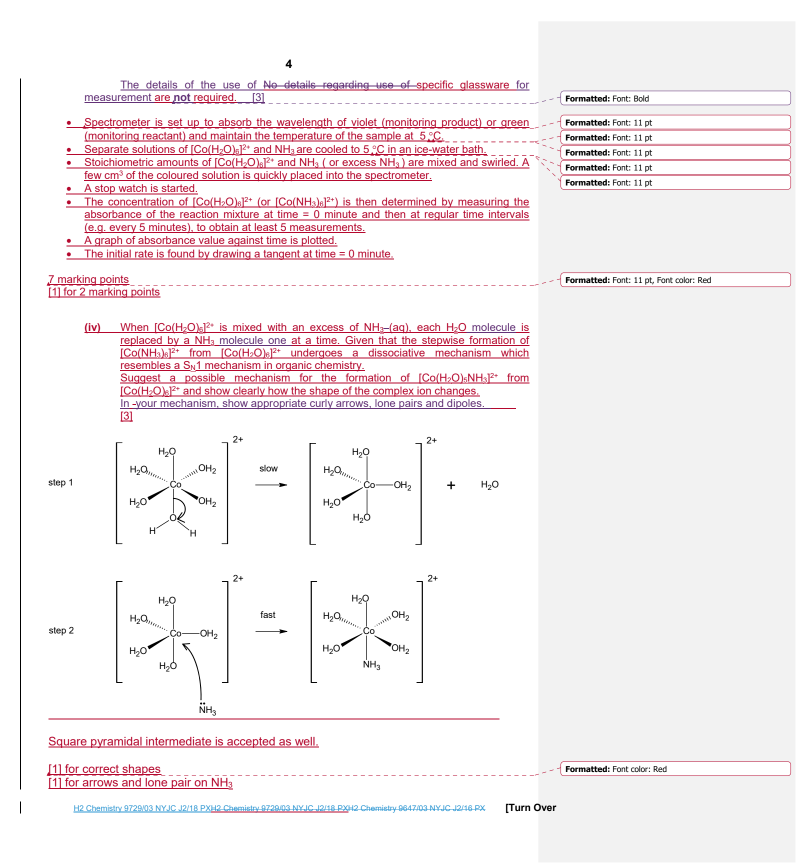
This document consists of **<u>13-14</u>** printed pages and **<u>10</u>** blank page.

2

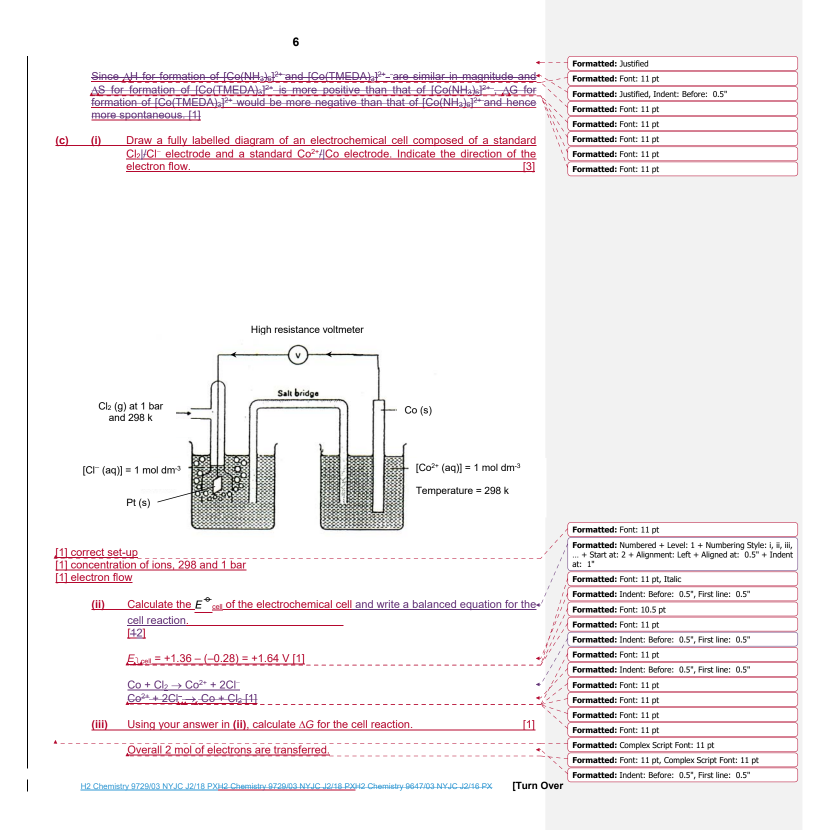
[Turn over

Formatted: Left

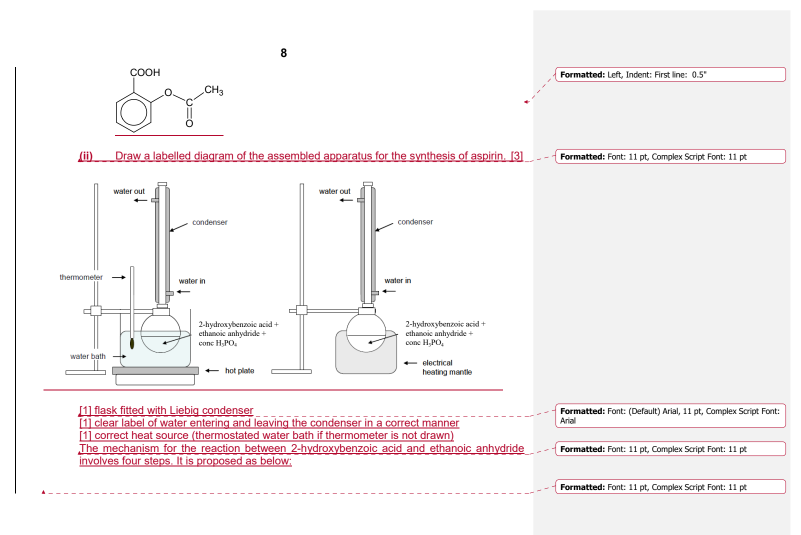
3	
Section A	Formatted: Font: 11 pt, Complex Script Font: 11 pt
Answer all questions in this section.	
$\frac{1}{1} \qquad \begin{array}{c} \mbox{Cobalt is a typical transition element which is commonly used as a catalyst and metal for electroplating. Cobalt also forms complex ions with ligands such as H_2O and NH_3 to give various coloured octahedral complexes such as [Co(H_2O)_6]^{2+} and [Co(NH_3)_6]^{2+} which are pink and yellow respectively. \end{array}$	
(a) The ligand exchange in octahedral complexes is one of the most extensively studied reactions in transition metals.	
An example of a ligand exchange reaction involving cobalt-(II) ions is:	
$\underline{[Co(H_2O)_6]^{2+} + 6NH_3} \rightleftharpoons \underline{[Co(NH_3)_6]^{2+} + 6H_2O}$	
(i) Explain why cobalt forms coloured complexes. [3]	
 The presence of ligands causes the energy level of the five 3d orbitals to be split into two different levels (crystal field splitting). 	(Formatted: Font: 11 pt
 The energy difference, <u>AE</u>, corresponds to wavelengths in the visible spectrum. 	Formatted: Font: 11 pt
When light energy is absorbed by the substance, an electron is promoted from a d orbital of	Formatted: Font: 11 pt
 <u>lower to one of higher energy (d-d transition)</u> <u>Unabsorbed wavelengths are transmitted and the colour of the complex is complementary</u> to the colour absorbed. 	
[3] for 4 marking points	Formatted: Font: 11 pt, Font color: Red
[2] for 3 marking points	
[1] for 2 marking points	Formatted: Font: 10.5 pt
(ii) Suggest why $[Co(NH_3)_6]^{2+}$ is of a different colour from $[Co(H_2O)_6]^{2+}$. [1]	
• Different ligands give rise to a difference in the splitting of the <i>d</i> -orbitals, cause a different	Formatted: Font: 11 pt
energy gap (ΔE).	Formatted: Font: 11 pt
A different wavelengths of visible light is absorbed and thus a different wavelength of light is	Formatted: Font: 11 pt
observed.	
[1] for 2 marking points	Formatted: Font: 11 pt, Font color: Red
(iii) A student wishes to investigate the kinetics of the ligand exchange reaction of	
$[Co(H_2O)_6]^{2+}$ to form $[Co(NH_3)_6]^{2+}$ by using a spectrometer. This machine measures the amount of light that is absorbed when a specific wavelength of visible light is shone through a few cm ³ of the coloured solution. It does this by comparing the amount of light passing through the sample with the amount of light passing through the pure solvent.	
The spectrometer is set to use the wavelength of light that is absorbed most strongly by the complex ion. The amount of light absorbed is expressed as an absorbance value. The more concentrated the solution, the higher the absorbance value. The temperature of the sample in the spectrometer can be thermostatically controlled for reaction rate analysis for which the sample has to be kept at a constant temperature.	
Outline the experimental procedure on how the student would accurately determine	
the initial rate of the ligand exchange reaction at 5 °C.	
[3] H2 Chemistry 9729/03 NYJC J2/18 PX H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn (Dver



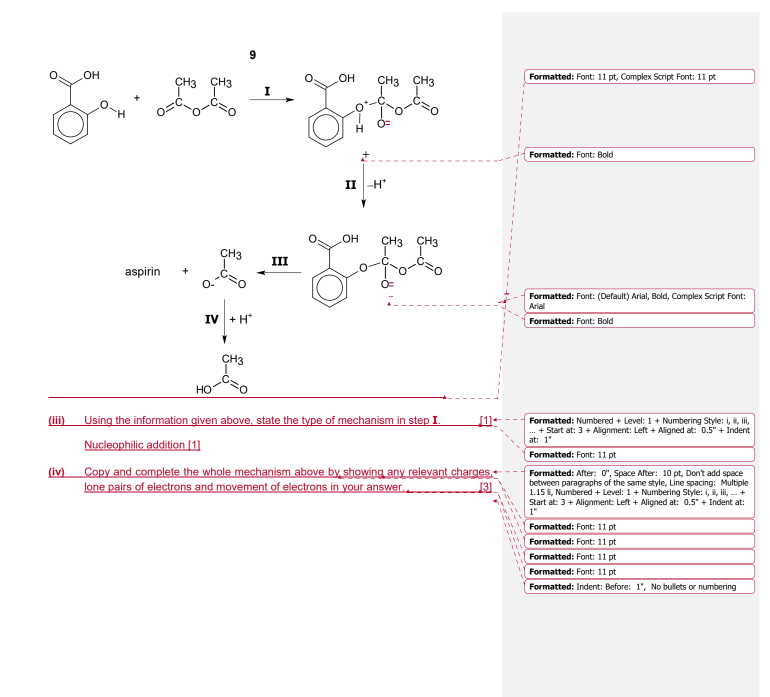
		5		
[1] fo	r intern	nediate and side product		
[1]10				
	<u>(v)</u>	State the rate equation for the above ligand exchange reaction. [1]		
		rate = 1/ [[Co/(1,10),124] [4]		
		<u>rate = k [[Co(H₂O)₆]²⁺] [1]</u>	<[Formatted: Font: 11 pt
				Formatted: Indent: Before: 0.5", First line: 0.5"
	<u>(vi)</u>	Hence, predict and explain the effect on the rate of reaction, if any, when the		
		ammonia ligand is replaced by with a fluoride ion.		
		[1]		
		The rate of reaction remains constant as the rate is independent of the incoming		Formatted: Font: 11 pt
		ligand. [1]	ς,	Formatted: Indent: Before: 1"
		<u>udeneri ul</u>		
(b)	[Co(H	$[_2O)_6]^{2+}$ can also undergo ligand exchange reactions with TMEDA to form		
(0)		$\underline{y}_{\underline{y}}$ MEDA) ₃] ²⁺ .		
	<u> </u>			
		Ň		
	By co	nsidering the entropy and enthalpy changes during the formation of [Co(TMEDA) ₃] ²⁺		
		[Co(H2O)6]2+ and that of [Co(NH3)6]2+ from [Co(H2O)6]2+, suggest how the standard		
	<u>Gibbs</u>	free energy change of the two reactions will compare in sign and in magnitude.		
	Hono	e prediet which reaction will be the more exectance up. Evalein your responsing		
[3]	Henc	e, predict which reaction will be the more spontaneous. Explain your reasoning.		
	[Co(H	$[_{2}O)_{6}]^{2+} + 6NH_{3} \rightleftharpoons [Co(NH_{3})_{6}]^{2+} + 6H_{2}O$		Formatted: Font: 11 pt
		<i>t</i>	57.	Formatted: Font: 11 pt
	<u>[Co(</u> H	$[20]_{6}]^{2+} + 3TMEDA_{2+} [Co(TMEDA)_{3}]^{2+} + 6H_{2}O_{2+}$. N.	Formatted: Font: 11 pt
	۸Ц fo	r formation of [Co(NH ₃) ₆] ²⁺ and [Co(TMEDA) ₃] ²⁺ is similar in magnitude and sign due to	17.	Formatted: Font: 11 pt
		reaking of 6 similar Co–O bonds and forming of 6 similar To–N bonds. [1]	, È	Formatted: Font: 11 pt
	<u></u>			
		or formation of [Co(TMEDA) ₃] ²⁺ would be more positive than that of [Co(NH ₃) ₆] ²⁺		
		use there is an increase in number of aqueous particles when [Co(TMEDA) ₃] ²⁺ is		
	<u>forme</u>	d, allowing more ways of arranging the particles. [1]		
	Since	ΔH for formation of [Co(NH ₃) ₆] ²⁺ and [Co(TMEDA) ₃] ²⁺ are similar in magnitude and		
		and ΔS for formation of [Co(TMEDA) ₃] ²⁺ is more positive than that of [Co(NH ₃) ₆] ²⁺ , ΔG		
		rmation of $[Co(TMEDA)_3]^{2+}$ would be more negative than that of $[Co(NH_3)_6]^{2+}$ and		
	-	e more spontaneous. [1]	1	Formatted: Font: 11 pt
		o r formation of [Co(NH₃)₀]²⁺ and [Co(TMEDA)₃]²⁺ is similar in magnitude due to+	×	Formatted: Justified, Indent: Before: 0.5"
	break	ing of similar Co-O bonds and forming of similar Co-N bonds. [1]		Formatted: Font: 11 pt
	AC fo	← <u>vr_formation_of_[Co(TMEDA)₃]²⁺</u> <u>would_be_more_positive_than_that_of_[Co(NH₂)₆]²⁺ </u>		Formatted: Justified
		se there is an increase in number of aqueous particles as [Co(TMEDA)₃] ²⁺ is formed.	1	Formatted: Font: 11 pt
		ng more ways of arranging the particles. [1]	N	Formatted: Font: 11 pt
			WOR	Formatted: Justified, Indent: Before: 0.5"
1		stry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn O	ver	



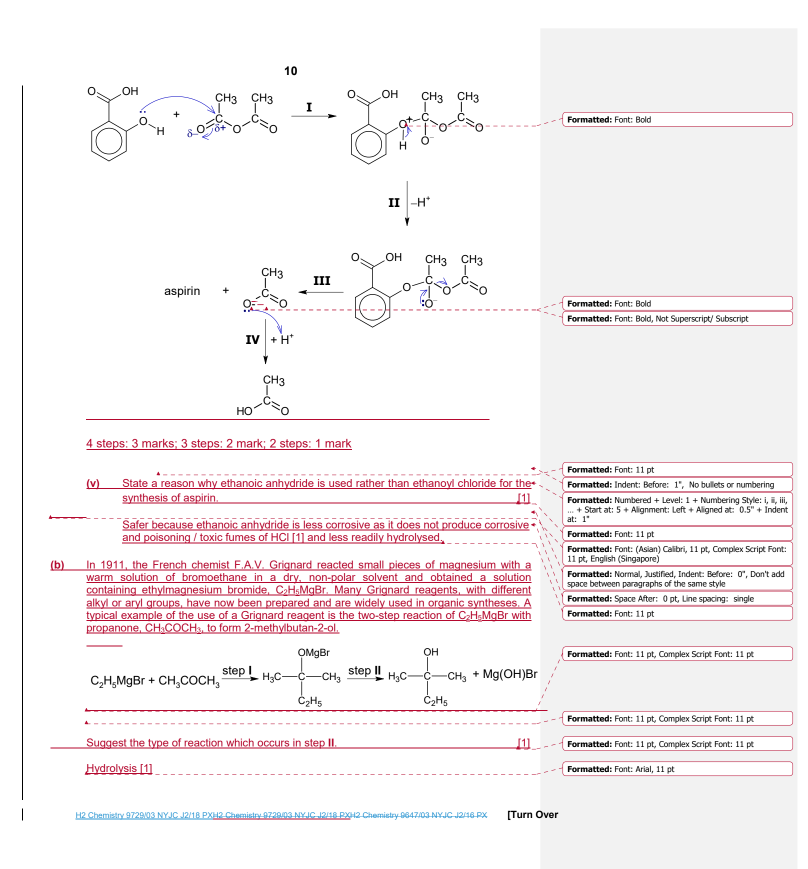
7	1	Formatted	
	11	Formatted	
<u>ΔG = – nFE_{cell} = – 2 x 96500 x 1.64 = – 317 000 J mol^{_1} = – 317 kJ mol^{_1}</u>	14	Formatted	
(iv) Use the Data Booklet to suggest the effect on the cell potential of this cell of adding		Formatted	
excess aqueous ammonia to the Co ²⁺ //Co half cell. Explain your answer. [1]	1.11.1	Formatted	
	111	Formatted	
$Co^{2+} + 2e \rightleftharpoons Co$ $E^{\odot} = -0.28 \vee$		Formatted	
$\underline{Co^{2^+} + 2e \rightleftharpoons Co} \qquad \underline{E^{\bullet}} = -0.28 \text{ V}$ $\underline{[Co(\text{NH}_3)_6]^{2^+} + 2e \rightleftharpoons Co + 6\text{NH}_3} \qquad \underline{E^{\bullet}} = -0.43 \text{ V}$	N. N.	Formatted	
		Formatted	
In the presence of ammonia ligands, Co ²⁺ undergoes ligand exchange to form a		Formatted	[
more stable complex ion of $[Co(NH_3)_6]^{2+}$. The $E^{\bullet}(Co^{2+} Co)$ becomes more negative		Formatted	[]
and the Ecell becomes more positive.	- 100 1111	Formatted	[]
	11001111	Formatted	[]
<u>Or</u>	- 1 1 MIL 11	Formatted	(]
	1.0.10011	Formatted	[]
The formation of [Co(NH ₃) ₆] ²⁺ decreases the concentration of Co ²⁺ . This caused the		Formatted	<u> </u>
position of equilibrium for $Co^{2+} + 2e \rightleftharpoons Co$ to shift left. The $E^{\diamond}(Co^{2+} Co)$ becomes	1 1 10 100	Formatted	
more negative and the Ecell becomes more positive.	1.1.0.00	Formatted	[[
		Formatted	[[.
		Formatted	[]
[Total: 242]	1 10 1 1 1 1		
	ь 1 Wi 111	Formatted]]
Answer any four guestions.	110011	Formatted	[[
	1111001	Formatted	[
A policy is seen of the most widely used down in the world. It is a powerful englancie (noise	11 1 1 1011		
Aspirin is one of the most widely used drug in the world. It is a powerful analgesic (pain reliever) antipyretic (fever reducer) and anti-inflammatory drug		Formatted	
Aspirin is one of the most widely used drug in the world. It is a powerful analgesic (pain reliever), antipyretic (fever reducer) and anti-inflammatory drug.	-0.1.1.0011 -0.1.1.1011 -0.1.1.1011 -0.1.1.1011 -0.1.1.1.101 -0.1.1.1.101	Formatted]]
reliever), antipyretic (fever reducer) and anti-inflammatory drug.	40000000000000000000000000000000000000]]
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated	1000 1 1 11 - 1000 1 1 1	Formatted]]]]]
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85%	1000 1 1 11 - 1000 1 1 1	Formatted	[] [] []]
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid.	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted]]]] [] []]
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted]]]]]
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted) ۱) ۱) ۱ ۱
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted Formatted	۱) ۱) ۱) ۱) ۱] ۱]
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted Formatted Formatted	اَسَ) اِسَ) اَسَ) اَسَ) اَسَ) اَسَ) اَسَ)
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH COOH CH3 CH3 CH3 CH3 CH3 CH3 CH3 CH	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted Formatted Formatted	ا…) ا…) ا…) ا…) ا…) ا…) ا…) ا…) ا…) ا…)
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH COOH CH3 CH3 CH3 L COOH CH3 L CH3 L	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH COOH CH3 CH3 CH3 CH3 CH3 CH3 CH3 CH	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH COOH CH3 CH3 CH3 L COOH CH3 L CH3 L	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH COOH CH3 CH3 CH3 CH3 CH3 CH3 CH3 CH	1000 1 1 11 - 1000 1 1 1	Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH COOH CH3 CH3 CH3 CH3 CH3 CH3 CH3 CH		Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $	1000 1 1 11 - 1000 1 1 1	Formatted Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. COOH COOH CH3 CH3 CH3 CH3 CH3 CH3 CH3 CH		Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $		Formatted Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $		Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $		Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $		Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $		Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $		Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $		Formatted	
reliever), antipyretic (fever reducer) and anti-inflammatory drug. It is synthesised using 2-hydroxybenzoic acid and ethanoic anhydride. 8 – 10 drops of 85% phosphoric acid which catalyses the reaction is added. The reaction mixture is then heated under reflux for around fifteen minutes. The other product of this reaction is ethanoic acid. $ \begin{array}{c} $		Formatted	

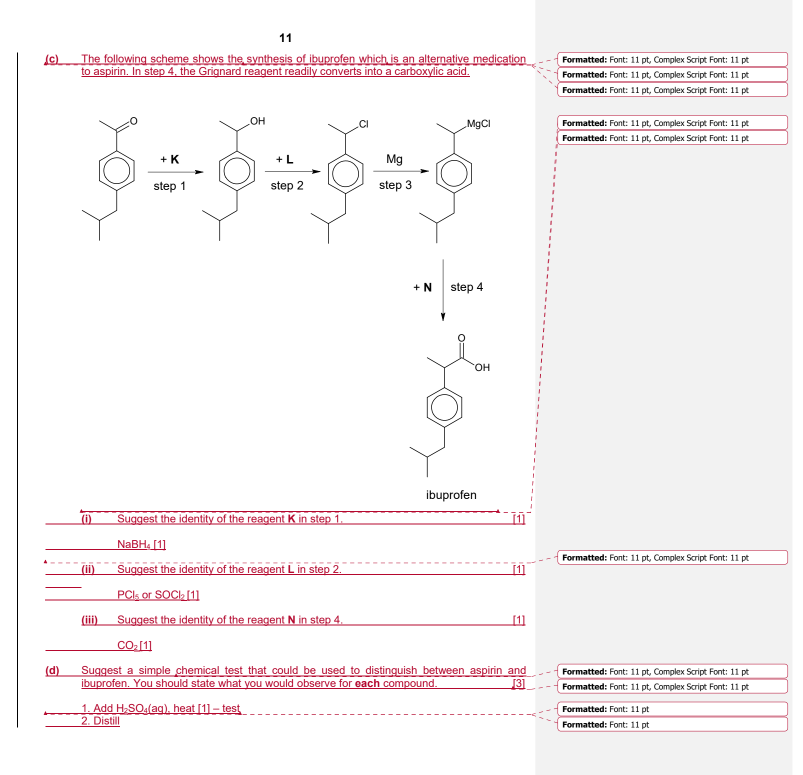


H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn Over



[Turn Over





H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX

[Turn Over

	12	
	СООН	Formatted: Font: 11 pt
	Дон	Formatted: Font: 11 pt
	Product obtained from aspirin: CH ₃ COOH.	
	Orange Br ₂ decolourises with aspirin; Orange Br ₂ remains for ibuprofen. [1]	Formatted: Font: 11 pt
Copp	er(I) sulfate, Cu ₂ SO ₄ , can be made from copper(I) oxide under non-aqueous conditions. On adding this salt to water, it immediately undergoes a disproportionation reaction.	Formatted: Indent: First line: 0"
	(i) Suggest, with a reason, the colour of copper(I) sulfate. [2]	
	(ii) Using suitable data from the Data Booklet, explain why the disproportionation reaction occurs, and write an equation for it. [3]	
(b) —	Palladium(II) salts can form square planar complexes. Successive addition of ammonia and hydrogen chloride to an aqueous palladium(II) salt produces, under different conditions, three compounds with empirical formula $PdN_2H_6Cl_2$. Two of these, A and B , are non-ionic, with $M_F = 211$. A has a dipole moment, whereas B has none. The third compound, C , is ionic, having $M_F = 422$, and contains palladium in both its cation and anion.	
	For each A, B and C, deduce a structure that fits the above data, explaining your reasons fully. fully. [6]	
(c)	Benzene ring is often represented as a structure that has a ring within the hexagon. Alternatively, chemists have also represented the structure of benzene in the following forms, known as resonance structures.	
	Benzene (two resonance forms)	
	The resonance relationship is indicated by the double headed arrow between them. The only difference between resonance forms is the placement of the pi electrons and non-bonding electrons.	
	Most aromatic compounds undergo electrophilic substitution. However aryl halides undergo a limited number of substitution reactions with strong nucleophiles.	
	An example of a reaction is as follows:	
	$A \longrightarrow X \longrightarrow A \longrightarrow Nu + X^{-}$	
	where A is an electron withdrawing group and X is a halogen	
	The mechanism of this reaction has two steps: addition of the nucleophile elimination of the halogen leaving group 	
1		

13 3 Step 1 involves the addition of the nucleophile (Nu⁻). The Nu⁻attacks the carbon atom bonded to a halogen, causing the pi bond to break. A resonance stabilised carbanion with a new C-Nu bond is formed. The aromatic ring is destroyed in this step. Formatted: Justified, Indent: Before: 0.5" Nu Field Code Changed Formatted: Indent: Before: 0.5" Step 2 involves the loss of the halogen X, reforming the aromatic ring. Formatted: Indent: Before: 0.5", First line: 0" Formatted: Indent: Before: 0.5" Nu Formatted: Justified, Indent: Before: 0.5" **Field Code Changed** Formatted: Justified, Indent: Before: 0.5", First line: 0" Two other resonance structures of the intermediate in Step 1 are shown below: Formatted: Justified, Indent: Before: 0.5" Nu Nu **Field Code Changed** 7 Formatted: Indent: Before: 0.5" Copy the above diagram and draw the resonance structure, Z. In your answ Formatted: Indent: Before: 0.5", First line: 0" show any relevant charges, lone pairs of electrons and movement of electrons in forming Z. [2] Formatted: Justified, Indent: Before: 0.5" The reaction below shows the synthesis of compound, D. Formatted: Indent: Before: 0.5", First line: 0" Formatted: Font: (Default) Arial, Complex Script Font: Arial N(CH₃)₂ N(CH₃)₂ Formatted: Indent: Before: 0.5" O_2N Formatted: Font: (Default) Arial, Complex Script Font: Arial 02 potassium metal + ΗÇI CI но D Formatted: Font: (Default) Arial, Complex Script Font: Arial Suggest the role of potassium metal in the reaction. 111 (ii) Formatted: Indent: Before: 0.5", First line: 0" Formatted: Indent: Before: 0.5" (iii) Use the information given above to draw out the full mechanism for the reaction that+ Formatted: Indent: Before: 0.5", First line: 0" forms D, labelling the slow and fast steps. In your answer, showing any relevant charges, lone pair of electrons and movement of electrons [3] Formatted: Indent: First line: 0", After: 0" Describe and explain the relative ease of hydrolysis followin (d) Formatted: Indent: Before: 0.5", First line: 0", Tab stops: chloro-compounds -[3] Not at 0.49 COCI CH₂CI CI Formatted: Font: (Default) Arial, 11 pt, Complex Script Font: Arial, 11 pt H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn Over

[Total: 20]

1

14

Formatted: Indent: First line: 0"

2 Sulfuric acid, H₂SO₄, can behave as an acid, an oxidising agent or as a dehydrating agent in various reactions.

(a) Draw a diagram to illustrate the shape of pure sulfuric acid and indicate the bond≁ angle about the sulfur atom. [2]

(b) The Contact Process is used for the manufacture of sulfuric acid. One of thereactions that takes place is the following reversible reaction:

 $\frac{2SO_2(g) + O_2(g) \cancel{4} 2SO_3(g)}{\Delta H = -197 \text{ kJ mol}^4}$

Sulfur dioxide and oxygen in a 2:1 molar ratio at a total initial pressure of 3 atm is passed over a catalyst in a fixed volume vessel at 400 °C. When equilibrium is established, the percentage of sulfur trioxide in the mixture of gases is found to be 30%.

(i) Write an expression for the equilibrium constant, K_e, of the reaction.

(ii) Calculate the value of K_e at 400 °C, stating its units.

(iii) How would the percentage conversion of SO₂ into SO₃ be affected when the+ pressure is raised? Explain. [2]

State the observations, if any, to indicate the differences in their reaction when water is added to each solid followed by dilute sulfuric acid. [4]

(d) Sulfur dioxide is a major pollutant from sulfuric acid plants. The SO₂-emitted into the atmosphere is oxidised in the air, which then reacts with water to form sulfuric acid, hence causing acid rain:

 $\frac{2SO_2(g) + O_2(g) + 2H_2O(l)}{\Delta H_4} \rightarrow \frac{2H_2SO_4(l)}{\Delta H_4}$

Using the data below and data from (b), construct an energy cycle to calculate

(i) the enthalpy change of formation of SO₂(g), and hence

(ii) the enthalpy change of reaction, ΔH_1 for the above reaction.

Enthalpy change of formation of H ₂ O(I)	– –286 kJ mol^{–1}
Enthalpy change of formation of H ₂ SO ₄ (I)	<u> </u>
Enthalpy change of formation of SO ₃ (g)	<u> </u>

Formatted: Justified, Indent: Before: 0.5", After: 0	"
Formatted: Indent: First line: 0", After: 0"	

Formatted: Indent: First line: 0", Automatically adjust right indent when grid is defined, Allow hanging punctuation, Adjust space between Latin and Asian text, Adjust space between Asian text and numbers, Font Alignment: Auto

Formatted: Indent: First line: 0" Formatted: Font: Arial

[1] - - -

Formatted: Indent: Before: 0.5"
Formatted: Indent: Before: 0.5", No bullets or numbering
Formatted: Indent: Before: 0.5"
Formatted: Indent: Before: 0.5", No bullets or numbering
Formatted: Indent: Before: 0.5"
Formatted: Indent: Before: 0.5", No bullets or numbering
Formatted: Indent: First line: 0", After: 0"
Formatted: Indent: Before: 0.5", First line: 0",

Automatically adjust right indent when grid is defined, Allow hanging punctuation, Adjust space between Latin and Asian text, Adjust space between Asian text and numbers, Font Alignment: Auto

Formatted: Indent: Before: 0.5", After: 0"

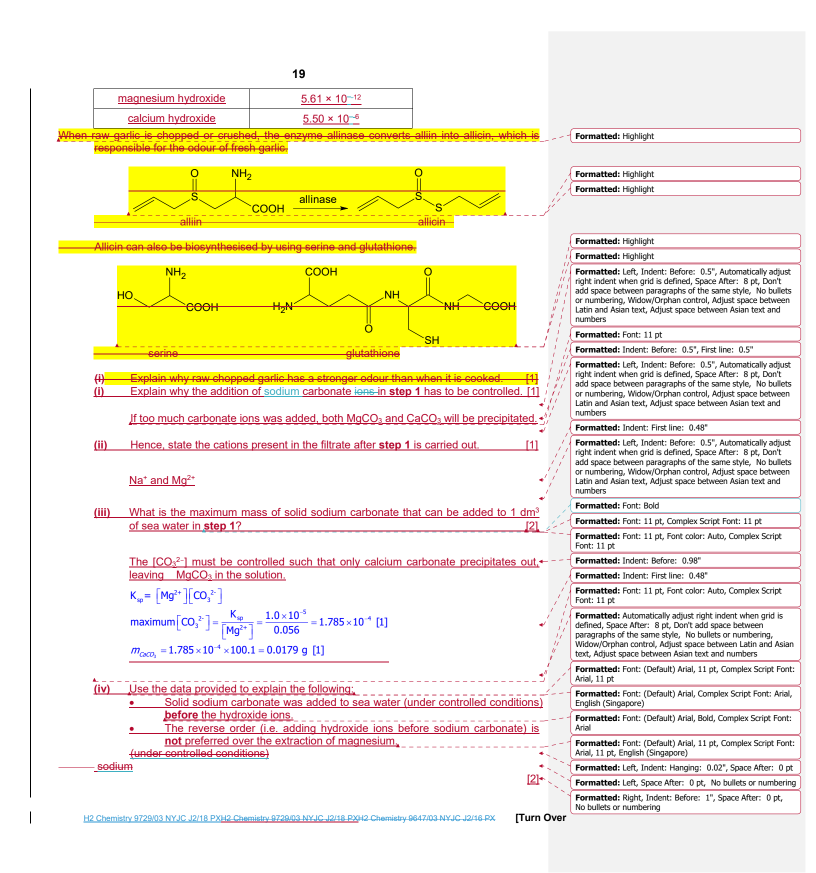
Formatted: Indent: Before: 0.5", Automatically adjust right indent when grid is defined, Allow hanging punctuation, Adjust space between Latin and Asian text, Adjust space between Asian text and numbers, Font Alignment: Auto

H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn Over

16 Alcohols react with concentrated sulfuric acid at high temperatures to form alkenes. (e) common side reaction that can happen is the formation of ethers, which is also catalysed by concentrated sulfuric acid. R Formatted: Font: Arial, 11 pt, Complex Script Font: 11 pt R R conc H₂SO H_O 2 R heat Ŕ Ŕ Ŕ R R Field Code Changed R conc H₂SO₄ 10 2 R heat Ŕ Ŕ Ŕ Formatted: Add space between paragraphs of the same The mechanism occurs via 3 steps: style Step Formatted: Justified, Add space between paragraphs of the same style An acid base reaction in which H* from H₂SO₄ protonates the oxygen atom in alcohol. This step is very fast and reversible. Formatted: Add space between paragraphs of the same style Step 2: molecule functions as the nucleophile and attacks the product from Formatted: Space Before: 0 pt, After: 0 pt, Add space between paragraphs of the same style step 1. The C-O bond is cleaved and a water molecule leaves the molecule. This creates R Field Code Changed R H_{-intermediate} an oxonium Formatted: Add space between paragraphs of the same style Step 3: Another acid base reaction in which the proton in the oxonium ion is removed by a suitable base (in this case a water molecule) to give the ether product. This step is very fast and reversible Formatted: Indent: Before: 0.5", Add space between paragraphs of the same style (i) Draw the ether formed when cyclopentanol undergoes the above reaction. Formatted: Indent: Before: 0.5", Automatically adjust right indent when grid is defined, Space After: 0 pt, Add space OH between paragraphs of the same style, No bullets or numbering, Allow hanging punctuation, Adjust space between Latin and Asian text, Adjust space between Asian text and numbers, Font Alignment: Auto Formatted: Justified, Indent: Before: 0.5", Add space between paragraphs of the same style cyclopentanol Field Code Changed 伯 Draw out the full mechanism for the reaction between two cyclopentanol molecules+ (iii) Formatted: Indent: Before: 0.5", Automatically adjust right to form an ether. In your answer, show any relevant charges, lone pairs of electrons and indent when grid is defined, Add space between paragraphs of the same style, No bullets or numbering, Allow hanging movement of electrons -[3] punctuation, Adjust space between Latin and Asian text, Adjust space between Asian text and numbers, Font [Total: 20] Alianment: Auto Formatted: Indent: First line: 0", After: 0" Formatted: Justified, Indent: Before: 0.5", After: 0" H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn Over

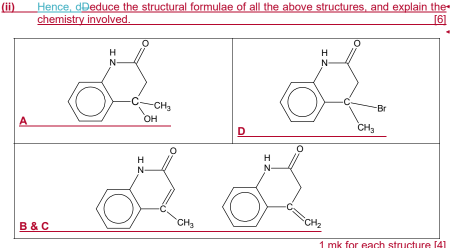


 Magnesium is present as dissolved magnering directly extracted from sea water. There is 	<u>sium ions in sea water and is</u> enough magnesium dissolvec	the only metal	
oceans to supply all of our magnesium needs			Formatted: Font: (Default) Arial, Complex Script Font: Arial, English (Singapore), Highlight
Garlic contains many amino acids, minerals and er compounds like alliin, allicin and ajoene. Th	ne sulfur compounds are respo		Formatted: English (United States)
garlic's pungent odour and many of its medic			
(a) <u>Apart_from_magnesium_ions, the tw</u> water are sodium and calcium ions,	o other most abundant cation	s found in sea	Formatted: Font: (Default) Arial, Complex Script Font: Arial Formatted: Normal, Indent: Before: 0", Hanging: 0.5",
Magnesium can be extracted from sea water	by the following steps:	· · · · · · · · · · · · · · · · · · ·	Space After: 0 pt, No bullets or numbering, Widow/Orphan control
Step 1	Step 2		Formatted: Font: (Default) Arial, Complex Script Font: Arial, Highlight
1. controlled addition of Na ₂ CO ₃ (s)	filtroto 1. addition of OH⁻(aq)	Ma(OH) reaidu	Formatted: Indent: First line: 0.25"
sea water2	filtrate 2. filter	Mg(OH) ₂ residue	Tormatted. Fond: 11 pt, complex Selipt Fond: 11 pt
		Step 3	Formatted: Font: (Default) Arial, 11 pt, Complex Script Font: Arial, 11 pt
		heat 🔶	Formatted: Justified, Indent: Before: 0.25", First line: 0.25", Space After: 0 pt, Line spacing: single
Mg Step 5 electrolysis	MgCl ₂ Step 4 HCl(g), heat unc anhydrous cond		
sea water $\frac{1. \text{ controlled addition of Na}_2\text{CO}_3(s)}{2. \text{ filter}} \text{ filtr}$	ate 1. addition of OH (aq) Mg(C 2. filter Step 3 heat	DH) ₂ residue	
Mg Step 5 electrolysis	MgCl ₂ → Step 4 → MgO HCl(g), heat under anhydrous conditions		
Concentration of common ions in sea water:			
jon concentr	ration / mol dm ⁻³		Formatted: Font: Bold
magnesium	<u>0.056</u>		Formatted: Font: Symbol, Bold Formatted: Font: Bold
calcium	<u>0.010</u>		Formatteu: Font. Bolu
sodium	<u>0.457</u>		
chloride	<u>0.535</u>		
The numerical values of solubility products a	re given below:	<u>ـ</u> ـ	
compound value of s	olubility product		Formatted: Font: Bold
magnesium carbonate <u>1.</u>	<u>00 × 10⁻⁵</u>		
calcium carbonate 8	. <u>70 ×10^{−9}</u>		
H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NY	<u>'JC J2/18 PX</u> H2 Chemistry 9647/03 NYJC (J <u>2/16 PX</u> [Turn O\	/er



	20		
	Comparing the K_{sp} of the 2 carbonates, CaCO ₃ is less soluble and will be precipitated out first, leaving the Mg ²⁺ ions in solution, and can be further purified via other steps. [1]		- Formatted: Indent: Before: 1"
	<u>If hydroxide ions was added before carbonate ions, $Mg(OH)_2$ having a smaller K_{sp} value than $Ca(OH)_2$ will be less soluble and precipitate out first, together with other solid impurities in sea water, leading to an impure product. [1]</u>		
	(v) Calculate the minimum pH of the hydroxide solution required for precipitation		Formatted: Right, Indent: Before: 1", Space After: 0 pt, No bullets or numbering
	of magnesium hydroxide in step 2 if an equal volume of hydroxide ions was added to the filtrate. Give your answer to 2 decimal places. [2]		- Formatted: Font: Bold, Font color: Auto
	Assume [Mg ²⁺] = 0.056 mol dm ⁻³ (from table)	·	Formatted: Indent: First line: 0.5"
	After adding equal volume of hydroxide ions, $[Mg^{2+}]_{new} = \frac{0.056}{2} = 0.028 \text{ mol dm}^{-3}$		
	$K_{sp} = \left[Mg^{2+}\right] \left[OH^{*}\right]^{2}$	1	(Formatted: Indent: First line: 0.48"
	$\left[OH^{-}\right] = \sqrt{\frac{K_{sp}}{\left[Mg^{2^{+}}\right]}} = \sqrt{\frac{5.61 \times 10^{-12}}{0.028}} = 1.415 \times 10^{-5} [1]$		
	$\begin{bmatrix} OH^{-} \end{bmatrix} required = 1.415 \times 10^{-5} \times 2$		
	$pOH=-log(1.415 \times 10^{-5} x 2) = 4.55$ pH = 9.45 [1] 2dp ans		
<u>(i)</u>	Write the equations that occur during the electrolysis of magnesium chloride in Sstep 5 . State clearly the reactions that occur at the cathode and the anode, and include state symbols.	 ```````	Formatted: Default, Indent: Before: 0.5", Hanging: 0.5", Automatically adjust right indent when grid is defined, Space After: 0 pt, Line spacing: single, No bullets or numbering, Adjust space between Latin and Asian text, Adjust space between Asian text and numbers
	[2]		Formatted: Font: Not Bold
	Cathode: $Mg^{2+}(l) + 2e \rightarrow Mg(l)$ [1] Anode : $2Cl^{-}(l) \rightarrow Cl_{2}(g) + 2e$ [1]		- Formatted: Indent: Before: 0.73", First line: 0.25"
<u>(ii)</u>	In a factory, a current of 95 kA was passed through a suitable setup for 24 hours.		Formatted: Indent: Before: 0.5", Hanging: 0.5", Space After: 0 pt, No bullets or numbering
	Assuming that the procedure is 90% efficient, calculate the mass of Mg that can be produced. [2]	t	Formatted: Space After: 0 pt, Numbered + Level: 1 + Numbering Style: i, ii, iii, + Start at: 2 + Alignment: Left + Aligned at: 0.5" + Indent at: 1"
	Q= It = nzF		Formatted: Indent: Before: 1", Space After: 0 pt, No bullets or numbering
	$\frac{95 \times 1000 \times 24 \times 60 \times 60}{100} \times 90 = n \times 2 \times 96500$		Formatted: Indent: Before: 0.5"
	$n = 38.28 \times 10^4$ [1] m= $38.28 \times 10^4 \times 24.3 = 930,000g = 930kg$ [1]		
(iii)	Give a reason why electrolysis of magnesium chloride is preferred to that of		Formatted: Space After: 0 pt, Numbered + Level: 1 + Numbering Style: i, ii, iii, + Start at: 2 + Alignment: Left +
<u>(m)</u>	magnesium oxide in this industrial process. [1]		Aligned at: 0.5" + Indent at: 1" Formatted: Default, Indent: Before: 0.5", Hanging: 0.5",
	MgCl ₂ has a lower melting point than MgO, hence the electrolysis can be carried out at a lower temperature, saving operation costs. [1]		Automatically adjust right indent when grid is defined, Line spacing: single, No bullets or numbering, Adjust space between Latin and Asian text, Adjust space between Asian text and numbers
	stry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX	Over	Formatted: Indent: Before: 1"

21	
A is an organic compound. When 0.4678 g of an organic compound A was vaporated in a syringe, the volume of the vapour produced after correction to s.t.p was 60 n ³ . On heating with aqueous sodium hydroxide, A gives a compound that dissolves in	Formatted: Indent: Before: 0", Hanging: 0.5", Space Afte 0 pt, No bullets or numbering
ater.	
reacts with aluminium oxide to give two products B and C . Both B and C react with HBr give the same product D . D exhibits enantiomerism and exists as a pair of enantiomers.	Formatted: Indent: Before: 0.5", Space After: 0 pt
gives E when reacted with lithium aluminium hydride in dry ether.	Formatted: Indent: First line: 0.25", Space After: 0 pt
OH CH ₃	Formatted: Centered, Indent: First line: 0.25"
	Formatted: Space After: 0 pt, Numbered + Level: 1 + / Numbering Style: i, ii, iii, + Start at: 1 + Alignment: Left +
E	/ Aligned at: 0.5" + Indent at: 1"
	Formatted: Indent: Before: 1", Space After: 0 pt, No bullets or numbering
Prove that the molarecular mass of A is 177 g mol ⁻¹ .	Formatted: Indent: First line: 0.25"
[1]	Formatted: Lowered by 20 pt
*/	Formatted: Superscript
From Data Booklet, at s.t.p. $V_m = 22.7 \text{ dm}^3 \text{ mol}^{-1}$ at 10 ⁵ Pa and 273 K	Formatted: Font: Italic, Complex Script Font: Italic
Either	Formatted: Font: Arial
$10^5 \times 60 \times 10^{-6} = \frac{0.468}{M} \times 8.31 \times 273$	Formatted: Font: 10 pt, Font color: Auto, Complex Script
$M = 176.9 = 177 \text{ g mol}^{-1} [1]$	Formatted: Indent: Before: 1", Space After: 0 pt, Bulleted
	Image: Property of the second seco
$\frac{\text{Or}}{n(\text{A})} = 60 / 22700 = 2.643 \times 10^{-7} \text{ mol}$	Formatted: Font: 10 pt, Font color: Auto, Complex Script
$\underline{M_{\rm t}(A)} = 0.468 / 2.643 \times 10^{-7} = 177.06 \approx 177 \text{ g mol}^{-1}$	Formatted: Indent: Before: 1", Space After: 0 pt, Bulleted
$\left(\begin{array}{c} \eta_{j} \\ \eta_{j} \end{array} \right)$	Formatted: Font: 10 pt, Complex Script Font: 10 pt
Comments	Formatted: Indent: Before: 1.25", Space After: 0 pt
Generally well-done.	Formatted: Indent: Before: 1", Space After: 0 pt, Bulleted
• <u>Common mistake include:</u> <u>Using incorrect V_m such as V_m = 24 dm³ or V_m = 22.4 dm³ • V_m</u>	+ Level: 3 + Aligned at: 1.25" + Indent at: 1.5" Formatted: Indent: Before: 1.25", First line: 0.25", Space
• A small number of students did not use the numerical data of 0.468 g and 60 cm ³ to τ_{1}^{\prime}	/ After: 0 pt / Formatted: Indent: Before: 1.25", Space After: 0 pt
prove the molar mass of A. These students used the following approach, which is NOT	Formatted: Font: 10 pt, Complex Script Font: 10 pt
acceptable for this question.	Formatted: Indent: Before: 1.25", First line: 0.25", Space
1 mol of A undergoes reduction to give 1 mol of E . molar mass of E = 163.0 g mol ⁻¹	Formatted: Font: 10 pt, Bold, Complex Script Font: 10 pt, Bold
<u>A has amide.</u> molar mass of $\mathbf{A} = 163.0 - 2(1.0) + 16.0 = 177.0 \text{ g mol}^{-1}$	Formatted: Indent: Before: 1.25", Space After: 0 pt
	Formatted: Font: 10 pt, Complex Script Font: 10 pt
Note that you are expected to use the numerical data of 0.468 g and 60 cm ³ to prove+1. the molar mass of A.	Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt
A	Formatted: Indent: Before: 1", Space After: 0 pt



1/2 mk for each point (max of 2 mks):

- 1) From the molar mass of 177 g mol⁻¹, the molecular formula is likely to be $C_{10}H_{11}O_2N$.
- 2) A undergoes basic hydrolysis with NaOH (aq) to give -COO⁻ (-COOH + NaOH) and -NH₂. -COO⁻ is an ionic salt that is soluble indissolves in water by forming ion-dipole interactions with water. \Rightarrow A contains an amide group to give an amine and carboxylic salt when hydrolysed.

Some students wrote that the compounds form from the hydrolysis of A dissolves in water to via formation of hydrogen bonds with water. It may seem correct, but note that A contains an amide in a cyclic structure, hence after hydrolysis, there will be only one single product

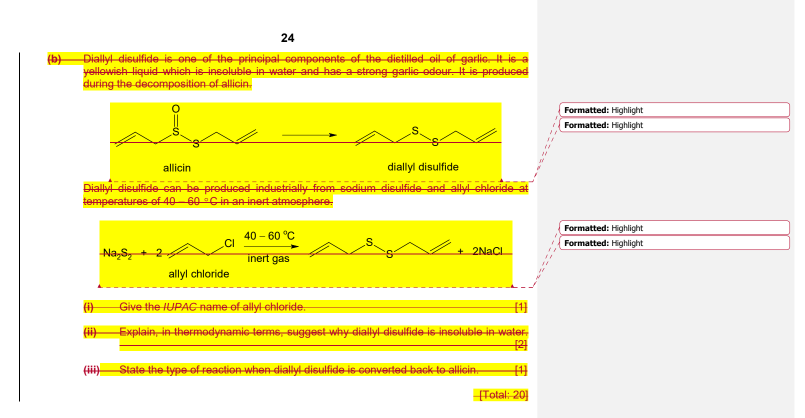
- 3) A undergoes elimination with Al_2O_3 to give alkenes **B** and **C** \Rightarrow **A** contains an alcohol aroup
- 4) The amide group in A undergoes reduction with LiAIH₄ to give an amine in E
- Alkenes B and C undergo electrophilic addition with HBr to give halogenoalkane/alkyl halide/bromoalkane D.
- 6) D contains a chiral carbon and hence exists as a pair of enantiomers.

Out of the above 6 statements, the following key points are expected to gain credit. <mark>∖ undergoes (basic) hydrolysis with NaOH ⇒,A contains an amide group</mark> **A** undergoes elimination with Al₂O₃ to give alkenes **B** and **C** \Rightarrow A contains aroup The amide group in A undergoes reduction with LiAIH4 to give an amine in E Alkenes B and C undergo electrophilic addition with HBr halogenoalkane/alkyl halide/bromoalkane D. OR D contains a chiral carbon and hence exists as a pair of enantiomers 4 bullet pts correct - 2 marks 2 bullet pts correct (minimum) - 1 mark

H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn Over

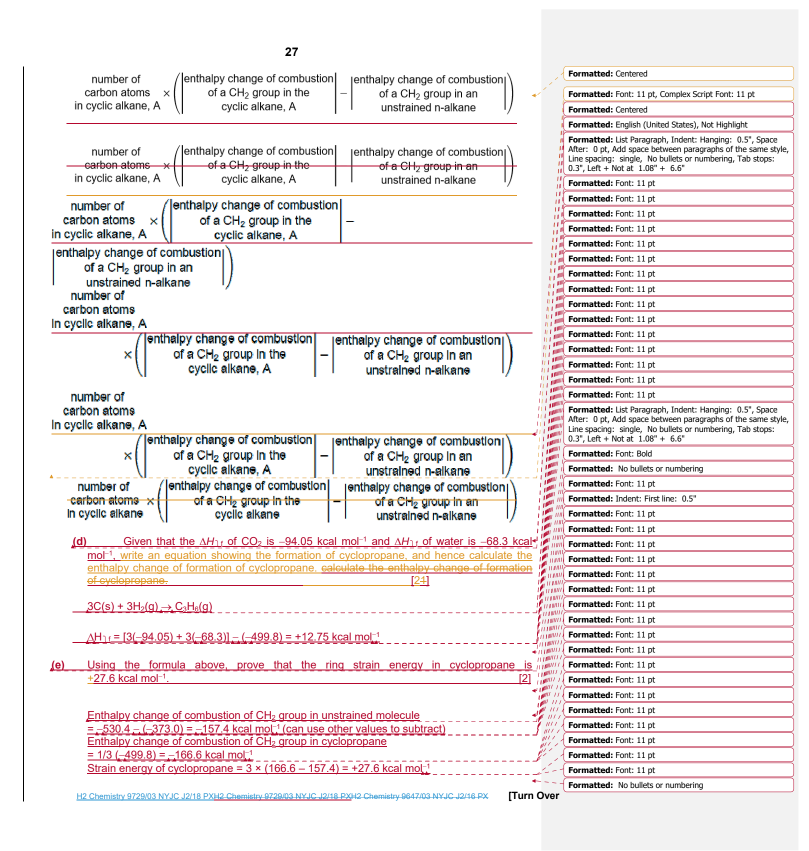
Formatted: Space After: 0 pt Formatted: Indent: Before: 1", Space After: 0 pt, No bullets or numbering Formatted: Font: Not Bold, Complex Script Font: Bold Formatted: Indent: Before: 0.75", No bullets or numbering Formatted: Complex Script Font: Bold Formatted: Indent: Before: 1", No bullets or numbering Formatted: Highlight Formatted: Indent: Before: 1", Space After: 0 pt, Bulleted + Level: 3 + Aligned at: 1.25" + Indent at: 1.5", Widow/Orphan control Formatted: Highlight Formatted: Highlight Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt, Highlight Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt, Highlight Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt, Highlight Formatted: Highlight Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt Formatted: Indent: Before: 1" Formatted: Font: 10 pt, Bold, Underline, Font color: Auto, Complex Script Font: 10 pt, Bold Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt Formatted: Font: Not Bold Formatted: Indent: Before: 1.25", No bullets or numbering Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt Formatted: Indent: Before: 1" Formatted: Font: 10 pt, Bold, Underline, Font color: Auto, Complex Script Font: 10 pt, Bold Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt Formatted: Font: 10 pt, Bold, Underline, Font color: Auto, Complex Script Font: 10 pt, Bold Formatted: Font: 10 pt, Underline, Font color: Auto, Complex Script Font: 10 pt, Bold Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt Formatted: List Paragraph, Left, No bullets or numbering Formatted: Font: Bold, Complex Script Font: Bold Formatted: Indent: Before: 1.25", No bullets or numbering Formatted: Font: 10 pt, Underline, Font color: Auto, Complex Script Font: 10 pt Formatted: Indent: Before: 1" Formatted: Font: 10 pt, Bold, Underline, Font color: Auto, Complex Script Font: 10 pt Formatted: Font: 10 pt, Font color: Auto, Complex Script Font: 10 pt Formatted: Font: 10 pt, Complex Script Font: 10 pt Formatted: Indent: Before: 1", Space After: 0 pt, No bullets or numbering

	23		
<u>(iii)</u>	State the type of isomerism exhibited by B and C . Explain why B and C both give- the same product D when reacted with HBr. [2].		Formatted: Space After: 0 pt, Numbered + Level: 1 + Numbering Style: i, ii, iii, + Start at: 1 + Alignment: Left Aligned at: 0.5" + Indent at: 1"
		1	Formatted: Font color: Blue
	Constitutional isomerism. [1]		Formatted: Indent: Before: 1"
	When alkenes B and C undergo electrophilic addition with HBr, both form the same carbocation. The carbocation will be attacked by Br ⁻ , which leads to the formation of D.		
	$ \underbrace{ \begin{array}{c} H \\ H $		Formatted: Indent: First line: 0.25"
	B or C carbocation		
	[1] for explanation & structure of carbocation		Formatted: Indent: Before: 0.5", First line: 0.5"
	<u>,[Total: 22]+</u>	~ - '	Formatted: Not Highlight
(ii)	When one molecule of serine reacts with one molecule of glutathione, it is possible to form two esters with different structural formulae. Draw the structural formula of each of these esters. [2]		Formatted: Right
(iii)	Draw the structural formulae of the products when glutathione is hydrolysed. [3]		
	Alliin has pK₂-values of 1.84 and 8.45.		Formatted: Right
(iv) -	Make use of these pKa values to suggest the major species present in solutions of alliin with the following pH values. [3] •pH 1 •pH 7 •pH 11 •pH 11		
	ť		Formatted: Not Highlight
v)	Calculate the pH of 0.10 mol dm ⁻³ solution of alliin. [1]		
(vi)	With reference to the pK _a values, identify the major species formed when 10 cm ³ of 0.10 mol dm ⁻³ NaOH is added to 10 cm ³ of 0.10 mol dm ⁻³ protonated alliin. Hence, deduce whether the solution is acidic, neutral or alkaline. [2]		
(vii) —	Sketch the pH-volume added curve you would expect to obtain when 30 cm ³ of 0.10 mol dm ³ -NaOH is added to 10 cm ³ of 0.10 mol dm ³ -protonated alliin. Briefly describe how you have calculated the various key points on the curve. [4]		
		·	Formatted: Highlight
2 Chemistr	y 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn (Over	

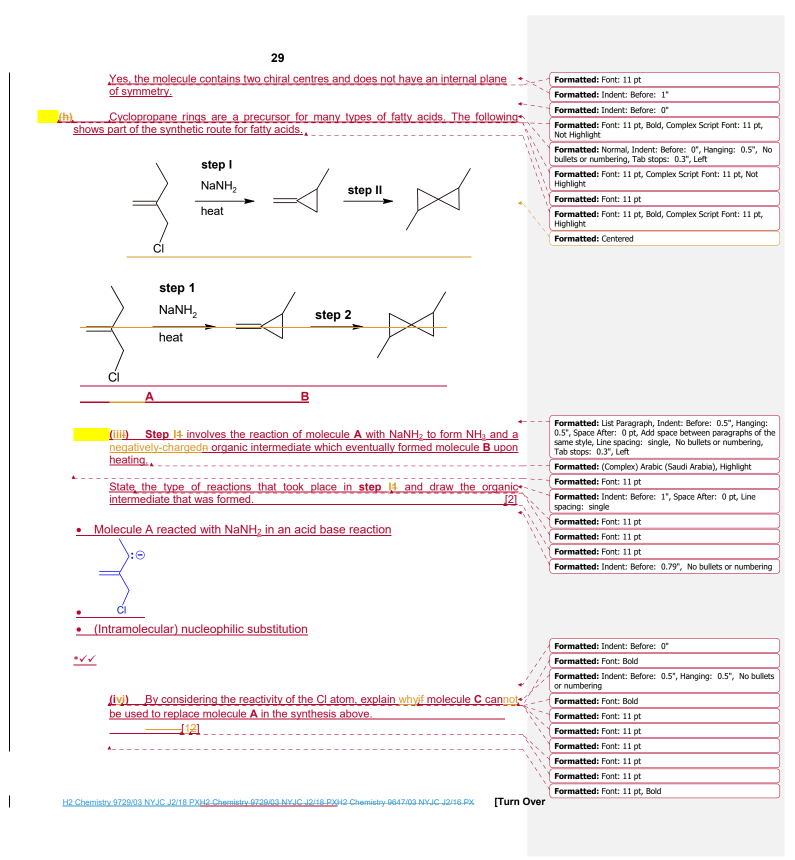


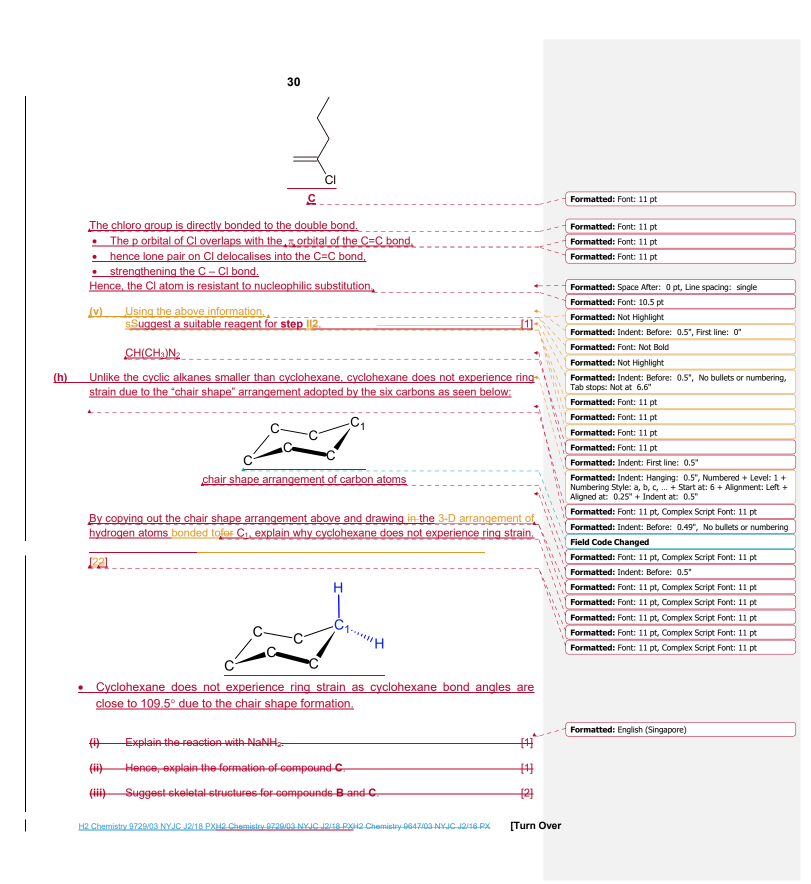
		Secti	on B				Formatted: Font: 11 pt
	Ar	nswer one questio	on from this section	<u>n.</u>			Formatted: Centered
							Formatted: Font: 10 pt
Cycloalkane	<u>s are a hom</u>	ologous series o alkanes are a	<u>f cyclic saturated</u>	hydrocarbo	ns with the gene	<u>ral</u> •	Formatted: Indent: Before: 0", Hanging: 0.49", Tab stop
		eneral formula C _n H		es or straig	gnt-chain saturat	<u>ea</u> ```	0.3", Left
nyarocarbor	is with the ge		<u>12n+2.</u>				Formatted: Font: 11 pt
[T	Enthalpy			Enthalpy	T	Formatted: Font: 11 pt, Bold, Complex Script Font: 11 pt, Bold, Italic, English (United Kingdom), Highlight
n-alkanes	<u>Boiling</u>	change of	cycloalkanes	<u>Boiling</u>	change of		Bold, Italic, English (Onited Kingdom), Highlight
<u>II-aikanes</u>	<u>point / °C</u>	<u>combustion /</u>	Cycloaikaries	point / °C	combustion /		
		<u>kcal mol⁻¹</u>			<u>kcal mol⁻¹</u>	_	
<u>ethane</u>	<u>–89</u>	<u>-373.0</u>	=	=	=	_	
propane	<u>-42</u>	<u>-530.4</u>	<u>cyclopropane</u>	<u>-33</u>	<u>-499.8</u>	_	
butane	<u>_1</u>	<u>–687.8</u>	<u>cyclobutane</u>	<u>12</u>	<u>-656.0</u>	_	
<u>pentane</u>	<u>36</u>	<u>-845.2</u>	<u>cyclopentane</u>	<u>49</u>	<u> </u>	_	
<u>hexane</u>	<u>69</u>	<u>-1002.6</u>	<u>cyclohexane</u>	<u>81</u>	<u>-944.6</u>		
<u>heptane</u>	<u>98</u>	<u>–1160.0</u>	<u>cycloheptane</u>	<u>119</u>	<u>–1108.3</u>		
_(a) _Expla _[1]	ain the term	homologous serie	<u>es".</u>				Formatted: Font: 11 pt, Complex Script Font: 11 pt, Not Highlight
						С. N.	Formatted: Normal, No bullets or numbering
Froup of organ							Formatted: Font: 11 pt, Complex Script Font: 11 pt
	ne functional					1	Formatted: Font: 11 pt
• Diff	er by a CH ₂ o	group				< <.	Formatted: Indent: Before: 0.55", Hanging: 0.25", Spac After: 0 pt, Line spacing: single
(b) Expl	ain the incre	ase in magnitude	es of both boiling	point and	enthalpy change	<u>of</u> ≁ ``	Formatted: Normal, No bullets or numbering
combustion	from ethane	to heptane.				3]] () ()	Formatted: Font: 11 pt, Bold
						NN Y	Formatted: Normal, Indent: Before: 0", Hanging: 0.5",
	s have simp	la molecular etri					
					ous dipole-induc	<u>ed</u> _ `\`\	bullets or numbering
	d-id) forces t	petween molecule	<u></u> <u>8</u>				Formatted: Format, indent. before: 0, Hanging: 0.5, bullets or numbering
• From e	id-id) forces t thane to her	petween molecule ptane, the electro	<u></u> <u>8</u>				bullets or numbering
From end hence id	id-id) forces t thane to hep d-id forces be	between molecule otane, the electro become stronger.	<u>es</u> In cloud gets big	ger and mo	re easily polarise		bullets or numbering Formatted: Font: 11 pt
From end hence id	id-id) forces t thane to hep d-id forces be	petween molecule ptane, the electro	<u>es</u> In cloud gets big	ger and mo	re easily polarise		bullets or numbering Formatted: Font: 11 pt Formatted: Font: 11 pt
From e hence id More er	id-id) forces t thane to hep d-id forces be hergy needec	between molecule otane, the electro become stronger.	<u>s</u> <u>n cloud gets big</u> forces, hence bo	ger and mo	re easily polarise		bullets or numbering Formatted: Font: 11 pt Formatted: Font: 11 pt
 From e hence id More er From et 	id-id) forces t thane to hep d-id forces be nergy needed hane to hept	between molecule otane, the electro ecome stronger. I to break the id-io	<u>is</u> on cloud gets big d forces, hence bo ∋ increases by a C	ger and mo biling point in CH ₂ group	<u>re easily polarise</u> Icreases	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Font: 11 pt
 From e hence is More er From et This res 	id-id) forces t thane to hep d-id forces be nergy needec hane to hept sults in more	between molecule btane, the electro ecome stronger. I to break the id-id ane, the molecule	<u>s</u> <u>on cloud gets big</u> d forces, hence bo ∋ increases by a C C – H bonds to b	ger and mo biling point ir CH ₂ group preak and m	<u>re easily polarise</u> Icreases	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Font: 11 pt
From e hence ic More er From et This res bonds to	id-id) forces t thane to hep d-id forces be nergy needec hane to hept sults in more	between molecule otane, the electro ecome stronger. I to break the id-ic ane, the molecule C – C bonds and	<u>s</u> <u>on cloud gets big</u> d forces, hence bo ∋ increases by a C C – H bonds to b	ger and mo biling point ir CH ₂ group preak and m	<u>re easily polarise</u> Icreases	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Font: 11 pt Formatted: Font: 11 pt
 From e hence is More er From et This res 	id-id) forces t thane to hep d-id forces be nergy needec hane to hept sults in more	between molecule otane, the electro ecome stronger. I to break the id-ic ane, the molecule C – C bonds and	<u>s</u> <u>on cloud gets big</u> d forces, hence bo ∋ increases by a C C – H bonds to b	ger and mo biling point ir CH ₂ group preak and m	<u>re easily polarise</u> Icreases	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Font: 11 pt
 From e hence is More er From et This res bonds to 	Id-Id) forces to thane to hep d-id forces be nergy needed hane to hept sults in more o form, hence	between molecule otane, the electro ecome stronger. I to break the id-id ane, the molecule C – C bonds and e more heat given	is on cloud gets big d forces, hence bo e increases by a C C – H bonds to b off during combu	ger and mo biling point in CH ₂ group break and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Font: 11 pt Formatted: Font: 11 pt
From e hence ic More er From et This res bonds to *********************************	d-id) forces b thane to hep d-id forces be nergy needed hane to hept sults in more o form, hence c molecules	between molecule btane, the electro ecome stronger. I to break the id-id ane, the molecule C – C bonds and e more heat given which contain of	Son cloud gets big d forces, hence bo ⇒ increases by a C C – H bonds to b off during combu arbon carbon trip	ger and mo biling point in CH ₂ group break and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Font: 11 pt Formatted: Font: 11 pt Formatted: Font: 11 pt
From e hence ic More er From et This res bonds to *********************************	d-id) forces b thane to hep d-id forces be nergy needed hane to hept sults in more o form, hence c molecules	between molecule otane, the electro ecome stronger. I to break the id-id ane, the molecule C – C bonds and e more heat given	Son cloud gets big d forces, hence bo ⇒ increases by a C C – H bonds to b off during combu arbon carbon trip	ger and mo biling point in CH ₂ group break and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt
From e hence ic More er From et This res bonds to *********************************	d-id) forces b thane to hep d-id forces be nergy needed hane to hept sults in more o form, hence c molecules	between molecule btane, the electro ecome stronger. I to break the id-id ane, the molecule C – C bonds and e more heat given which contain of	Son cloud gets big d forces, hence bo ⇒ increases by a C C – H bonds to b off during combu arbon carbon trip	ger and mo biling point in CH ₂ group break and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt
From e hence ic More er From et This res bonds to *********************************	d-id) forces b thane to hep d-id forces be nergy needed hane to hept sults in more o form, hence c molecules	between molecule btane, the electro ecome stronger. I to break the id-id ane, the molecule C – C bonds and e more heat given which contain of	Son cloud gets big d forces, hence bo ⇒ increases by a C C – H bonds to b off during combu arbon carbon trip	ger and mo biling point in CH ₂ group break and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt
From e hence ic More er From et This res bonds to *********************************	d-id) forces to thane to hep d-id forces be hergy needed hane to hept sults in more b form, hence c molecules series with f	between molecule btane, the electro ecome stronger. I to break the id-id ane, the molecule C – C bonds and e more heat given which contain of	is in cloud gets big d forces, hence bo increases by a C C – H bonds to b off during combu- carbon-carbon-trip i.e. ■C—R ₂	ger and mo biling point in CH ₂ group oreak and m istion.	re easily polarise creases ore C=O and O –	:d.	Formatted: Font: 11 pt
From et hence in More er From et This res bonds tr *********************************	d-id) forces to thane to hep d-id forces be hergy needed hane to hept sults in more o form, hence c molecules series with f	etween molecule otane, the electro ecome stronger. I to break the id-id ane, the molecule C – C bonds and e more heat given which contain c formula of C _n H _{2n-2}	is in cloud gets big d forces, hence bo increases by a C C – H bonds to b off during combu- carbon-carbon-trip i.e. ■C—R ₂ or alkyl or aryl gro	ger and mo biling point in CH ₂ group oreak and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Normal, Indent: Before: 0", Hanging: 0.5", bullets or numbering
From et <u>hence is</u> <u>More er</u> <u>From et</u> <u>This res</u> <u>bonds ts</u> */*/*/ es are organi homologous	id-id) forces to thane to hep d-id forces be hergy needed hane to hept sults in more o form, hence c molocules series with f whe ibit similar ch	between molecule otane, the electro ecome stronger. I to break the id-ic ane, the molecule C – C bonds and e more heat given which contain c formula of C_nH_{2n-2} R_1 —C re R_4 and R_2 = H	s n cloud gets big d forces, hence bo e increases by a C C – H bonds to b off during combu arbon carbon trip i.e. C — R ₂ or alkyl or aryl gro	ger and mo biling point in CH ₂ group preak and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Normal, Indent: Before: 0", Hanging: 0.5", bullets or numbering
From et <u>hence is</u> <u>More er</u> <u>From et</u> <u>This res</u> <u>bonds ts</u> */*/*/ es are organi homologous	id-id) forces to thane to hep d-id forces be hergy needed hane to hept sults in more o form, hence c molocules series with f whe ibit similar ch	between molecule otane, the electro- ecome stronger. I to break the id-ic ane, the molecule C – C bonds and e more heat given which contain c formula of C_nH_{2n-2} R_1-C= re R_1 and R_2 = H memical properties ith electrophiles i.	rss rss n cloud gets big rss forces, hence bo rss increases by a C C - H bonds to t rst off during combu- rst of rst	ger and mo biling point in CH ₂ group preak and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Normal, Indent: Before: 0", Hanging: 0.5", bullets or numbering
From et <u>hence is</u> <u>More er</u> <u>From et</u> <u>This res</u> <u>bonds ts</u> */*/*/ es are organi homologous	id-id) forces to thane to hep d-id forces be hergy needed hane to hept sults in more o form, hence c molocules series with f whe ibit similar ch	between molecule otane, the electro- ecome stronger. I to break the id-ic ane, the molecule C – C bonds and e more heat given which contain c formula of C_nH_{2n-2} R_1-C= re R_1 and R_2 = H memical properties ith electrophiles i.	s n cloud gets big d forces, hence bo e increases by a C C – H bonds to b off during combu arbon carbon trip i.e. C — R ₂ or alkyl or aryl gro	ger and mo biling point in CH ₂ group preak and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Normal, Indent: Before: 0", Hanging: 0.5", bullets or numbering Field Code Changed
From et <u>hence is</u> <u>More er</u> <u>From et</u> <u>This res</u> <u>bonds ts</u> */*/*/ es are organi homologous	id-id) forces to thane to hep d-id forces be hergy needed hane to hept sults in more o form, hence c molocules series with f whe ibit similar ch	between molecule otane, the electro- ecome stronger. I to break the id-ic ane, the molecule C – C bonds and e more heat given which contain c formula of C_nH_{2n-2} R_1-C= re R_1 and R_2 = H memical properties ith electrophiles i.	rss rss n cloud gets big rss forces, hence bo rss increases by a C C - H bonds to t rst off during combu- rst of rst	ger and mo biling point in CH ₂ group preak and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Normal, Indent: Before: 0", Hanging: 0.5", bullets or numbering Field Code Changed
From et <u>hence is</u> <u>More er</u> <u>From et</u> <u>This res</u> <u>bonds ts</u> */*/*/ es are organi homologous	id-id) forces to thane to hep d-id forces be hergy needed hane to hept sults in more o form, hence c molocules series with f whe ibit similar ch	between molecule otane, the electro- ecome stronger. I to break the id-ic ane, the molecule C – C bonds and e more heat given which contain c formula of C_nH_{2n-2} R_1-C= re R_1 and R_2 = H memical properties ith electrophiles i.	rss rss n cloud gets big rss forces, hence bo rss increases by a C C - H bonds to t rst off during combu- rst of rst	ger and mo biling point in CH ₂ group preak and m istion.	re easily polarise creases ore C=O and O –	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Normal, Indent: Before: 0", Hanging: 0.5", bullets or numbering Field Code Changed
From et <u>hence in</u> <u>More er</u> <u>From et</u> <u>This res</u> <u>bonds tr</u> <u>*/*///</u> es are organi homologous <u>Alkynes exh</u> 	id-id) forces to thane to hep d-id forces be hergy needed hane to hept sults in more o form, hence o form, hence c moloculos series with f whe ibit similar ch reactions wi	between molecule otane, the electro- ecome stronger. I to break the id-ic ane, the molecule C – C bonds and e more heat given which contain c formula of C_nH_{2n-2} R_1-C= re R_1 and R_2 = H memical properties ith electrophiles i.	$\frac{1}{1} \frac{1}{1} \frac{1}$	ger and mo piling point in CH ₂ group preak and m istion. 	re easily polarise icreases ore C=O and O – nd are part of t	:d.	bullets or numbering Formatted: Font: 11 pt Formatted: Normal, Indent: Before: 0", Hanging: 0.5", bullets or numbering Field Code Changed Formatted: Font: 11 pt, Complex Script Font: 11 pt

	e.g. oxidation by hot concentrated KMnO₄ to form mixture of carboxylic acids		
_			
	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	ľ	Field Code Changed
	However, unlike alkenes, terminal alkynes are able to react with strong bases like sodium amide.		
	$\frac{RC = CH + NaNH_2}{RC = C^{-}Na^{+} + NH_3}$		Field Code Changed
a)	Ethyne, C_2H_2 , is heated with excess sodium bromide and concentrated sulfuric acid to produce a dihalide, $C_2H_4Br_2$. The overall reaction may be considered to take place in two stages, the first between inorganic reagents only and the second involving the organic reagent.		
	(i) Write an equation for the first stage. [1]		
	(ii) Suggest a structure for the dihalide formed. [1]		
	(iii) When the concentrated sulfuric acid is added to the reaction mixture, cooling is necessary to prevent the formation of inorganic by products.		
	Write an equation to explain the formation of these inorganic by-products. [1]		
b)	Compound A , is an enyne chloride (i.e. compounds that contains chloro, alkyne and alkene functional groups).		
			Field Code Changed
		/	
		- /	
	Ci	/	
		í –	
	Compound A		
	One mole of compound A reacts with two moles of Br ₂ to produce a mixture of 4 storeoisomers. Draw structures of the storeoisomers formed. [3]		
(c			Formatted: Font: 11 pt, Complex Script Font: 11 pt
A 9	treated with sodium amide, NaNH ₂ followed by heating under reflux to form compound C , C ₈ H ₁₀ . Compound C reacts with hot concentrated KMnO ₄ to produce butane 1,4 dioic acid	< `	Formatted: Normal, Indent: Before: 0", Hanging: 0.5", N bullets or numbering, Tab stops: 0.3", Left + Not at 1.08" 6.6"
	only.Suggest why combustion tends to be incomplete as the alkane increases in molecular mass.		Formatted: Font: 11 pt, Complex Script Font: 11 pt
	As the alkane increases in molecular mass, the alkane becomes liquid and liquid phase		Formatted: Font: 11 pt, Complex Script Font: 11 pt, Highlight
	reactions are slower than gas phase reactions. Hence combustion tends to be incomplete.	5	Formatted: Font: 11 pt
Velo	propane is a colourless gas with a "petroleum-like" odour. Unlike its straight-chain		Formatted: Indent: Before: 0.5"
	erpart, it is considered to be highly strained and unstable. The instability of cyclic alkanes can		Formatted: Font: 11 pt
	asured by calculating its "ring strain energy" using the formula below:		Formatted: Font: 11 pt
	<u>Ring strain energy = </u>		Formatted: Font: 11 pt
	<u></u>	<	Formatted: Space After: 0 pt, Line spacing: single



	28		
<u>(f)</u>	Due to the presence of ring strain, cyclopropane undergoes an addition reaction with +>>		Formatted: Font: 11 pt, Complex Script Font: 11 pt
	bromine in the absence of ultraviolet radiation.		Formatted: Indent: Hanging: 0.5", Numbered + Level: 1 + Numbering Style: a, b, c, + Start at: 6 + Alignment: Left + Aligned at: 0.25" + Indent at: 0.5", Tab stops: Not at 1.08"
	(i) Suggest the skeletal structure of the molecule formed after reaction with Br₂. [1] ← Br		Formatted: Indent: Hanging: 0.55", Numbered + Level: 1 + Numbering Style: i, ii, iii, + Start at: 1 + Alignment: Left + Aligned at: 0.54" + Indent at: 1.04", Tab stops: Not at 6.6"
	· · · · · · · · · · · · · · · · ·	{	Formatted: Indent: Before: 1.04", No bullets or numbering, Tab stops: Not at 6.6"
	(ii) Hence, using VSEPR theory, explain why the presence of ring strain causes		Formatted: Font: 11 pt, Complex Script Font: 11 pt
	cyclopropane to undergo addition reactions. [1]	{	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	Cyclopropane has a C – C – C angle of 60° which is smaller than the optimal angle -		Formatted: Font: 11 pt
	of 109.5° in sp ³ carbons. As a result, bond pairs are much closer to each other and		Formatted: Indent: Before: 0.98"
	experience greater repulsion, hence weakening the C – C bonds, which breaks	\sum	Formatted: Font: 11 pt
	easily during addition.	$\left(\right)$	Formatted: Font: 11 pt
		\sim	Formatted: Font: 11 pt
A		Ì	Formatted: Font: 11 pt
<u>(g)</u>	Cyclopropane rings can be formed using a technique called "cyclopropanation".]	Formatted: English (Singapore)
	One such cyclopropanation technique involves the 2 mechanistic steps stated below;	11	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	Step 1: Dissociation of diazomethane, CH ₂ N ₂ to form methylene, CH ₂ , and N ₂ . Formation of		Formatted: Indent: Hanging: 0.5", Numbered + Level: 1 + Numbering Style: a, b, c, + Start at: 6 + Alignment: Left + Aligned at: 0.25" + Indent at: 0.5", Tab stops: Not at 1.08"
metnyk	ene, CH ₂ , and N ₂ from diazomethane, CH ₂ N ₂ .	$\langle \cdot \rangle$	Formatted: Indent: First line: 0.2"
	Step 2: Addition of methylene, CH ₂ , to trans-but-2-ene to form the cyclic ring. The reaction	\sim	Formatted: Font: 11 pt, Complex Script Font: 11 pt
4	leaves the stereochemistry of the molecule unchanged.	. 'Y	Formatted: Indent: Before: 0", First line: 0.5"
	A	\sum	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	(i) It is observed that the diazomethane molecule is trigonal planar in shape. By	$\langle \rangle$	Formatted: Indent: Before: 0.49", Hanging: 0.57"
	considering the shape, dDraw a dot-and-cross diagram of diazomethane, CH ₂ N ₂ , $$		Formatted: Font: 11 pt, Complex Script Font: 11 pt
	clearly showing the type of bonds formed within the molecule.	N N N N N N N N N	Formatted: Indent: Before: 0.49", Hanging: 0.49", Numbered + Level: 1 + Numbering Style: a, b, c, + Start at: 9 + Alignment: Left + Aligned at: 1" + Indent at: 1.25", Tab stops: Not at 6.6"
		111	Formatted: Font: 11 pt, Complex Script Font: 11 pt
		111	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	H	11	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	•x xx	- Y	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	H X C X IN IX NX	ĺ	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	(ii) _Draw the structure of the cyclic molecule formed in sStep 2, showing the	(Formatted: Indent: Before: 1.25", No bullets or numbering, Tab stops: Not at 6.6"
	stereochemical arrangement clearly. State and explain if the molecule can rotate		Formatted: Font: 11 pt, Complex Script Font: 11 pt
	plane-polarised light. [2]	1	Formatted: Font: 11 pt, Complex Script Font: 11 pt
		$\langle \rangle$	Formatted: Font: 11 pt, Complex Script Font: 11 pt
		1	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	H ₃ C H	Ì	Formatted: Indent: Before: 0"
	H ČH ₃		





15 mi	a current of 1.0 A was passed through aqueous potassium maleate (KO ₂ CCH=CHCO ₂ K) for nutes, it was found that 110 cm ³ H ₂ , measured at r.t.p, was collected at the cathode. The ng reaction took place.	
	$2H_2O + 2e^- \rightarrow H_2 + 2OH^-$	
(d)	State the relationship between the Faraday constant, F and the Avogadro's constant, L. [1]	
(e)	Using the data above and the Data Booklot, calculate a value for Avogadro's constant. [3]	
(f)	Ethyne and CO ₂ -gas were produced at the anode. In order to determine the stoichiometry of the anode reaction, the volume of the gases collected at the anode was measured. The anode gas was first passed through aqueous NaOH before being collected in a gas syringe. The following data was collected:	
	mass of NaOH before experiment = 10.501 g mass of NaOH after experiment = 10.904 g initial reading on syringe = 10.0 cm ³ final reading on syringe = 120.0 cm ³	
	(i) State the oxidation state of carbon in ethyne. [1]	
	(ii) With the help of an equation, explain the purpose of passing the anode gas through NaOH. [1]	
	(iii) Calculate the volume of CO ₂ produced, assuming r.t.p conditions. [1]	
	(iv) Hence, suggest an ionic equation for the reaction that occurred at the anode. [1]	
(g)	When aqueous potassium maleate was acidified, maleic acid, HO ₂ CCH=CHCO ₂ H ($pK_{g_1} = 1.90$ and $pK_{g_2} = 6.07$) was liberated. Fumaric acid ($pK_{g_1} = 3.03$ and $pK_{g_2} = 4.44$) is a stereoisomer of maleic acid. With a suitable illustration, suggest a reason why maleic acid has a lower pK_{g_1} -but higher pK_{g_2} than fumaric acid.	Formatted: Font: Not Italic, Complex Script Font: Not Italic, Subscript
	[2] [Total: 20]	
▲		Formatted: Highlight

5(a) Depamine is an organic compound of the catecholamine and phenethylamine families that plays several important roles in the brain and body. Its name is derived from its chemical structure: it is an amine synthesised by removing a carboxyl group from a molecule of its precursor compound L DOPA.

its precursor compound, L-DOPA. The halogens and their compounds, show many similarities and trends in their properties. Some data are given for the elements fluorine, chlorine and iodine.

<u>Element</u>	Bond Energy / kJ mol ⁻¹	Standard enthalpy change of atomisation / kJ mol ⁻¹
Fluorine	<u>158</u>	<u>79</u>
<u>Chlorine</u>	<u>242</u>	<u>121</u>
Bromine	<u>193</u>	<u>112</u>
lodine	<u>151</u>	<u>107</u>

(i) For fluorine and chlorine, their enthalpy changes of atomisation are half the value of their respective bond energies. For bromine and iodine, their enthalpy changes of atomisation are much more than half the value of their respective bond energies.

Explain in detail for this difference.

Fluorine and chlorine are gases but bromine is a liquid and iodine is a solid at roomtemperature. The enthalpy change of atomisation includes the energy required to change $Br_2(I) \rightarrow Br_2(g)$ and $I_2(g) \rightarrow I_2(g)$ [to vapourise bromine and iodine to the gaseous state].

(ii) The standard enthalpy change of formation of iodine monochloride, I–Cl, is −24.0 kJ mol⁻¹.

Use this information and the data from the table above to calculate the I–Cl bond energy. [1]

 $\underline{\Delta H_r} = \underbrace{\sum \text{Bonds broken } -\sum \text{Bonds formed}}_{=24.0 = \frac{1}{2} \text{BE}(\text{CI-CI}) + \frac{1}{2} \text{BE}(\text{I-I}) - \text{BE}(\text{I-CI})}_{\text{BE}(\text{I-CI}) = +220.5}$

= +221 kJmol⁻¹

Below is a synthetic route involving L-DOPA and dopamine:

Formatted: English (Singapore)

 $\mbox{Formatted:}$ Indent: Before: 0", Hanging: 0.49", No bullets or numbering

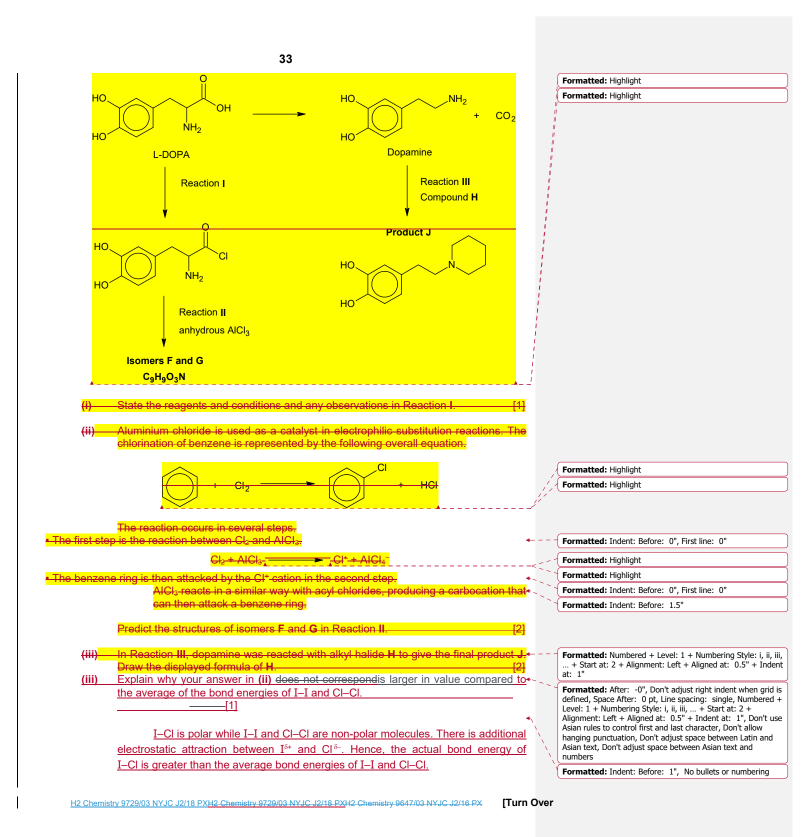
-		Formatted: Space Before: 3 pt, After: 3 pt
		Formatted Table
+		Formatted: Space Before: 3 pt, After: 3 pt
-		Formatted: Space Before: 3 pt, After: 3 pt
-		Formatted: Space Before: 3 pt, After: 3 pt
-		Formatted: Space Before: 3 pt, After: 3 pt
<u>the</u> +		Formatted: Font: Bold, Not Highlight
<u>llpy</u> ond		Formatted: Indent: Before: 0.5", Hanging: 0.5", Space After: 10 pt, Line spacing: Multiple 1.15 li, No bullets or numbering
	. · ·	Formatted: Not Highlight
[1]	1	Formatted: Highlight
1-1		
<u>om</u> ≁-		Formatted: Indent: Before: 1"
l to		
the		
is≉.		Formatted: Indent: Before: 0.5", First line: 0.5", Space After: 0 pt, Line spacing: single
<u>- 10</u> * *		Formatted: Numbered + Level: 1 + Numbering Style: i, ii, iii, + Start at: 2 + Alignment: Left + Aligned at: 0.5" + Indent at: 1"
ond [1]		Formatted: Indent: Before: 1", Space After: 0 pt, Line spacing: single
		Formatted: Indent: First line: 0.25"

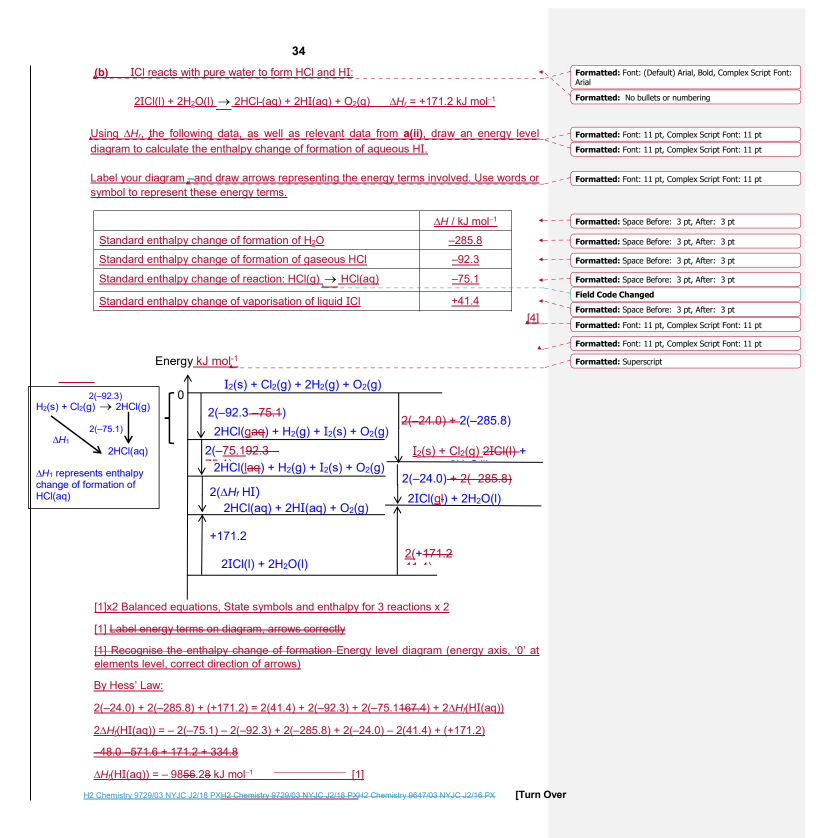
Formatted: Indent: Before: 1.25"

Formatted: Highlight

[Turn Over

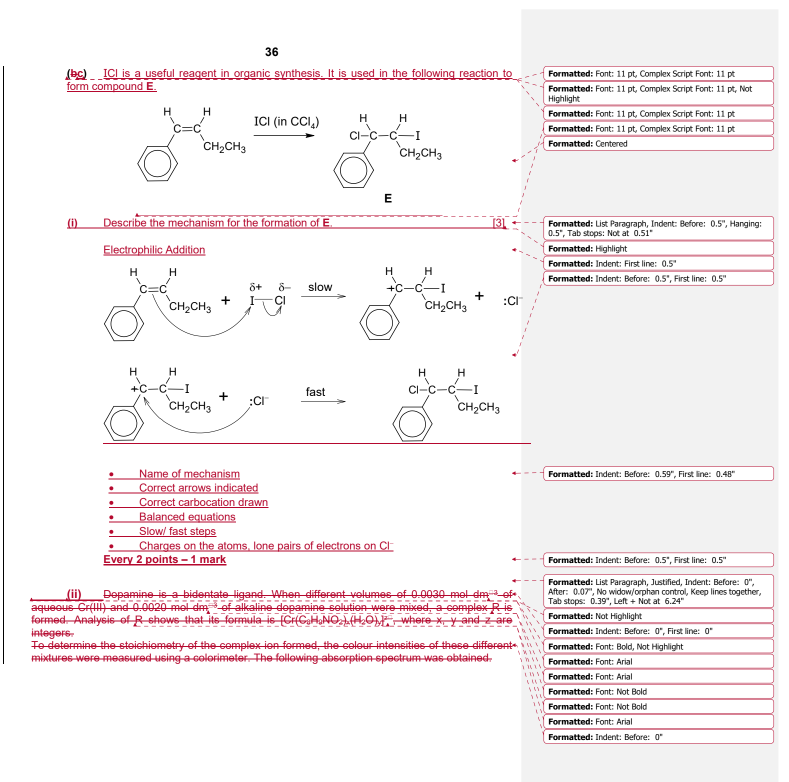
H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX



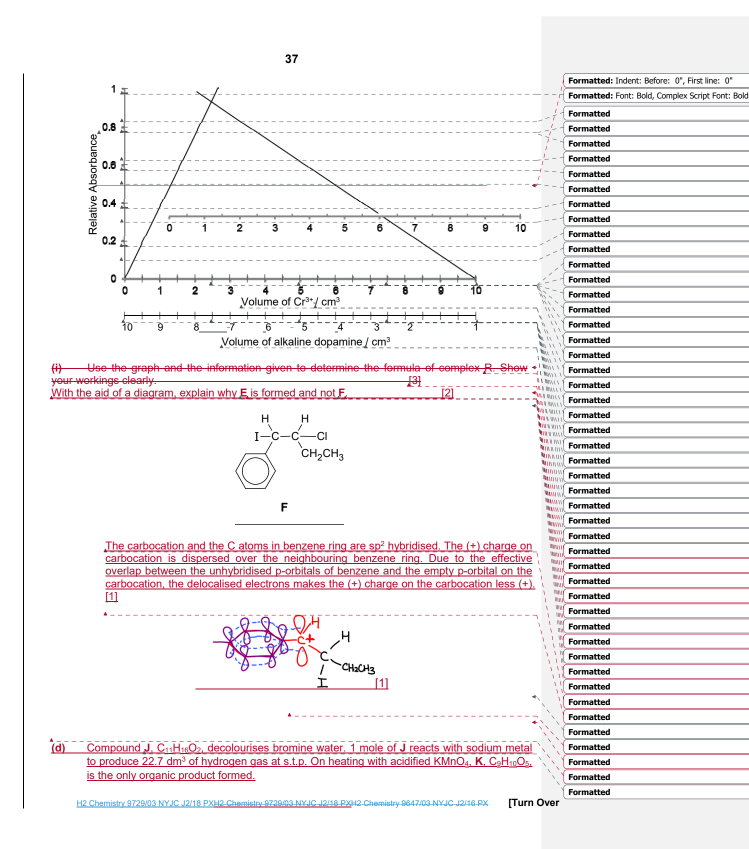




H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX [Turn Over



[Turn Over



[... [77]

[... [75]

... [68]

... [76]

... [73]

... [74]

[... [71]

[... [72]

... [69]

[... [70]

... [79]

[... [80]

... [78]

... [83]

... [84]

... [81]

... [82]

... [85]

... [86]

... [91]

[... [92]

... [89]

... [90]

... [95]

... [96]

... [93]

[... [87]

... [88]

... [51]

... [52]

... [53]

... [54]

... [55]

... [56]

[... [57]

[... [58]

... [59]

... [60]

... [61]

... [62]

... [63]

[... [64]

[... [65]

[... [66]

... [67]

.. [94]

	odium carbonate and 2,4-DNPH. When K reacts with alkaline aqueou	<u>is</u>	Formatted: Font: 11 pt, Complex Script Font: 11 pt
iodine, L is forme	d upon acidification.		Formatted: Font: 11 pt, Complex Script Font: 11 pt
A	0 	\`\`\	Formatted: Font: 11 pt, Complex Script Font: 11 pt, Not Highlight
			Formatted: Font: 11 pt, Complex Script Font: 11 pt
	HO' Y Y	` .	Formatted: Font: 11 pt, Complex Script Font: 11 pt
	HO, 🙏 🗼		Formatted: Font: 11 pt, Complex Script Font: 11 pt
	$\gamma \sim 0$		Formatted: Font: 11 pt, Complex Script Font: 11 pt
	⁰ L		
Suggest structure	es for J and K and explain the reactions described.	<u>51 _</u>	Formatted: Font: (Default) Arial, 11 pt, Complex Script Fond Arial, 11 pt
	ОН		Formatted: Font: 11 pt, Complex Script Font: 11 pt
		1	Formatted: Font: 11 pt, Complex Script Font: 11 pt
		×	Formatted: List Paragraph, Justified, Indent: Before: 0", After: 0.07", No widow/orphan control, Keep lines together, Tab stops: 0.39", Left + Not at 6.24"
	CH ₃		
	з к		
ervations	Deductions		
11 <u>H16</u> O2	J could be alkene or phenol.		
olourises bromine	An alkene undergoes electrophilic addition reaction with Br ₂ (aq) to		
er ′ dm³ of hydrogen	form halogenoalkane.		
rmed at s.t.p when	$\underline{n(H_2) = \frac{22.7}{22.7} = 1 \text{ mol of } H_2 \text{ is formed.}}$		
reacted with			
um metal	$\underline{\text{ROH} + \text{Na}} \rightarrow \underline{\text{RO}} \underline{\text{Na}} + \frac{1}{2}\underline{\text{H}}_2$		
	Since 1 FG produces $\frac{1}{2}$ mole of H ₂ , there must be 2 –OH groups		
	present. –COOH group is absent as it will only produce only ½ mole of H ₂ and 2 –COOH groups or 1 –OH & 1 –COOH groups cannot be		
	present as it will not correspond to the molecular formula of J .		
heating with	Secondary alcohol and the alkene in J undergoes oxidation with		
lified KMnO₄, K ,	KMnO ₄ to form ketones and carboxylic acid.		
$1_{10}O_5$, is the only	There is a decrease in 2C atoms – which suggest that ethane-1,2-		
anic product formed	dioc acid was oxidised to form CO ₂ .		
	From the given structure of L, it can be seen that K is unlikely a		
	phonol and should not contain honzone as there will not be ovidation		

H2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX

that leads to a reduction of 2 C atoms.

ketones.

present.

K reacts with sodium carbonate and 2,4

K reacts with alkaline

aqueous iodine to form

3 marks for explanation 5 points – 3 marks 3 to 4 points – 2 marks

2 points – 1 mark

1 mark for each correct structure

DNPH

phenol and should not contain benzene as there will not be oxidation

K undergoes condensation with 2,4 DNPH → -confirms presence of

K undergoes mild oxidation with alkaline $I_2(aq) \rightarrow -COCH_3$ present.

K undergoes acid-base reaction with alkali to form salt → -COOH

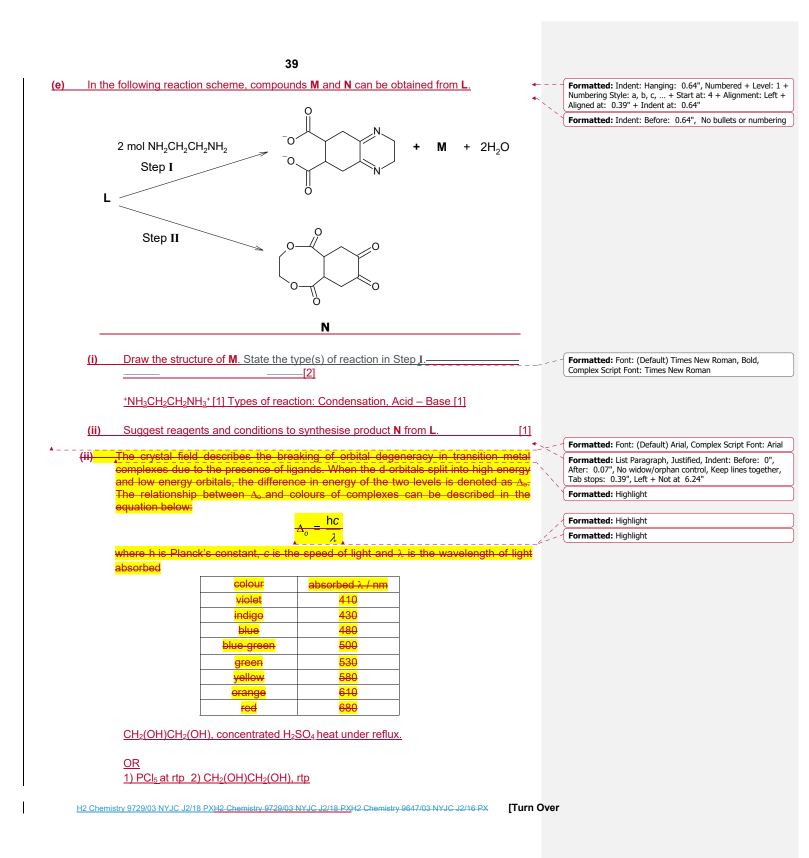
K undergoes acid-base with Na₂CO₃ \rightarrow –COCH₃ present.

38

Formatted: List Paragraph, Justified, Indent: Before: 0", After: 0.07", No widow/orphan control, Keep lines together, Tab stops: 0.39", Left + Not at 6.24"

Formatted: Space After: 0 pt

Tab stops: 0



		40		
		Given that <u>A</u> , for complex R is 4.125 x 10 ⁻²² kJ and using relevant data from the Data Booklet, calculate the wavelength of light. Deduce the colour of complex R		Formatted: Highlight
(6)	equilib			
	(i)	Draw a fully labelled experimental set-up for a voltaic cell made up of a Cr_2Oq^{2-1} Cr ³⁺ half-cell and a F_2/I^{2-} half-cell under standard conditions. Indicate clearly the anode and cathode and show the flow of electrons.		
	(ii)	By using appropriate values from the <i>Data Booklet</i> , predict what, if anything, will happen when a small amount of acidified vanadium(II) chloride is added to the I ₂ /I ⁻ half cell.		
(d)	- Explai	n the following statements.		
	(i)	BrF ₃ is a covalent compound which exhibits electrical conductivity in liquid state at room temperature. With the aid of an equation, suggest an explanation for its electrical conductivity. [2]		
	(ii)	SiCl ₄ reacts violently in water but CCl ₄ has no reaction with water. [1]		
	(iii) <mark> </mark>	Compounds NeF ₂ and NeF ₄ do not exist but XeF ₂ and XeF ₄ exist. [1]		
		- [Total: 20] -		Formatted: After: -0.01"
		4- 4-		Formatted: Right, After: -0.01" Formatted: After: -0.01"
		+		Formatted: Indent: Before: 0", First line: 0", After: -0.0
		•		Formatted: After: -0.01"
		[End of Paper]		Formatted: Font: (Default) Arial, Complex Script Font: Aria
		ry 9729/03 NYJC J2/18 PXH2 Chemistry 9729/03 NYJC J2/18 PXH2 Chemistry 9647/03 NYJC J2/16 PX	NOT	

Page 7: [1] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt, Complex Script Font: 11	pt	
Page 7: [2] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:23:00 PM
Indent: Before: 0.5", First line: 0.5"		
Page 7: [3] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: (Default) Arial, 11 pt, Complex	Script Font: Arial, 11 pt	
Page 7: [4] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt, Complex Script Font: 11	pt	
Page 7: [5] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt, Complex Script Font: 11	pt	
Page 7: [6] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt, Complex Script Font: 11	pt	
Page 7: [7] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt, Complex Script Font: 11	pt	
Page 7: [8] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt, Complex Script Font: 11	pt	
Page 7: [9] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Complex Script Font: 11 pt		
Page 7: [10] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 12 pt		
Page 7: [11] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [12] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Indent: Before: 0.5", First line: 0.5"		
Page 7: [13] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [14] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [15] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [16] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [17] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [18] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [19] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		

Font: 11 pt

Page 7: [20] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [21] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Indent: Before: 1"		
Page 7: [22] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [23] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [24] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Indent: Before: 0.5", First line	: 0.5"	
Page 7: [25] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Indent: Before: 1"		
Page 7: [26] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [27] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [28] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [29] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
Font: 11 pt		
Page 7: [30] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:24:00 PM
English (United States)		
Page 7: [31] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:45:00 PM
After: -0.03"		
Page 7: [32] Formatted	Eunice Sim Kwei Yen	05-Aug-18 4:15:00 PM
Font: Bold		
Page 7: [33] Formatted	Eunice Sim Kwei Yen	05-Aug-18 4:15:00 PM
Font: Bold		
Page 7: [34] Formatted	Eunice Sim Kwei Yen	05-Aug-18 4:15:00 PM
Font: 11 pt, Bold, Complex Sc	ript Font: 11 pt	
Page 7: [35] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:45:00 PM
After: -0.03"		
Page 7: [36] Formatted	Eunice Sim Kwei Yen	05-Aug-18 4:15:00 PM
Font: Bold		
Page 7: [37] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Fo	ont: 11 pt	
Page 7: [38] Formatted	Eunice Sim Kwei Yen	05-Aug-18 4:03:00 PM
Indant: First line: O" After: (0.02	

Indent: First line: 0", After: -0.03"

Page 7: [39] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Font: 1	1 pt	
Page 7: [40] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Bold, Complex Script I (United States), Do not check spelli		rabic (Saudi Arabia), English
Page 7: [41] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Font: 1	1 pt	
Page 7: [42] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Font: 1	1 pt	
Page 7: [43] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Font: 1	1 pt	
Page 7: [44] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Font: 1	1 pt	
Page 7: [45] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Font: 1	1 pt	
Page 7: [46] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Font: 1	1 pt	
Page 7: [47] Formatted	Eunice Sim Kwei Yen	15-Aug-18 3:28:00 PM
Centered		
Page 7: [48] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt, Complex Script Font: 1	1 pt	
Page 7: [49] Formatted	Eunice Sim Kwei Yen	15-Aug-18 3:37:00 PM
Numbered + Level: 1 + Numbering 0.5" + Indent at: 1"	Style: i, ii, iii, + Start at: 1 +	Alignment: Left + Aligned at:
Page 7: [50] Formatted	Eunice Sim Kwei Yen	05-Aug-18 3:51:00 PM
Font: 11 pt		
Page 37: [51] Formatted	Eunice Sim Kwei Yen	06-Aug-18 12:02:00 AM
No bullets or numbering, Tab stops	s: Not at 0.39"	
Page 37: [52] Formatted	Eunice Sim Kwei Yen	06-Aug-18 12:02:00 AM
Font: Not Bold, Complex Script Fo	nt: Arial	
Page 37: [53] Formatted	Eunice Sim Kwei Yen	06-Aug-18 12:02:00 AM
Complex Script Font: Arial		
Page 37: [54] Formatted	Eunice Sim Kwei Yen	06-Aug-18 12:02:00 AM
Font: Arial		
Page 37: [55] Formatted	Eunice Sim Kwei Yen	06-Aug-18 12:02:00 AM
List Paragraph, Justified, Indent: Be	efore: 0", After: 0.07", No wid	ow/orphan control, Keep lines
together, Tab stops: Not at 6.24"		
together, Tab stops: Not at 6.24" Page 37: [56] Formatted	Theresia Line Ishak	13-Aug-18 8:08:00 AM

Font: Arial

Page 37: [57] Formatted	Eunice Sim Kwei Yen	06-Aug-18 12:02:00 AM
Font: Arial		
Page 37: [58] Formatted	Theresia Line Ishak	13-Aug-18 8:08:00 AM
Font: Arial		
Page 37: [59] Formatted	Eunice Sim Kwei Yen	06-Aug-18 12:02:00 AM
Font: Arial	Lunce Sin Kwei Ten	00-Aug-10 12.02.00 AM
	Thoracia Line Tehek	12 Aug 19 7.51.00 AM
Page 37: [60] Formatted List Paragraph, Centered, Indent: E	Theresia Line Ishak	13-Aug-18 7:51:00 AM
lines together, Tab stops: Not at 6.		ow/orphan control, Keep
Page 37: [61] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:59:00 PM
Font: 11 pt		
Page 37: [62] Formatted	Eunice Sim Kwei Yen	14-Sep-18 1:59:00 PM
Font: 11 pt		
Page 37: [63] Formatted	Theresia Line Ishak	13-Aug-18 7:51:00 AM
List Paragraph, Centered, Indent: E lines together, Tab stops: Not at 6.		ow/orphan control, Keep
		06-Aug-18 12:02:00 AM
Page 37: [64] Formatted	Eunice Sim Kwei Yen	
Page 37: [64] Formatted Font: Arial		
	Eunice Sim Kwei Yen	06-Aug-18 12:00:00 AM
Font: Arial	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide	06-Aug-18 12:00:00 AM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide	06-Aug-18 12:00:00 AM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24"	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24"	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Scr	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Scr Page 37: [68] Formatted	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Script Page 37: [68] Formatted Font: 11 pt, Complex Script Font:	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Script Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [69] Formatted Font: Bold, Complex Script Font: 1	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM 24-Aug-16 7:04:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Scri Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [69] Formatted Font: Bold, Complex Script Font: I Page 37: [70] Formatted	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Script Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [69] Formatted Font: Bold, Complex Script Font: I Font: Bold, Complex Script Font: I	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold Theresia Line Ishak Bold, English (Singapore)	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM 24-Aug-16 7:04:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Scri Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [69] Formatted Font: Bold, Complex Script Font: I Page 37: [70] Formatted	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold Theresia Line Ishak Bold, English (Singapore) Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM 24-Aug-16 7:04:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Scri Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [69] Formatted Font: Bold, Complex Script Font: I Page 37: [70] Formatted Font: Bold, Complex Script Font: I Page 37: [71] Formatted Font: Bold, Complex Script Font: I	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold Theresia Line Ishak Bold, English (Singapore) Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM 24-Aug-16 7:04:00 PM 24-Aug-16 7:04:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Script Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [69] Formatted Font: Bold, Complex Script Font: I Page 37: [71] Formatted Font: Bold, Complex Script Font: I Page 37: [72] Formatted	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold Theresia Line Ishak Bold, English (Singapore) Theresia Line Ishak Bold	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM 24-Aug-16 7:04:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Script Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [70] Formatted Font: Bold, Complex Script Font: 1 Page 37: [71] Formatted Font: Bold, Complex Script Font: 1 Page 37: [72] Formatted Font: Bold, Complex Script Font: 1	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold Theresia Line Ishak Bold, English (Singapore) Theresia Line Ishak Bold Theresia Line Ishak Bold	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM 24-Aug-16 7:04:00 PM 24-Aug-16 7:04:00 PM 24-Aug-16 7:04:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Script Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [69] Formatted Font: Bold, Complex Script Font: I Page 37: [71] Formatted Font: Bold, Complex Script Font: I Page 37: [72] Formatted Font: Bold, Complex Script Font: I Page 37: [73] Formatted	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold Theresia Line Ishak Bold, English (Singapore) Theresia Line Ishak Bold English (Singapore) Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM 24-Aug-16 7:04:00 PM 24-Aug-16 7:04:00 PM
Font: Arial Page 37: [65] Formatted List Paragraph, Justified, Indent: B together, Tab stops: 0.39", Left + 1 Page 37: [66] Formatted Font: Arial Page 37: [67] Formatted Font: (Default) Arial, Complex Scri Page 37: [68] Formatted Font: 11 pt, Complex Script Font: 1 Page 37: [70] Formatted Font: Bold, Complex Script Font: 1 Page 37: [71] Formatted Font: Bold, Complex Script Font: 1 Page 37: [72] Formatted Font: Bold, Complex Script Font: 1	Eunice Sim Kwei Yen Before: 0", After: 0.07", No wide Not at 6.24" Eunice Sim Kwei Yen Theresia Line Ishak ript Font: Arial Theresia Line Ishak 11 pt Theresia Line Ishak Bold Theresia Line Ishak Bold, English (Singapore) Theresia Line Ishak Bold English (Singapore) Theresia Line Ishak	06-Aug-18 12:00:00 AM ow/orphan control, Keep lines 06-Aug-18 12:02:00 AM 13-Aug-18 8:22:00 AM 24-Aug-16 7:20:00 PM 24-Aug-16 7:04:00 PM 24-Aug-16 7:04:00 PM 24-Aug-16 7:04:00 PM

Font: Bold, Complex Script Font: Bold, English (Singapore)

Page 37: [75] Formatted	Theresia Line Ishak	24-Aug-16 7:04:00 PM
Font: Bold, Complex Script Font: Bold		
Page 37: [76] Formatted	Theresia Line Ishak	24-Aug-16 7:04:00 PM
Font: Bold, Complex Script Font: Bold	, English (Singapore)	
Page 37: [77] Formatted	Theresia Line Ishak	24-Aug-16 7:04:00 PM
Font: Bold, Complex Script Font: Bold	, English (Singapore)	
Page 37: [78] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold	, English (Singapore)	
Page 37: [79] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold		
Page 37: [80] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold	, English (Singapore)	
Page 37: [81] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold		
Page 37: [82] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold	, English (Singapore)	
Page 37: [83] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold		
Page 37: [84] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold	, English (Singapore)	
Page 37: [85] Formatted	Theresia Line Ishak	24-Aug-16 7:20:00 PM
Font: 11 pt, Complex Script Font: 11 p	t	
Page 37: [86] Formatted	Theresia Line Ishak	24-Aug-16 7:20:00 PM
Font: 11 pt, Complex Script Font: 11 p	t	
Page 37: [87] Formatted	Theresia Line Ishak	24-Aug-16 7:20:00 PM
Font: 11 pt, Complex Script Font: 11 p	t	
Page 37: [88] Formatted	Theresia Line Ishak	24-Aug-16 7:20:00 PM
Font: 11 pt, Complex Script Font: 11 p	t	
Page 37: [89] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold		
Page 37: [90] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold	, English (Singapore)	
Page 37: [91] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold		
Page 37: [92] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold, Complex Script Font: Bold	, English (Singapore)	
Page 37: [93] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM
Font: Bold Complex Script Font: Bold		

Font: Bold, Complex Script Font: Bold

Page 37: [94] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM		
Font: Bold, Complex Script Font: Bold, English (Singapore)				
Page 37: [95] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM		
Font: Bold, Complex Script Font: Bold				
Page 37: [96] Formatted	Theresia Line Ishak	24-Aug-16 6:52:00 PM		
Font: Bold Complex Script Fo	ont: Bold English (Singapore)			

Font: Bold, Complex Script Font: Bold, English (Singapore)

	NANYANG JUNIOR COLLEGE JC2 PRACTICAL PRELIMINARY EXAMINATION Higher 2		
CANDIDATE NAME			
CLASS 1	7 REG TUTOR		
CHEMIST Paper 4 Practic	al	2	9729/04 28 August 2018 hour 30 minutes
Candidates ans	wer on the Question Paper		
Additional Mate	rials: As listed in the Confidential Instructions		
Write your nam Give details of t Write in dark blu You may use an Do not use stap	NSTRUCTIONS FIRST e and class on all the work you hand in. he practical shift and laboratory, where appropriate, in the boxes pro- ue or black pen. h HB pencil for any diagrams or graphs les, paper clips, glue or correction fluid. stions in the spaces provided on the Question Paper.	vided.	
The use of an a	pproved scientific calculator is expected, where appropriate.		Shift
appropriate unit			
Qualitative Analysis Notes are printed on pages 22 and 23.			aboratory
	e examination, fasten all your work securely together. marks is given in brackets [] at the end of each question or part		
		For Exam	iner's Use
		1	/ 28
		2	/ 11

/16

/55

3

Total

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

1 To determine the concentrations of aqueous sodium hydroxide and aqueous ethanedioic acid and hence the percentage by mass of sodium ethanedioate in a mixture of sodium ethanedioate and ethanedioic acid.

You are provided with the following reagents.

- **FB 1** is a mixture of approximately 0.01 mol dm⁻³ aqueous sodium ethanedioate, C₂O₄Na₂, and approximately 0.1 mol dm⁻³ ethanedioic acid, C₂O₄H₂
- **FB 2** is aqueous sodium hydroxide, NaOH
- **FB 3** is 0.0755 mol dm⁻³ aqueous sodium carbonate
- thymolphthalein indicator

Both sodium hydroxide and sodium carbonate are bases which will neutralise ethanedioic acid as shown in **reactions 1** and **2**.

$$2NaOH + C_2O_4H_2 \rightarrow C_2O_4Na_2 + 2H_2O$$
 reaction 1

$$Na_2CO_3 + C_2O_4H_2 \rightarrow C_2O_4Na_2 + CO_2 + H_2O$$
 reaction 2

In this experiment, you will add different volumes of **FB 3** to identical samples of the mixture of aqueous sodium ethanedioate, $C_2O_4Na_2$, and ethanedioic acid, $C_2O_4H_2$, **FB 1**. You will then complete the neutralisation of $C_2O_4H_2$ in each mixture by titrating with dilute sodium hydroxide, **FB 2**.

Each titration is to be performed **once only**, so great care should be taken that you do not exceed the end-points.

Graphical analysis of your results will enable you to determine the concentrations of the ethanedioic acid in the mixture of aqueous $C_2O_4Na_2$, and $C_2O_4H_2$, **FB 1** and of the sodium hydroxide, **FB 2**.

Read through the whole method of conducting the experiment before starting any practical work.

The experiment

(a) Titrations

Experiment 1

- 1. Fill a burette with **FB 3**.
- 2. Pipette 25.0 cm³ of **FB 1** into a 250 cm³ conical flask.
- 3. Run 5.00 cm³ of **FB 3** into the flask.
- 4. Add 10 15 drops of thymolphthalein indicator to the flask.
- 5. Fill a second burette with **FB 2**.
- 6. Titrate the mixture in the conical flask against **FB 2** until the first permanent pale blue colour remains in the solution.
- 7. Rinse the conical flask thoroughly before it is used for the next experiment.

The end-point should be found after the addition of approximately 14 cm³ of **FB 2**.

One titration, performed accurately, will be sufficient.

Experiment 5

1. Repeat **Experiment 1** but run 30.00 cm³ of **FB 3** into the flask containing 25.0 cm³ of **FB 1**.

Experiments 2 – 4

Select three other suitable volumes of **FB 3** for use in **Experiments 2 – 4**. Your selected volumes must be between the volumes used in **Experiments 1 and 5**.

Do not use volumes of FB 3 outside the range of 5 – 30 cm³.

Repeat the procedures used in **Experiment 1** but, in each case, add 25.0 cm³ of **FB 1** and your chosen volume of **FB 3** into the conical flask. You should perform your titrations in order of **increasing** volume of **FB 3**.

Rinse this conical flask thoroughly between each titration.

Prepare a table in the space below and use it to record the titration results for each volume of **FB 3** added. You should record your titration results in order of **increasing** volume of **FB 3** added.

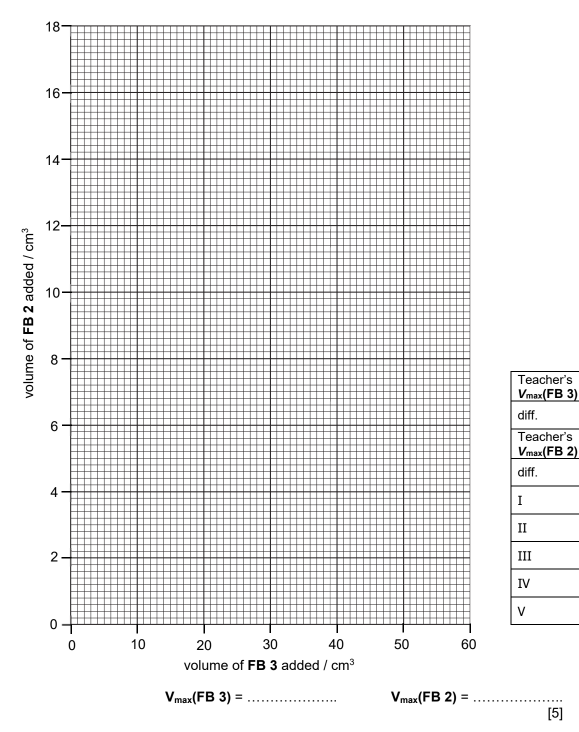
Ι	
Π	

[2]

(b) (i) Plot, on the grid below, your values for the FB 2 titre (y-axis) against the volume of **FB 3** added (*x-axis*).

Draw the line of best fit, taking into account all of your plotted points. Hence obtain values for

- the volume of FB 3 required, $V_{\text{max}}(\text{FB 3}),$ to react completely with • 25.0 cm³ of **FB 1** if no **FB 2** is added;
- the volume of FB 2 required, V_{max}(FB 2), to react completely with • 25.0 cm^3 of **FB 1** if no **FB 3** is added.



[5]

gradient = [1]

(iii) Explain, in terms of the chemistry involved, the direction of the slope of your graph.

• • • • •
 [1]

Calculations

Show your working and appropriate significant figures in **all** of your calculations.

(c) (i) Using appropriate data from your graph, calculate the concentration of ethanedioic acid, $C_2O_4H_2$ in **FB 1**.

concentration of $C_2O_4H_2$ in **FB 1** =[1]

(ii) Using your answer to (c)(i) and appropriate data from your graph, calculate the concentration of sodium hydroxide in FB 2.

(d) Use the expression below to calculate the concentration of sodium hydroxide in **FB 2**.

$$[NaOH] = \frac{2[Na_2CO_3]}{|gradient|}$$

[Turn over

(e) Student B repeats the experiment described in (a). However, student A had used up FB 3. As such student B has to prepare another 250 cm³ of FB 3. Unknowingly, he weighs a sample of solid sodium carbonate which is slightly damp for the preparation of FB 3.

Suggest and explain what effect this will have on the value of $V_{\text{max}}(\mbox{FB 3})$ he obtains.

effect on V_{max}(FB 3)

(f) A student performs the original experiment using solutions of different concentration to those you have used. He determines the gradient of his line using a small triangle.

He calculates the concentration of NaOH in **FB 2**. In part (c)(ii) he obtains a value of 0.589 mol dm⁻³. In part (d) he obtains a value of 0.581 mol dm⁻³.

Suggest which of these two values is likely to be more accurate. Explain your answer.

the more accurate value is
explanation
[1]

(g) Plan an experiment to determine the percentage by mass of sodium ethanedioate, $C_2O_4Na_2$ in **FB 1**.

Acidified potassium manganate(VII) oxidises ethanedioate ions, $C_2O_4^{2-}$ ions, as shown below.

$$2MnO_4^{-}(aq) + 5C_2O_4^{2-}(aq) + 16H^{+}(aq) \rightarrow 2Mn^{2+}(aq) + 10CO_2(g) + 8H_2O(I)$$

However, the reaction takes place slowly at first. The rate of the reaction increases as more products are produced. This is because one of the products acts as a catalyst for the reaction. Hence this reaction is an example of 'autocatalysis'.

(i) Identify the product that acts as the catalyst in the reaction.

......[1]

(ii) Using the information given above, complete the procedures for this additional experiment which is written partially as shown below.

Do **NOT** carry out this experiment.

- 1. Fill a burette with 0.020 mol dm⁻³ of an aqueous solution of potassium manganate(VII).
- 2. Pipette 10.0 cm³ of **FB 1** into a 250 cm³ conical flask.
- 3. Using a 10 cm³ measuring cylinder, add in 10 cm³ of sulfuric acid into the same conical flask.
- 4.

(h) Planning

The volume of aqueous sodium hydroxide needed to completely neutralise all the ethanedioic acid, $C_2O_4H_2$ present in **FB 1** can also be obtained by carrying out a thermometric titration.

The reaction between an acid and an alkali is exothermic. It is possible to make use of this fact to determine the equivalence-point of a neutralisation reaction without the use of an indicator. This process is known as *thermometric titration* and can be used to calculate the concentration of an acid solution and the value of the enthalpy change of neutralisation, ΔH_n .

The concentration of an acid solution can be determined by adding various volumes of acid and alkali and measuring the change in temperature. A series of about six experiments were conducted, where different volumes of alkali were added to a fixed volume (for example 25.0 cm³) of acid while keeping the total volume of the solution constant by adding appropriate volumes of water.

The data obtained is plotted and two best-fit graph lines are drawn. One line is drawn using data before the equivalence-point and the second line using the remaining data. These lines are then extrapolated (extended) until they cross.

(i) Using the information given above, you are required to write a plan for a thermometric titration in which a known concentration of an aqueous solution of sodium hydroxide is added to **FB 1**.

You are to determine the volume of aqueous sodium hydroxide required to completely neutralise the ethanedioic acid present in **FB 1**.

You may also assume that you are provided with:

- **FB 1** which is a mixture of approximately 0.01 mol dm⁻³ aqueous sodium ethanedioate, $C_2O_4Na_2$, and approximately 0.1 mol dm⁻³ ethanedioic acid, $C_2O_4H_2$.
- 0.35 mol dm⁻³ aqueous sodium hydroxide, NaOH
- graph paper;
- the equipment normally found in a school or college laboratory.

Your plan should include:

- calculation of the approximate volume of aqueous sodium hydroxide required to completely neutralise the ethanedioic acid present in 25.0 cm³ of FB 1;
- brief, but specific, details of the apparatus you would use, bearing in mind the levels of precision they offer;
- an outline of how the results would be obtained;
- a table containing the volumes of each reagent to be added
- how you would recognise that the equivalence-point had been passed;
- on the grid provided, a sketch of the graph you would expect to obtain;
- an explanation of the shape of your graph;

9

.....

.....

Explanation of the shape of the graph:

(ii) In another experiment, hydrochloric acid is used instead of FB 1.
 If the hydrochloric acid has the same concentration as the ethanedioic acid present in FB 1, draw on your graph in (i) another pair of lines to show the results you would expect to obtain.

Explain your answer.

[3] [Total: 28]

2 You are required to determine the enthalpy change of combustion of ethanol.

When an exothermic reaction takes place in a container such as a metal calorimeter, some of the evolved heat energy is absorbed by the metal calorimeter.

When an endothermic reaction takes place some of the required heat energy is supplied by the metal calorimeter.

The amount of heat energy evolved or supplied for a 1 °C change in temperature is known as the heat capacity of the metal calorimeter.

In preparation for your experiment to determine the enthalpy change of combustion of ethanol, you will first need to determine the approximate heat capacity of a 100 cm³ metal calorimeter.

Before starting any practical work read through the instructions in (a) and draw up a table to record your results.

(a) Determining the approximate heat capacity of the 100 cm³ metal calorimeter

When samples of hot and cold water are mixed in the 100 cm³ metal calorimeter, some heat is lost to the metal calorimeter in raising its temperature. To determine the approximate heat capacity of your 100 cm³ metal calorimeter, you will determine the maximum temperature rise when a sample of hot water is added to cold water in the metal calorimeter.

- Use a 50 cm³ measuring cylinder to transfer 50 cm³ of cold water into the 100 cm³ metal calorimeter.
- Use the 50 cm³ measuring cylinder to transfer 50 cm³ of cold water into a 100 cm³ beaker.
- Note the temperature of the water in this 100 cm³ beaker and heat it **carefully and gently** until the temperature of the water in it has increased by 45–50 °C then stop heating, *e.g. if the water is at 20.0* °C you should warm it to 65–70 °C.
- Stir the cold water in the 100 cm³ metal calorimeter with the thermometer.
- Record the temperature of the cold water (this is the temperature at t = 0 min).
- Record the temperature each minute for 3 minutes.
- After you have taken the reading at t = 3 min, use the thermometer to stir the hot water in the 100 cm³ beaker.
- At t = 4 min, measure the temperature of the hot water and record this value in the box below.
- **Immediately** add the hot water from the 100 cm³ beaker to the cold water in the 100 cm³ metal calorimeter.
- Stir with the thermometer but do not record the temperature.
- Continue to stir the water throughout the experiment.
- Record the temperature at t = 5 min, and then every $\frac{1}{2}$ minute until t = 8 min.
- Record all measurements of time and temperature obtained on page 13.

The temperature, T_1 , of the hot water at t = 4 min is°C.

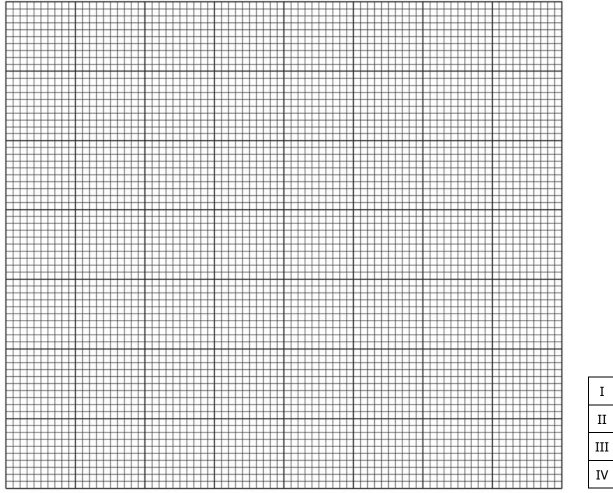
Table of results

Ι	
II	
III	

[3]

(b) Graph plotting

- 1. Plot a graph of the temperature of the water in the 100 cm³ metal calorimeter (y-axis) against time (x-axis) on the grid below. Do not plot the temperature, T_1 , of the hot water at t = 4 min.
- 2. Draw two straight lines of best fit; one through the points up to t = 3 min; the second through the points from t = 5 min to t = 8 min. Extrapolate both lines to t = 4 min.
- From the extrapolated lines read the minimum and the maximum temperatures at t = 4 min. Record these values in the spaces provided below.
- 4. Determine the values for the two temperature changes at t = 4 min.



Ι	
II	
III	
IV	

Minimum temperature, T_2 , at t = 4 min is°C.Maximum temperature, T_3 , at t = 4 min is°C.Temperature rise for 50 cm³ of cold water in the 100 cm³ metal calorimeter, $(T_3 - T_2)$ is°C.Temperature fall for 50 cm³ of hot water from the 100 cm³ beaker, $(T_1 - T_3)$ is°C.[4]

(c) Calculations

Working should be shown in all calculations. [4.2 J are absorbed or released when the temperature of 1.0 cm^3 of water changes by 1.0 °C.]

(i) Calculate the heat energy gained by the 50 cm³ of cold water in the 100 cm³ metal calorimeter.

The heat energy gained by the cold water =J.

(ii) Calculate the heat energy lost by the 50 cm³ of hot water from the 100 cm³ beaker.

The heat energy lost by the hot water =J.

(iii) The difference between the values calculated in (i) and (ii) is an approximate value for the total heat energy absorbed by the 100 cm³ metal calorimeter during the experiment. The heat capacity of the metal calorimeter is the amount of heat energy absorbed for a 1 °C change in temperature.

approximate heat capacity of the 100 cm³ metal calorimeter = $\frac{(heat energy lost) - (heat energy gained)}{(T_3 - T_2)}$ J °C⁻¹

Use your answers to (i) and (ii) and the temperature rise from (b) to calculate the approximate heat capacity of the 100 cm³ metal calorimeter.

The approximate heat capacity of the 100 cm³ metal calorimeter = J $^{\circ}C^{-1}$. [1]

(d) Determining the enthalpy change of combustion of ethanol, C₂H₅OH

An experiment was carried out as follows to determine the enthalpy change of combustion of ethanol.

The 100 cm³ metal calorimeter used in (a) was used to contain some water. The water was heated using ethanol placed in a spirit lamp. The temperature rise was recorded. The spirit lamp was weighed before and after the experiment to determine the mass of ethanol used. The following results were obtained.

mass of ethanol burned = 0.391 g mass of water heated = 40.0 g temperature rise = 19.5 °C

(i) Complete the following diagram to show the experimental set-up needed to carry out the experiment to determine the enthalpy change of combustion of ethanol.

You are to use the following apparatus to complete the diagram:

- stand and clamp
- 100 cm³ metal calorimeter
- thermometer
- tripod stand and heat-proof mat



spirit lamp containing ethanol

(ii) Use your answer in (c)(iii) and the results obtained to calculate the enthalpy change of combustion of ethanol.
[Ar: O, 16.0; C, 12.0; H, 1.0]
[specific heat capacity of water = 4.2 J °C⁻¹ cm⁻³]
(If you were unable to answer (c)(iii), you may assume that the heat capacity of the 100 cm³ metal calorimeter is 50 J °C⁻¹.)

enthalpy change of combustion of ethanol =[1]

(iii) Calculate the maximum percentage error in the measurement of each mass used in the experiment.

mass measured	maximum error in a single reading	maximum percentage error / %
0.391 g of ethanol burned	0.0005 g	
40.0 g of water	0.05 g	

[1]

[Total: 11]

3 Qualitative Analysis

At each stage of any test you are to record details of the following.

- colour changes seen
- the formation of any precipitate
- the solubility of such precipitates in an excess of the reagent added

Where gases are released they should be identified by a test, **described in the** appropriate place in your observations.

You should indicate clearly at what stage in a test a change occurs. No additional tests for ions present should be attempted.

If any solution is warmed, a boiling tube MUST be used.

Rinse and reuse test-tubes and boiling tubes where possible.

Where reagents are selected for use in a test, the name or correct formula of the element or compound must be given.

FB 4 is a solution containing **one** cation and **one** anion from the Qualitative Analysis Notes in pages 22 and 23.

Solid **FB 5** contains the sodium ion and **one** anion from the Qualitative Analysis Notes in pages 22 and 23.

(a) Carry out the following tests on **FB 4** and state the observations you make.

	Test	Observations
(i)	To 1 cm depth of aqueous FB 4 in a test-tube, add aqueous ammonia, then	
	add sulfuric acid dropwise until there is no further observations.	
(ii)	To 1 cm depth of aqueous FB 4 in a test-tube, add silver nitrate,	
	followed by aqueous ammonia.	

For Examiner's Use			
Obs points		marks	

	Test	Observations
(i)	Place half of the solid FB 5 in a boiling tube. Heat strongly until the solid melts and a gas is given off.	
	Test and identify the gas.	
	Continue the strong heating for 2-3 min,	
	then leave the tube to cool and retain the residue for the test in (iv) .	
(ii)	To the remaining half of FB 5 in the boiling tube, add 3 cm depth of aqueous sodium hydroxide.	
	Warm gently and retain for the test in (iii) .	
(iii)	To the solution from the test in (ii) , add a piece of aluminium foil and warm gently.	
(iv)	Dissolve the residue from test in (i) in distilled water and divide the solution into three parts. Use these for the tests in (v) to (vii).	
(v)	To one part of the solution from (iv) , add aqueous potassium iodide followed by dilute sulfuric acid.	
(vi)	To the second part of the solution from (iv), add aqueous potassium manganate (VII) followed by dilute sulfuric acid.	
(vii)	To the third part of the solution (iv), add aqueous sodium hydroxide and a piece of aluminium foil. Warm gently.	

(b) Carry out the following tests on **FB 5** and state the observations you make.

(c)	 c) (i) Suggest the identities of the cation and anion present in FE observations from the tests in (a). 		3 4 from your	
		cation: anion:	[1]	
	(ii)	Write equations, with state symbols to explain your observations for te in a(i).	est	
			[3]	
(d)	(i)	Suggest the identity of the anion present in FB5 before it was heated	d.	
			[1]	
	(ii)	Suggest the identity of the anion present in FB5 after it was heated.		
			[1]	
	(iii)	In (b)(v) , the anion is behaving as	[1]	
	(iv)	In (b)(vi) , the anion is behaving as	[1]	
	(v)	Suggest the purpose of carrying out test (b)(ii).		
			[1]	

(e) Student C carried out the following experiment on **FB 5**, predict the observations that will be observed and write your answer in the blanks.

Test	Observations
To solid FB 5 in a boiling tube. Heat heat strongly for 4 min. Let the residue cool. Add dilute sulfuric acid to it.	

[1]

[Total: 16]

Qualitative Analysis Notes [ppt. = precipitate]

(a) Reactions of aqueous cations

ootion	reaction with					
cation	NaOH(aq)	NH₃(aq)				
aluminium, A <i>l</i> ³⁺(aq)	white ppt. soluble in excess	white ppt. insoluble in excess				
ammonium, NH₄⁺(aq)	ammonia produced on heating	_				
barium, Ba ²⁺ (aq)	no ppt. (if reagents are pure)	no ppt.				
calcium, Ca²⁺(aq)	white ppt. with high [Ca ²⁺ (aq)]	no ppt.				
chromium(III), Cr³⁺(aq)	grey-green ppt. soluble in excess giving dark green solution	grey-green ppt. insoluble in excess				
copper(II), Cu ²⁺ (aq)	pale blue ppt. insoluble in excess	blue ppt. soluble in excess giving dark blue solution				
iron(II), Fe ²⁺ (aq)	green ppt., turning brown on contact with air insoluble in excess	green ppt., turning brown on contact with air insoluble in excess				
iron(III), Fe³⁺(aq)	red-brown ppt. insoluble in excess	red-brown ppt. insoluble in excess				
magnesium, Mg²⁺(aq)	white ppt. insoluble in excess	white ppt. insoluble in excess				
manganese(II), Mn²⁺(aq)	off-white ppt., rapidly turning brown on contact with air insoluble in excess	off-white ppt., rapidly turning brown on contact with air insoluble in excess				
zinc, Zn ²⁺ (aq)	white ppt. soluble in excess	white ppt. soluble in excess				

(b) Reactions of anions

anion	reaction
carbonate, CO ₃ ^{2–}	CO ₂ liberated by dilute acids
chloride , C <i>l</i> È(aq)	gives white ppt. with Ag⁺(aq) (soluble in NH₃(aq))
bromide, BrÈ(aq)	gives pale cream ppt. with Ag ⁺ (aq) (partially soluble in $NH_3(aq)$)
iodide, IÈ(aq)	gives yellow ppt. with Ag⁺(aq) (insoluble in NH₃(aq))
nitrate, NO₃È(aq)	NH_3 liberated on heating with $OHE(aq)$ and Al foil
nitrite, NO₂È(aq)	NH ₃ liberated on heating with OH $\dot{\mathbb{E}}$ (aq) and A <i>l</i> foil; NO liberated by dilute acids (colourless NO \rightarrow (pale) brown NO ₂ in air)
sulfate, SO ₄ ²⁻(aq)	gives white ppt. with Ba ²⁺ (aq) (insoluble in excess dilute strong acids)
sulfite, SO ₃ ²⁻(aq)	SO ₂ liberated with dilute acids; gives white ppt. with Ba ²⁺ (aq) (soluble in dilute strong acids)

(c) Tests for gases

gas	test and test result
ammonia, NH₃	turns damp red litmus paper blue
carbon dioxide, CO ₂	gives a white ppt. with limewater (ppt. dissolves with excess CO ₂)
chlorine, Cl_2	bleaches damp litmus paper
hydrogen, H ₂	"pops" with a lighted splint
oxygen, O ₂	relights a glowing splint
sulfur dioxide, SO ₂	turns aqueous acidified potassium manganate(VII) from purple to colourless

(d) Colour of halogens

halogen	colour of element	colour in aqueous solution	colour in hexane
chlorine, Cl ₂	greenish yellow gas	pale yellow	pale yellow
bromine, Br ₂	reddish brown gas / liquid	orange	orange-red
iodine, I	black solid / purple gas	brown	purple

NANYANG JUNIOR COLLEGE JC2 PRACTICAL PRELIMINARY EXAMINATION Higher 2						
CANDIDATE NAME Mark Scheme						
CLASS 1 7 REG TUTOR						
CHEMISTRY 9729/04 Paper 4 Practical 28 August 2018 Candidates answer on the Question Paper 2 hour 30 minutes Additional Materials: As listed in the Confidential Instructions						
READ THESE INSTRUCTIONS FIRST						
Write your name and class on all the work you hand in. Give details of the practical shift and laboratory, where appropriate, in the boxes pro 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.	vided.					
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		Shift				
appropriate units. Qualitative Analysis Notes are printed on pages 22 and 23.	La	aboratory				
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					
	2	/ 28				
	3	/16				
	Total	/55				

1

This document consists of 23 printed pages and 1 blank page.

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

1 To determine the concentrations of aqueous sodium hydroxide and aqueous ethanedioic acid and hence the percentage by mass of sodium ethanedioate in a mixture of sodium ethanedioate and ethanedioic acid.

You are provided with the following reagents.

- **FB 1** is a mixture of approximately 0.01 mol dm⁻³ aqueous sodium ethanedioate, C₂O₄Na₂, and approximately 0.1 mol dm⁻³ ethanedioic acid, C₂O₄H₂
- **FB 2** is a dilute solution of sodium hydroxide, NaOH
- **FB 3** is 0.0755 mol dm⁻³ aqueous sodium carbonate
- thymolphthalein indicator

Both sodium hydroxide and sodium carbonate are bases which will neutralise ethanedioic acid as shown in **reactions 1** and **2**.

$$2NaOH + C_2O_4H_2 \rightarrow C_2O_4Na_2 + 2H_2O$$
 reaction 1

$$Na_2CO_3 + C_2O_4H_2 \rightarrow C_2O_4Na_2 + CO_2 + H_2O$$
 reaction 2

In this experiment, you will add different volumes of **FB 3** to identical samples of the mixture of aqueous sodium ethanedioate, $C_2O_4Na_2$, and ethanedioic acid, $C_2O_4H_2$, **FB 1**. You will then complete the neutralisation of $C_2O_4H_2$ in each mixture by titration with dilute sodium hydroxide, **FB 2**.

Each titration is to be performed **once only**, so great care should be taken that you do not overshoot the end-points.

Graphical analysis of your results will enable you to determine the concentrations of the ethanedioic acid in the mixture of aqueous $C_2O_4Na_2$, and $C_2O_4H_2$ and of the sodium hydroxide.

Read through the whole method of conducting the experiment before starting any practical work.

The experiment

(a) Titrations

Experiment 1

- 1. Fill a burette with **FB 3**.
- 2. Pipette 25.0 cm³ of **FB 1** into a 250 cm³ conical flask.
- 3. Run 5.00 cm³ of **FB 3** into the flask.
- 4. Add 10 15 drops of thymolphthalein indicator to the flask.
- 5. Fill a second burette with **FB 2**.
- 6. Titrate the mixture in the conical flask against **FB 2** until the first permanent pale blue colour remains in the solution.
- 7. Rinse the conical flask thoroughly before it is used for the next experiment.

The end-point should be found after the addition of approximately 14 cm³ of **FB 2**.

One titration, performed accurately, will be sufficient.

Experiment 5

1. Repeat **Experiment 1** but run 30.00 cm³ of **FB 3** into the flask containing 25.0 cm³ of **FB 1**.

Experiments 2 – 4

Select **three** other suitable volumes of **FB 3** for use in **Experiments 2 – 4**. Your selected volumes must be between the volumes used in **Experiments 1 and 5**.

Do not use volumes of FB 3 outside the range of 5 – 30 cm³.

Repeat the procedures used in **Experiment 1** but, in each case, add 25.0 cm³ of **FB 1** and your chosen volume of **FB 3** into the conical flask. You should perform your titrations in order of **increasing** volume of **FB 3**.

Rinse this conical flask thoroughly between each titration.

Prepare a table in the space below and use it to record the titration results for each volume of **FB 3** added. You should record your titration results in order of **increasing** volume of **FB 3** added.

Experiment	1	2	3	4	5
Volume of FB 3 added / cm ³	5.00	11.00	18.00	24.00	30.00
Initial burette reading / cm ³	0.00	20.00	0.00	11.70	21.60
Final burette reading / cm ³	15.80	34.00	11.70	21.60	29.80
Volume of FB 2 added / cm ³	15.80	14.00	11.70	9.90	8.20

[1]: Correct column or row headings and units for titration recordings

Acceptable forms of units:

Use of solidus, e.g. / cm³

Unit in brackets, e.g. (cm³)

In words, e.g. volume in cubic centimetres

AND

All burette readings and/or volumes/titres recorded to 2 dp or to nearest 0.05 cm³)

[1]: Follows instructions.

Selects three additional volumes of **FB 3** which are within the range 5 - 30 cm³ for experiments 2, 3, and 4.

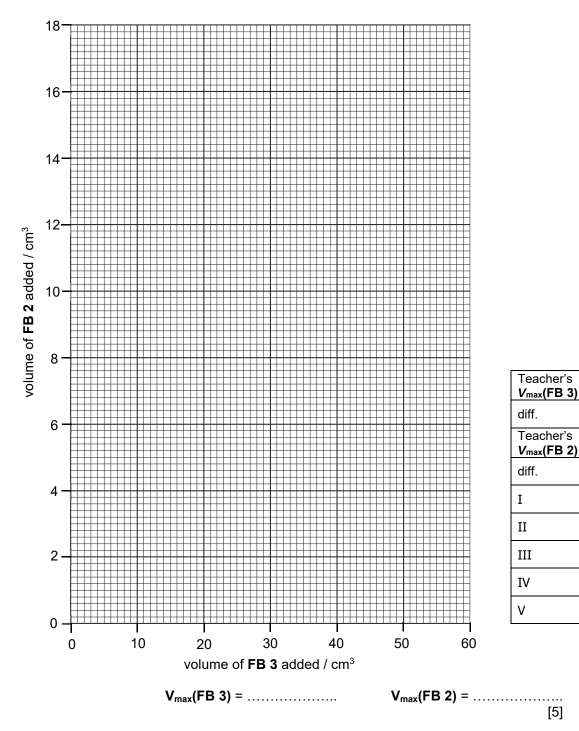
AND All gaps between adjacent **FB 3** volumes should be $\ge 5 \text{ cm}^3$ and $\le 10 \text{ cm}^3$ **AND** Completes a total of 5 experiments.

[2]

(b) (i) Plot, on the grid below, your values for the FB 2 titre (y-axis) against the volume of **FB 3** added (*x-axis*).

Draw the line of best fit, taking into account all of your plotted points. Hence obtain values for

- the volume of FB 3 required, V_{max}(FB 3), to react completely with • 25.0 cm³ of **FB 1** if no **FB 2** is added;
- the volume of FB 2 required, V_{max}(FB 2), to react completely with • 25.0 cm^3 of **FB 1** if no **FB 3** is added.

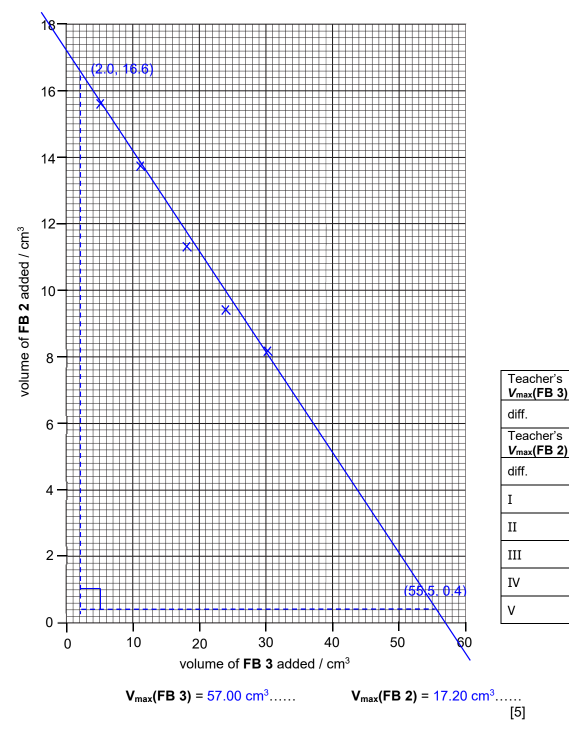


[5]

(b) (i) Plot, on the grid below, your values for the **FB 2** titre (*y*-axis) against the volume of **FB 3** added (*x*-axis).

Draw the line of best fit, taking into account all of your plotted points. Hence obtain values for

- the volume of FB 3 required, V_{max}(FB 3), to react completely with 25.0 cm³ of FB 1 if no FB 2 is added;
- the volume of FB 2 required, V_{max}(FB 2), to react completely with 25.0 cm³ of FB 1 if no FB 3 is added.



[1]: All points plotted to within 1/2 small square in either direction and in the correct square.

[1]: The graph line is the <u>straight</u> best-fit line AND

Line is correctly extrapolated to intersect both axes. i.e. one continuous straight line **AND**

no point is further than 1 cm³ away from the line in either direction

[2]: Accuracy for V_{max}(FB 3) and V_{max}(FB 2)

[1] Reads correctly, to $\pm \frac{1}{2}$ small square, the value for $V_{max}(FB 3)$ and the value for $V_{max}(FB 2)$

If the student has not shown the extrapolation on the graph, the student may still earn this mark if the values quoted correspond to correct extrapolation of the student's graph line.

(ii) Calculate the gradient of your graph line, showing clearly how you did this.

gradient =
$$\frac{16.6 - 0.4}{2 - 55.5}$$
 = -0.3028 \approx -0.303

gradient = -0.303.....[1]

To earn this mark, students must

- Use clearly stated co-ordinates <u>from graph</u> or correct values of *∆titre* and of *∆(volume* of *FB 3 used)* obtained from a clearly drawn triangle (allow measurements to ± ½ small square). Coordinates could be from V_{max} values.
- Use a triangle (or equivalent coordinates/∆ values) which covers at least <u>3 large squares</u> in each direction
- Correctly calculate gradient. (ignore sig figs for this mark)
- Do not award this mark if the graph line is not straight.

For this mark, ignore missing working and incorrect/missing units.

(iii) Explain, in terms of the chemistry involved, the direction of the slope of your graph.

[1]: Negative gradient because the more Na_2CO_3 added, the less NaOH is needed for neutralisation (or words to that effect)

Calculations

Show your working and appropriate significant figures in **all** of your calculations.

(c) (i) Using appropriate data from your graph, calculate the concentration of ethanedioic acid, $C_2O_4H_2$ in **FB 1**.

1 mol of Na₂CO₃ will completely neutralise 1 mol of C₂O₄H₂

(ii) Using your answer to (c)(i) and appropriate data from your graph, calculate the concentration of sodium hydroxide in FB 2.

2 mol of NaOH will completely neutralise 1 mol of C₂O₄H₂ V_{max}(**FB 2**) = 17.20 cm³ [NaOH] = $\frac{57.00/1000 \times 0.0755 \times 2}{17.20/1000}$ = 0.5004 \approx 0.500 mol dm⁻³

concentration of sodium hydroxide in **FB 2** = $0.500 \text{ mol } \text{dm}^{-3} \dots [1]$

(d) Use the expression below to calculate the concentration of sodium hydroxide in **FB 2**. Give your answer to three significant figures.

$$[NaOH] = \frac{2[Na_2CO_3]}{|gradient|}$$

 $[NaOH] = \frac{2(0.0755)}{0.3028} = 0.4986 \approx 0.499 \text{ mol dm}^{-3}$

concentration of sodium hydroxide in **FB 2** = $0.499 \text{ mol } \text{dm}^{-3}$[1]

[1 mark]
Correctly calculates [NaOH] using the given expression
Show working in (c)(ii), (d) and (e)
3 significant figures in final section answers to (c)(ii), (d) and (e)
Show appropriate units in answers in (c)(i), (c)(ii), (d)(ii), (d)(iii) and (e)

(e) Student B repeats the experiment described in (a). However, student A had used up FB 3. As such student B has to prepare another 250 cm³ of FB 3. Unknowingly, he weighs a sample of solid sodium carbonate which is slightly damp for the preparation of FB 3.

Suggest and explain what effect this will have on the value of $V_{\text{max}}(\text{FB 3})$ he obtains.

effect on V _{max} (FB 3) value of V _{max} (FB 3) too high
explanation
[1]

presence of water means mass of Na_2CO_3 weighed out is lower than expected, so $[Na_2CO_3]$ in **FB 3** is lower than expected, so more **FB 3** is needed to neutralise the ethanedioic acid in **FB 1**.

(f) A student performs the original experiment using solutions of different concentration to those you have used. He determines the gradient of his line using a small triangle.

He calculates the concentration of NaOH in **FB 2**. In part (c)(ii) he obtains a value of 0.589 mol dm⁻³. In part (d) he obtains a value of 0.581 mol dm⁻³.

Suggest which of these two values is likely to be more accurate. Explain your answer.

(g) Plan an experiment to determine the percentage by mass of sodium ethanedioate, $C_2O_4Na_2$ in **FB 1**.

Acidified potassium manganate(VII) oxidises ethanedioate ions, $C_2O_4^{2-}$ ions, as shown below.

$$2MnO_4^{-}(aq) + 5C_2O_4^{2-}(aq) + 16H^{+}(aq) \rightarrow 2Mn^{2+}(aq) + 10CO_2(g) + 8H_2O(I)$$

However, the reaction takes place slowly at first. The rate of the reaction increases as more products are produced. This is because one of the products acts as a catalyst for the reaction. Hence this reaction is an example of 'autocatalysis'.

(i) Identify the product that acts as the catalyst in the reaction.

......[1]

(ii) Using the information given above, complete the procedures for this additional experiment which is written partially as shown below.

Do **NOT** carry out this experiment.

- 1. Fill a burette with 0.020 mol dm⁻³ of an aqueous solution of potassium manganate(VII).
- 2. Pipette 10.0 cm³ of **FB 1** into a 250 cm³ conical flask.
- 3. Using a 10 cm³ measuring cylinder, add in 10 cm³ of sulfuric acid into the same conical flask.
- 4.

Heat the mixture in the conical flask to about 60/70/80 °C.

- 5. Titrate the mixture in the conical flask against potassium manganate(VII) until the first permanent pale pink..... colour remains in the solution.
- 6. Repeat the titration

.....

.....

Repeat the titration as many times necessary to achieve two consistent readings (to be within ± 0.10 cm³. The average of the two consistent readings will be the volume of potassium manganate(VII) added to achieve endpoint.)

(h) Planning

The volume of aqueous sodium hydroxide needed to completely neutralise all the ethanedioic acid, $C_2O_4H_2$ present in **FB 1** can also be obtained by carrying out a thermometric titration.

The reaction between an acid and an alkali is exothermic. It is possible to make use of this fact to determine the equivalence-point of a neutralisation reaction without the use of an indicator. This process is known as *thermometric titration* and can be used to calculate the concentration of an acid solution and the value of the enthalpy change of neutralisation, ΔH_n .

The concentration of an acid solution can be determined by adding various volumes of acid and alkali and measure the change in temperature. A series of about six experiments were conducted, where different volumes of alkali were added to a fixed volume (for example 25.0 cm³) of acid while keeping the total volume of the solution constant by adding appropriate volumes of water.

The data obtained is plotted and two best-fit graph lines are drawn. One line is drawn using data before the equivalence-point and the second line using the remaining data. These lines are then extrapolated (extended) until they cross.

(i) Using the information given above, you are required to write a plan for a thermometric titration in which a known concentration of an aqueous solution of sodium hydroxide is added to **FB 1**.

You are to determine the volume of aqueous sodium hydroxide required to completely neutralise the ethanedioic acid present in **FB 1**.

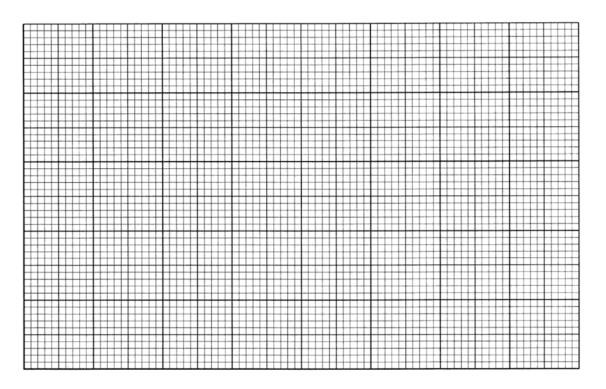
You may also assume that you are provided with:

- **FB 1** which is a mixture of approximately 0.01 mol dm⁻³ aqueous sodium ethanedioate, $C_2O_4Na_2$, and approximately 0.1 mol dm⁻³ ethanedioic acid, $C_2O_4H_2$.
- 0.35 mol dm⁻³ sodium hydroxide, NaOH
- graph paper;
- the equipment normally found in a school or college laboratory.

Your plan should include:

- calculation of the approximate volume of aqueous sodium hydroxide required to completely neutralise the ethanedioic acid present in FB 1;
- brief, but specific, details of the apparatus you would use, bearing in mind the levels of precision they offer;
- an outline of how the results would be obtained;
- a table containing the volumes of each reagent to be added
- how you would recognise that the equivalence-point had been passed;
- on the grid provided, a sketch of the graph you would expect to obtain;
- an explanation of the shape of your graph;

.....



......[8]

[1] Calculation:

Given approx. $[C_2O_4H_2] = 0.1 \text{ mol } dm^{-3} \text{ and } [NaOH] = 0.35 \text{ mol } dm^{-3}$ Given fix acid volume at 25.0 cm³

 $n(C_2O_4H_2) = \frac{25.0}{1000} \times 0.1 = 0.0025 \text{ mol}$

 $n(NaOH) = 2n(C_2O_4H_2) = 0.0025 \times 2 = 0.005 mol$

approximate $V_{max}(FB 2) = 0.005 / 0.35 \times 1000 = 14.29 \text{ cm}^3$ (ignore the precision when presenting this answer)

Procedure:

- Use a burette to transfer 25.00 cm³ of FB 1 into a styrofoam cup. (Place the cup in a 250 cm³ beaker to prevent it tipping over.) Record the initial temperature of FB 1 using a thermometer graduated in 0.2 °C.
- 2. From a second burette, measure the required volume of water as shown in the table below and add to the **FB 1** in the styrofoam cup. (Use the thermometer to stir the mixture gently.)
- 3. From a third burette, measure the required volume of NaOH as shown in the table below into a 100 cm³ beaker and add to **FB 1** in the styrofoam cup. Use the thermometer to stir the mixture gently.
- 4. Record the highest temperature reached.
- 5. Repeat steps **1** to **4** by varying the volume of water and NaOH added to **FB 1** as shown in the table below.
- 6. The temperature change, $\Delta T = T_{\text{final}} T_{\text{initial}}$, where T_{final} is the highest temperature measured and T_{initial} is the initial temperature of **FB 1**.
- 7. Plot a graph of the ΔT against the volume of NaOH added.
- 8. The equivalence-point is reached once the temperature is observed to stop rising.

Experiment	Volume of NaOH / cm ³	Volume of FB 1 / cm ³	Volume of water / cm ³	Initial temperature of FB 1 / °C	Highest temperature / °C	∆T / °C
1	8.00	25.00	22.00			
2	12.00	25.00	18.00			
3	14.00	25.00	16.00			
4	16.00	25.00	14.00			
5	18.00	25.00	12.00			
6	20.00	25.00	10.00			

Marking Scheme

1 mark for describing the followings in the procedure:

- state the use of burette to measure the volumes of water, NaOH and FB 1
- state the precision of the thermometer to be used
- use styrofoam cup as the container for the solution mixture

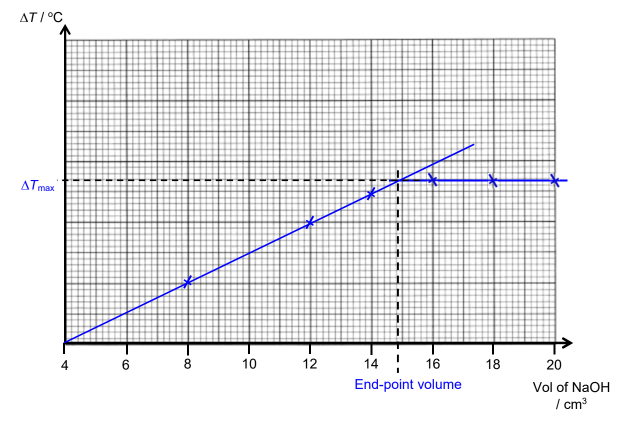
1 mark for describing the followings in the procedure:

- measure out a fixed volume of **FB 1** and determine its initial temperature
- mix **FB 1**, NaOH and water (either **FB 1** or NaOH should be the last reagent to be added) and record the highest temperature obtained
- repeat the experiment by varying the volumes of NaOH and water to be added and ensuring that the total volume of the reaction mixture remains constant

1 mark for selecting an appropriate fixed volume of **FB 1** and range of volume of water and NaOH taking into account the followings:

- The minimum fixed volume of reaction mixture is 25.00 cm³ so that the bulb of the thermometer is completely immersed into the solution when measuring the initial temperature
- Using the approximate V_{max}(FB 2) calculated to select the appropriate <u>range of the</u> volume of NaOH to be added, i.e. 3 volumes less than 14 cm³ and 3 volumes greater than 14 cm³
- Adding appropriate volumes of water to ensure that the <u>total volume of the reaction</u> <u>mixture is fixed</u> and preferably be less than 100 cm³ taking into account the maximum volume of the polystyrene cup used and to prevent spillage upon mixing

1 mark for writing step **8** in the above procedure



Sketch of graph expected to be obtained:

Marking Scheme

1 mark for

- labelling the axes correctly and indicating the scales on the x axis
- indicating the end-point volume of NaOH

1 mark for drawing a linear, increasing line and a linear, horizontal line. These lines are then extrapolated until they cross. The point of intersection is the equivalence point.

Explanation of the shape of the graph:

Before equivalence point, the limiting reagent is NaOH, hence both the number of moles of water and the total volume of the solution are increasing. According to

$$\Delta H_n = -\frac{m \times C/1000 \times \Delta T}{n_{H,0}}, \Delta T \text{ is directly proportional to the number of moles of water.}$$

Since the number of moles of water increases with increasing volume of NaOH so ΔT increases.

After the equivalence point, the limiting reagent is **FB 3**. The number of moles of water which depend on **FB 3** remain constant as a fixed volume of **FB 3** is used. Hence ΔT remains unchanged.

1 mark for sensible explanation

(ii) In another experiment, hydrochloric acid is used instead of FB 1.
 If the hydrochloric acid has the same concentration as the ethanedioic acid present in FB 1, draw on your graph in (i) another pair of lines to show the results you would expect to obtain.

Explain your answer.

[3]

[1]: Line rises more steeply and intersects second line at a higher temperature rise

[1]: Maximum is reached at ½V(NaOH) needed to completely neutralised the ethanedioic acid in **FB 1**

[1]: Some of the heat that would have been released is used to ionise the ethanedioic acid and the reacting mole ratio between hydrochloric acid and sodium hydroxide is 1 : 1.

[Total: 28]

2 You are required to determine the enthalpy change of combustion of ethanol.

When an exothermic reaction takes place in a container such as a metal calorimeter, some of the evolved heat energy is absorbed by the metal calorimeter.

When an endothermic reaction takes place some of the required heat energy is supplied by the metal calorimeter.

The amount of heat energy evolved or supplied for a 1 °C change in temperature is known as the heat capacity of the metal calorimeter.

In preparation for your experiment to determine the enthalpy change of combustion of ethanol, you will first need to determine the approximate heat capacity of a 100 cm³ metal calorimeter.

Before starting any practical work read through the instructions in (a) and draw up a table to record your results.

(a) Determining the approximate heat capacity of the 100 cm³ metal calorimeter

When samples of hot and cold water are mixed in the 100 cm³ metal calorimeter, some heat is lost to the metal calorimeter in raising its temperature. To determine the approximate heat capacity of your 100 cm³ metal calorimeter, you will determine the maximum temperature rise when a sample of hot water is added to cold water in the metal calorimeter.

- Use a 50 cm³ measuring cylinder to transfer 50 cm³ of cold water into the 100 cm³ metal calorimeter.
- Use the 50 cm³ measuring cylinder to transfer 50 cm³ of cold water into a 100 cm³ beaker.
- Note the temperature of the water in this 100 cm³ beaker and heat it **carefully and gently** until the temperature of the water in it has increased by 45–50 °C then stop heating, *e.g. if the water is at 20.0* °C you should warm it to 65–70 °C.
- Stir the cold water in the 100 cm³ metal calorimeter with the thermometer.
- Record the temperature of the cold water (this is the temperature at t = 0 min).
- Record the temperature each minute for 3 minutes.
- After you have taken the reading at t = 3 min, use the thermometer to stir the hot water in the 100 cm³ beaker.
- At t = 4 min, measure the temperature of the hot water and record this value in the box below.
- **Immediately** add the hot water from the 100 cm³ beaker to the cold water in the 100 cm³ metal calorimeter.
- Stir with the thermometer but do not record the temperature.
- Continue to stir the water throughout the experiment.
- Record the temperature at t = 5 min, and then every $\frac{1}{2}$ minute until t = 8 min.
- Record all measurements of time and temperature obtained.

The temperature, T_1 , of the hot water at t = 4 min is 72.0....°C.

Table of results

time / s	0	1	2	3	5	5.5	6	6.5
temperature / °C	31.5	31.5	31.5	31.5	49.0	49.0	48.5	48.0

time / s	7	7.5	8
temperature / °C	48.0	47.5	47.0

[3]

[1]: Follows instructions with regard to times and temperature readings 0-3 minutes at 1 minute intervals; 5-8 minutes at $\frac{1}{2}$ minute intervals, **and T**₁ recorded in box. (Ignore if also in table)

[1]: All columns correctly labelled with appropriate unit shown.

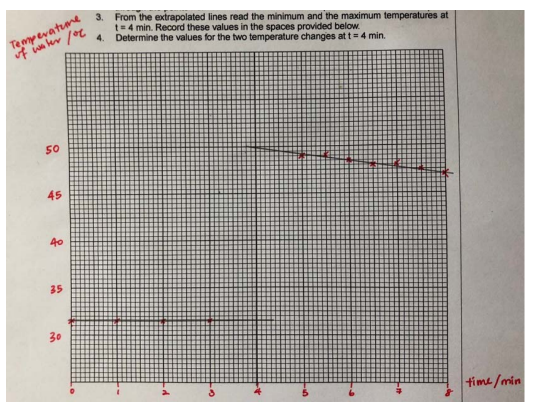
Must use solidus, brackets or describe fully in words.

If units not included in column headings every entry must have correct unit shown Accept min, mins, minutes

[1]: All **thermometer readings** (table and box) recorded to nearest 0.5 °C. *Ignore the precision for the recording of time.*

(b) Graph plotting

- 1. Plot a graph of the temperature of the water in the 100 cm³ metal calorimeter (y-axis) against time (x-axis) on the grid below. Do not plot the temperature, T_1 , of the hot water at t = 4 min.
- 2. Draw two straight lines of best fit; one through the points up to t = 3 min; the second through the points from t = 5 min to t = 8 min. Extrapolate both lines to t = 4 min.
- 3. From the extrapolated lines read the minimum and the maximum temperatures at t = 4 min. Record these values in the spaces provided below.
- 4. Determine the values for the two temperature changes at t = 4 min.



Minimum temperature, T_2 , at t = 4 min is 31.5......°C.
Maximum temperature, T_3 , at t = 4 min is 50.0......°C.
Temperature rise for 50 cm³ of cold water in the 100 cm³ metal calorimeter, $(T_3 - T_2)$ is
18.5.......°C.
Temperature fall for 50 cm³ of hot water from the 100 cm³ beaker, $(T_1 - T_3)$ is 22.0.......°C.
[4]

[1]: Temperature of water in the metal calorimeter plotted on y-axis against time on x-axis. NYJC mark scheme: Clearly labelled axes. Units required. Allow ECF from how headers are labelled in table in **(a)**.

CIE mark scheme and FYI only: Clearly labelled axes (ignore units) [temp/time are minimum acceptable labels] but accept **T** / °**C** and **t** / **min** as labels. The unit is necessary in this case

[1]: Uniform and sensible scales for candidate's choice of graph.

Plotted points must be in at least 4 large squares on the temperature axis and 5 large squares on the time axis.

Do **not** include any plotted value of **T**₁.

[1]: There should be a minimum of 5 plotted points between 5 and 8 minutes.

Examiner then checks plotting of points at t_0 min, t_5 min and t_8 min and the plotting of any suspect point.

If any of the $t_0 \min$, $t_5 \min$ and $t_8 \min$ points is missing check the adjacent point. Points should be within $\frac{1}{2}$ of a small square of the correct position and in the correct small square

[1]: Acceptable straight lines drawn – an acceptable straight line is one passing through the majority of points or has balanced points on either side of the line **and**

correct values of T_2 and T_3 read (to within $\frac{1}{2}$ small square) from the graph.

(c) Calculations

Working should be shown in all calculations.

[**4.2 J** are absorbed or released when the temperature of 1.0 cm³ of **water** changes by 1.0 °C.]

(i) Calculate the heat energy gained by the 50 cm³ of cold water in the 100 cm³ metal calorimeter.

heat energy gained by cold water = $50 \times 4.2 \times 18.5 = 3885 \approx 3890 \text{ J}$

The heat energy gained by the cold water = 3890......J.

(ii) Calculate the heat energy lost by the 50 cm³ of hot water from the 100 cm³ beaker.

heat energy lost by hot water = $50 \times 4.2 \times 22.0 = 4620$ J

The heat energy lost by the hot water = 4620.....J.

(iii) The difference between the values calculated in (i) and (ii) is an approximate value for the total heat energy absorbed by the 100 cm³ metal calorimeter during the experiment. The heat capacity of the metal calorimeter is the amount of heat energy absorbed for a 1 °C change in temperature.

approximate heat capacity of the 100 cm³ metal calorimeter = $\frac{(heat energy lost) - (heat energy gained)}{(T_3 - T_2)}$ J °C⁻¹

Use your answers to (i) and (ii) and the temperature rise from (b) to calculate the approximate heat capacity of the 100 cm^3 metal calorimeter.

approximate heat capacity of the 100 cm³ metal calorimeter $=\frac{4620 - 3885}{18.5}$ = 39.72 $\approx 39.7 \text{ J} ^{\circ}\text{C}^{-1}$

The approximate heat capacity of the 100 cm³ metal calorimeter = 39.7... J °C⁻¹. [1]

(i) and (ii) Award one mark if both of the following expressions are correctly evaluated. heat gained = $210 \times \text{candidate}$ value of $(T_3 - T_2)$ heat lost = $210 \times \text{candidate}$ value of $(T_1 - T_3)$ *Units should be consistent. Ignore any sign given.* (iii) no mark.

(d) Determining the enthalpy change of combustion of ethanol, C₂H₅OH

An experiment was carried out as follows to determine the enthalpy change of combustion of ethanol.

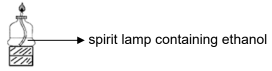
The 100 cm³ metal calorimeter used in (a) was used to contain some water. The water was heated using ethanol placed in a spirit lamp. The temperature rise was recorded. The spirit lamp was weighed before and after the experiment to determine the mass of ethanol used. The following results were obtained.

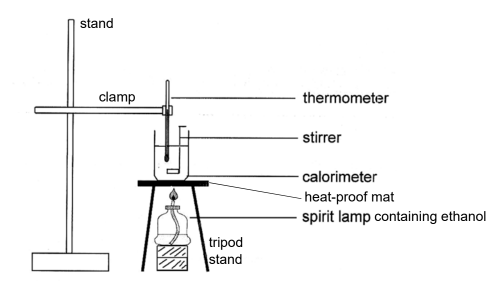
mass of ethanol burned = 0.391 g mass of water heated = 40.0 g temperature rise = 19.5 °C

(i) Complete the following diagram to show the experimental set-up needed to carry out the experiment to determine the enthalpy change of combustion of ethanol.

You are to use the following apparatus to complete the diagram:

- stand and clamp
- 100 cm³ metal calorimeter
- thermometer
- tripod stand and heat-proof mat





(ii) Use your answer in (c)(iii) and the results obtained to calculate the enthalpy change of combustion of ethanol. [Ar: O, 16.0; C, 12.0; H, 1.0]
[specific heat capacity of water = 4.2 J °C⁻¹ cm⁻³]
(If you were unable to answer (c)(iii), you may assume that the heat capacity of the 100 cm³ metal calorimeter is 50 J °C⁻¹.)

 $\Delta H_{\rm c}(\text{ethanol}) = -\frac{(40.0 \times 4.2 \times 10^{-3} \times 19.5) + (39.72 \times 10^{-3} \times 19.5)}{0.391/46.0}$ = -477 kJ mol⁻¹

enthalpy change of combustion of ethanol = -477 kJ mol^{-1}[1]

(iii) Calculate the maximum percentage error in the measurement of each mass used in the experiment.

mass measured	maximum error in a single reading	maximum percentage error / %
0.391 g of ethanol burned	0.0005 g	(0.0005×2)/0.391*100 = 0.256 %
40.0 g of water	0.05 g	0.05/40.0*100 = 0.125 %

[1]

[Total: 11]

3 Qualitative Analysis

At each stage of any test you are to record details of the following.

- colour changes seen
- the formation of any precipitate
- the solubility of such precipitates in an excess of the reagent added

Where gases are released they should be identified by a test, **described in the** appropriate place in your observations.

You should indicate clearly at what stage in a test a change occurs. No additional tests for ions present should be attempted.

If any solution is warmed, a boiling tube MUST be used.

Rinse and reuse test-tubes and boiling tubes where possible.

Where reagents are selected for use in a test, the name or correct formula of the element or compound must be given.

FB 4 is a solution containing **one** cation and **one** anion from the Qualitative Analysis Notes in pages 22 and 23.

Solid **FB 5** contains the sodium ion, and **one** anion from the Qualitative Analysis Notes in pages 22 and 23.

(a) Carry out the following tests on **FB 4** and state the observations you make.

(i)	To 1 cm depth of aqueous FB 4 in a test-tube, add aqueous ammonia the	White ppt soluble in excess NH ₃ (aq) [M1]
	add sulfuric acid dropwise until there is no further observations.	White ppt reforms, soluble in excess [M2]
(ii)	To 1 cm depth of aqueous FB 4 in a test-tube, add silver nitrate,	White ppt formed, soluble in excess NH ₃ (aq) [M3]
	followed by aqueous ammonia.	

For Examiner's Use			
Obs points	9-11 pts 6 mks 7-8 pts 5 mks 5-6 pts 4 mks 4 pts 3mks 3 pts 2 mks 1-2 pts 1 mk	marks	

(b) Carry out the following tests on **FB 5** and state the observations you make.

	test	observations
(i)	Place half of the solid FB 5 in a boiling tube. Heat strongly until the solid melts and a gas is given off.	White solid melts to give colourless liquid. [M4]
	Test and identify the gas.	Colourless, odourless gas relights a glowing splint. Gas is O ₂ . [M5]
	Continue the strong heating for $2 - 3$ min,	
	then leave the tube to cool and retain the residue for the test in (iv) .	White residue obtained on cooling [M6]
(ii)	To the remaining half of FB 5 is the boiling tube, add 3 cm depth of aqueous sodium hydroxide.	No NH₃ gas given off. [M7]
	Warm gently and retain for the test in (iii) .	
(iii)	To the solution from the test in (ii) , add a piece of aluminium foil and warm gently.	Pungent colourless gas turns moist red litmus blue. (Gas is NH ₃ .) [M8]
(iv)	Dissolve the residue from test in (i) in distilled water and divide the solution into three parts. Use these for the tests in (v) to (vii).	
(v)	To one part of the solution from (iv), add aqueous potassium iodide followed by dilute sulfuric acid.	Solution turns yellow/brown/black solid formed [M9]
(vi)	To the second part of the solution from (iv), add aqueous potassium manganate (VII) followed by dilute sulfuric acid.	Purple MnO ₄ ⁻ solution decolourised [M10]
(vii)	To the third part of the solution (iv), add aqueous sodium hydroxide and a piece of aluminium foil. Warm gently.	Pungent colourless gas turns moist red litmus blue. (Gas is NH ₃ .) [M11]

- (C) (i) Suggest the identities of the cation and anion present in **FB 4** from your observations from the tests in (a). cation: anion: [1] Zn²⁺ & Cl₇ [1] (ii) Write equations, with state symbols to explain your observations for test in a(i).
 -[3]

When aq NH_3 is added, a white ppt of $Zn(OH)_2$ is formed.

- Zn²⁺ + 2OH⁻ Ý Zn(OH)₂ (1)
- When excess aqueous NH₃ is added, ligand exchange takes place. NH₃ ligand replaces water ligand from $[Zn(H_2O)_6]^{2+}$ to form $[Zn(NH_3)_4]^{2+}$ which is a colourless solution.
- $[Zn(H_2O)_6]^{2+} + 4NH_3 \circ [Zn(NH_3)_4]^{2+} + 6H_2O$ (2) Accept $[Zn(NH_3)_6]^{2+}$ and $[Zn(NH_3)_4(H_2O]_2]^{2+}$ and pls mark equation (2) accordingly.
- This causes [Zn(H₂O)₆]²⁺ to drop, hence equilibrium (1) shifts to the left, and the precipitate, Zn(OH)₂ dissolves.

(The drop in $[Zn(H_2O)_6]^{2+}$ also causes the ionic product of $Zn(OH)_2$ to decrease to the extent that IP of $Zn(OH)_2 < Ksp$ of $Zn(OH)_2$. Hence the precipitate, $Zn(OH)_2$ dissolves.)

- When H₂SO₄(aq) was added, [NH₃] decreases, hence equilibrium (1) shifts to the left, and the precipitate, Zn(OH)₂ reforms.
- precipitate dissolves due to <u>further</u> removal of OHE by the <u>excess</u> $H_2SO_4(aq)$, causing the POE of reaction (2) to shift to the left.

3 marks for 6 bullets correct

- 2 marks for 3, 4-5 bullets correct
- 1 mark for 1-2 bullets correct

1 mark

(d)

(c) Student C carried out the following experiment on FB 5, predict the observations that will be observed and write your answer in the blanks.

Experiment	Observation
To solid FB 5 in a boiling tube. Heat heat strongly for 4 min. Let the residue cool. Add dilute sulfuric acid to it.	

[1]

[Total: 16]

Qualitative Analysis Notes [ppt. = precipitate]

(a) Reactions of aqueous cations

a a ti a m	reaction with		
cation	NaOH(aq)	NH ₃ (aq)	
aluminium, Al ³⁺ (aq)	white ppt. soluble in excess	white ppt. insoluble in excess	
ammonium, NH₄⁺(aq)	ammonia produced on heating	_	
barium, Ba ²⁺ (aq)	no ppt. (if reagents are pure)	no ppt.	
calcium, Ca ²⁺ (aq)	white ppt. with high [Ca ²⁺ (aq)]	no ppt.	
chromium(III), Cr³⁺(aq)	grey-green ppt. soluble in excess giving dark green solution	grey-green ppt. insoluble in excess	
copper(II), Cu ²⁺ (aq)	pale blue ppt. insoluble in excess	blue ppt. soluble in excess giving dark blue solution	
iron(II), Fe ²⁺ (aq)	green ppt., turning brown on contact with air insoluble in excess	green ppt., turning brown on contact with air insoluble in excess	
iron(III), Fe³⁺(aq)	red-brown ppt. insoluble in excess	red-brown ppt. insoluble in excess	
magnesium, Mg ²⁺ (aq)	white ppt. insoluble in excess	white ppt. insoluble in excess	
manganese(II), Mn²⁺(aq)	off-white ppt., rapidly turning brown on contact with air insoluble in excess	off-white ppt., rapidly turning brown on contact with air insoluble in excess	
zinc, Zn ²⁺ (aq)	white ppt. soluble in excess	white ppt. soluble in excess	

25

(b) Reactions of anions

anion	reaction
carbonate, CO ₃ ^{2–}	CO ₂ liberated by dilute acids
chloride, ClÈ (aq)	gives white ppt. with Ag⁺(aq) (soluble in NH₃(aq))
bromide, BrÈ(aq)	gives pale cream ppt. with Ag ⁺ (aq) (partially soluble in $NH_3(aq)$)
iodide, IÈ(aq)	gives yellow ppt. with Ag⁺(aq) (insoluble in NH₃(aq))
nitrate, NO₃È(aq)	NH_3 liberated on heating with $OH\dot{E}(aq)$ and Al foil
nitrite, NO₂È(aq)	NH ₃ liberated on heating with OH \dot{E} (aq) and A <i>l</i> foil; NO liberated by dilute acids (colourless NO \rightarrow (pale) brown NO ₂ in air)
sulfate, SO₄²⁻(aq)	gives white ppt. with Ba ²⁺ (aq) (insoluble in excess dilute strong acids)
sulfite, SO ₃ ²⁻(aq)	SO ₂ liberated with dilute acids; gives white ppt. with Ba ²⁺ (aq) (soluble in dilute strong acids)

(c) Tests for gases

gas	test and test result	
ammonia, NH₃	turns damp red litmus paper blue	
carbon dioxide, CO ₂	gives a white ppt. with limewater (ppt. dissolves with excess CO ₂)	
chlorine, Cl ₂	bleaches damp litmus paper	
hydrogen, H ₂	"pops" with a lighted splint	
oxygen, O ₂	relights a glowing splint	
sulfur dioxide, SO ₂	turns aqueous acidified potassium manganate(VII) from purple to colourless	

(d) Colour of halogens

halogen	colour of element	colour in aqueous solution	colour in hexane
chlorine, Cl ₂	greenish yellow gas	pale yellow	pale yellow
bromine, Br ₂	reddish brown gas / liquid	orange	orange-red
iodine, I	black solid / purple gas	brown	purple